Glycosylation and Cross-linking in Bone Type I Collagen

Monograph

Fibrillar type I collagen is the major organic component in bone, providing a stable template for mineralization. During collagen biosynthesis, specific hydroxylysine residues become glycosylated in the form of galactosyl- and glucosylgalactosyl-hydroxylysine. Furthermore, key glycosylated hydroxylysine residues, α1/2-87, are involved in covalent intermolecular cross-linking. Although cross-linking is crucial for the stability and mineralization of collagen, the biological function of glycosylation in cross-linking has not yet been understood. In this study, we quantitatively characterized glycosylation of non-cross-linked and cross-linked peptides by biochemical and nanoscale liquid chromatography-high resolution tandem mass spectrometric analyses. The results showed that glycosylation of non-cross-linked hydroxylysine is different from that involved in cross-linking. Among the cross-linked species involving α1/2-87, divergent cross-links were glycosylated with both monoglycosylated, whereas the mature, trivalent cross-links were primarily monoglycosylated. Markedly diminished diglycosylation in trivalent cross-links at this axis was also confirmed in type II collagen. The data, together with our recent report (Sricholpech, M., Perdivara, I., Yokoyama, M., Nagaoka, H., Terajima, M., Tomer, K. B., and Yamauchi, M. (2012) Lysyl hydroxylase 3-mediated glucosylation in type I collagen: molecular loci and biological significance. J. Biol. Chem. 287, 22998–23009), indicate that the extent and pattern of glycosylation may regulate cross-link maturation in fibrillar collagen.

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The abbreviations used are: Hyl, hydroxylysine; G, galactosyl; GG, glucosylgalactosyl; C-telo, C-telopeptide(s); N-telo, N-telopeptide(s); Pyl, pyridinoline; d-Pyr, deoxypyridinoline; Prl, pyrrole; DHLNL, dihydroxylysinonorleucine; HLNL, hydroxylysinonorleucine; EC, Ehrlich's chromogen; EIC, extracted ion chromatogram; Hyp, hydroxyproline; ESI, electrospray ionization; nanoLC, high performance/high resolution nanoscale liquid chromatography.
and α2 N-telo, respectively). The aldehyde can then form an iminium bond to a juxtaposed ε-amino group of Lys/Hyl in the helical domain of the staggered adjacent molecule. The possible combinations are as follows: α1-16G × α1-87, α1-16G × α1-87, α1-9N × α1-930, α1-9N × α2-933, α2-5N × α1-930, and α2-5N × α2-933. Some of these combinations are preferred for cross-linking (27). In skeletal tissues, these divergent cross-links can then mature into trivalent cross-links, pyridoline (Pyr), deoxypyridinoline (d-Pyr), and pyrrrole (Prl) (1, 28), tying two or three collagen molecules (29).

Despite current knowledge regarding type I collagen glycosylation (14, 30), the precise molecular loci and the extent/type of glycosylation in cross-linked and non-cross-linked residues are still not well characterized. This information, however, is essential to understand the role of glycosylation in collagen biosynthesis and function. Mass spectrometry (MS) has become increasingly used in the structural characterization of collagens from different sources, and several studies have utilized MS to characterize collagen cross-linked peptides (30–34). Although these studies pioneered the MS-based characterization of collagen cross-linked species, their major limitation is the use of low resolution mass analyzers. This renders accurate characterization of species bearing naturally occurring heterogeneity (e.g., incomplete hydroxylation of Lys or Pro) or of species with similar mass/charge ratio (m/z), difficult. In the present study, we used high performance/high resolution nanoscale liquid chromatography-tandem mass spectrometry (nanoLC/MS/MS) to comprehensively characterize the glycosylation at various molecular loci in non-cross-linked and cross-linked peptides in bovine bone type I collagen. A multistep chromatographic approach was employed to obtain highly purified cross-linked tryptic peptides. The analytical challenges associated with these large cross-linked species were overcome with the use of an alternative enzyme. The molecular distribution of glycosylation in non-cross-linked Hyl and immature and mature cross-links was quantitatively determined by nanoLC/MS. The results revealed a differential glycosylation pattern, depending on the involvement in cross-linking, molecular loci, and type and maturational stage of cross-linking.

**EXPERIMENTAL PROCEDURES**

**Collagen Preparation**—Fresh femoral bone samples from 2–3-year-old bovine animals were obtained commercially (Aries Scientific, Dallas, TX). After removing the surrounding connective tissues, both ends of the bone, and the bone marrow, the bones were cut into small pieces. All operations were carried out at 4 °C. The bone pieces were pulverized to a fine powder under liquid nitrogen using a Spex Freezer Mill (Spex, Inc., Metuchen, NJ). Pulverized samples were washed several times with cold phosphate-buffered saline (PBS), and cold distilled water, centrifuged at 4000 × g for 30 min, and lyophilized. Bone powder was then demineralized with 0.5 M EDTA, pH 7.5, for 2 weeks with several changes of the EDTA solution by centrifugation at 4000 × g. The EDTA-insoluble residue was thoroughly washed with cold distilled water by repeated centrifugation at 4000 × g and lyophilized.

**Reduction with NaBH₄**—Demineralized bone (~2.0 g) was suspended in buffer containing 0.15 M N-trimethyl-2-amino-ethanesulfonic acid and 0.05 M Tris-HCl, pH 7.4, and reduced with standardized NaBH₄. The specific activity of the NaBH₄ was determined by the method described previously (35, 36). The reduced samples were washed with cold distilled water several times by repeated centrifugation at 4000 × g and lyophilized. Upon reduction, the dehydrodihydroxylysinonorleucine (dehydro-DHLNL) and dehydrohydroxylysino norleucine (dehydro-HLNL) and their respective keto amine forms are reduced to stable secondary amines, DHLNL and HLNL, and radiolabeled simultaneously (+2 Da molecular mass increase). Hereafter, the terms DHLNL and HLNL will be used for both the unreduced and reduced forms.

**Cross-link Analysis**—Reduced collagen was hydrolyzed with 6 N HCl and subjected to cross-link analysis as described previously (37). The reducible cross-links were analyzed as their reduced forms (i.e. DHLNL and HLNL, respectively). The levels of the immature reducible (DHLNL and HLNL) and mature non-reducible cross-links (Pyr and d-Pyr) were quantified and expressed in mol/mol of collagen.

**Digestion with Trypsin**—Digestion with trypsin of reduced bone collagen was prepared by the procedure described previously (37) with slight modifications. Briefly, the reduced collagen was heated at 65 °C for 15 min, digested with 1% (w/w) trypsin for 16 h at 37 °C, rehydrated to 65 °C for 10 min, and retreated with 0.5% (w/w) trypsin for 3 h at 37 °C. Over 99% of the starting material was recovered in the supernatant of the trypsin digest.

**Molecular Sieve Chromatography**—Molecular sieve chromatography of the bone tryptic digest was performed on a HiLoad Superdex 75 preparative scale column (1.6 × 60 cm) equilibrated with 0.05 M ammonium bicarbonate (pH 7.9) at room temperature. Aliquots of the reduced bone tryptic digest (~200 mg) were injected and separated at a flow rate of 0.75 ml/min. Fractions of 1.5 ml were collected, and their absorbance (230 nm), radioactivity, and fluorescence (excitation 330 nm and emission 390 nm) were measured. Fluorescence measurement was employed to detect Pyr and d-Pyr, and radioactivity was used to detect DHLNL and HLNL. The main fluorescent (F1–F3) and radioactive (R1 and R2) fractions were recovered. An equal aliquot from each fraction was lyophilized and subjected to cross-link analysis as described above. The majority of aliquots were subjected to further purification by reversed phase chromatography or digestion with chymotrypsin (see below). To identify fractions containing the Prl cross-link, aliquots from each fraction were subjected to Ehrlich’s chromogen (EC) analysis following reported methods (30, 38). Briefly, the aliquots were lyophilized and rehydrated in distilled water, and 35 μl of a 5% solution (w/v) of p-dimethylaminobenzaldehyde in 4 M perchloric acid were added. The reaction was incubated for 5 min at room temperature. The absorbance (572 nm) was monitored with a F-2000 fluorescence spectrophotometer (Hitachi, Tokyo, Japan). The main EC-positive fractions (P1–P3) eluted at similar positions as F1–F3 and were further characterized by mass spectrometry (see below).

**Chymotrypsin Digestion**—Chymotrypsin digestion of the fractions F1–F3, R1, and R2 collected after molecular sieve chromatography was performed as follows. The lyophilized fractions were dissolved in 100 μl of 25 mM ammonium bicarbonate (pH 7.4). Aliquots of 50 μl from each fraction were incu-
bated with 2 μl of chymotrypsin solution (0.5 μg/μl in 25 mM ammonium bicarbonate) overnight at room temperature. These digests were further analyzed by nanolC/MS/MS.

Reversed Phase High Performance Liquid Chromatography—
The lyophilized fluorescent and radioactive fractions resolved by molecular sieve chromatography were further fractionated by reversed phase on a SOURCE 5RPC ST 4.6/150 column (GE Healthcare) using a Varian HPLC system (Prostar 240/310, Varian, Walnut Creek, CA). The solvents employed were 2% (v/v) acetonitrile in 10 mM ammonium acetate (solvent A) and 70% acetonitrile in deionized water (v/v) (solvent B). The samples were dissolved in solvent A and eluted with a linear gradient from 0 to 15% solvent B for the first 10 min, followed by a linear gradient from 15 to 50% solvent B over the next 50 min at a flow rate of 1.0 ml/min at room temperature. The effluent was monitored for absorbance (230 nm) and fluorescence (excitation 330 nm, emission 390 nm). Fractions of 1 ml were collected, and aliquots were subjected to radioactivity and fluorescence measurements. The radioactive and fluorescent fractions were pooled, lyophilized, and subjected to further analysis.

Ion-Exchange Chromatography—Ion-exchange chromatography of fractions collected after reversed phase separation was performed on a TSK-GEL DEAE-5PW column (8.0 × 75 mm, Tosoh Bioscience LLC, Montgomeryville, PA). The column was equilibrated with 0.01 M NH₄HCO₃ containing 1% isoprolyl alcohol. Elution was carried out with a linear gradient from 0.01 to 0.25 M NH₄HCO₃ at a flow rate of 1 ml/min, monitoring the absorbance at 230 nm. One-minute fractions were collected, and those corresponding to fluorescent and/or radioactive signals were pooled and lyophilized. Their structural characterization was performed by mass spectrometry.

Nanoscale Liquid Chromatography, Mass Spectrometry, and Data Analysis—Glycosylated, non-cross-linked collagen peptides were characterized by nanolC/MS/MS from a bovine bone trypptic digest on a nanoACQUITY UPLC-Q-Tof Premier mass spectrometer (Waters, Milford, MA). Their structure- and site-specific quantitative analyses were performed as described previously (24).

Flow Injection Analyses of Purified Cross-linked Tryptic Peptides—Flow injection analyses of purified cross-linked tryptic peptides were performed by positive ion nanoelectrospray (ESI⁺) on a Waters Micromass Q-Tof Micro mass spectrometer (Waters). The samples were desalted using a C18 ZipTip pipette tip (EMD Millipore, Billerica, MA) and reconstituted in 50% acetonitrile with 0.1% formic acid (v/v). Dilutions were performed as needed to ensure optimal signal intensity. Mass spectrometer parameters were as follows: capillary voltage, 3.4 kV; sampling cone, 30; source temperature, 80 °C; desolvation temperature, 20 °C.

NanolC/MS/MS Analyses—NanolC/MS/MS analyses of cross-linked peptides were carried out on a nanoACQUITY UPLC-Q-Tof Global mass spectrometer (Waters) with data-dependent acquisition and charge state selection of the top four ions. Separations of chymotryptic digests of fractions after molecular sieve chromatography were carried out on a C18 BEH column (1.7 μm, 75 μm × 100 mm) at 0.3 μl/min, with a gradient from 1 to 50% solvent B over 30 min. The solvents were as follows: solvent A (0.1% formic acid in deionized water) and solvent B (0.1% formic acid in acetonitrile). Mass spectrometer settings for (+)-nano-ESI were as follows: capillary voltage, 3.4 kV; sampling cone, 30; source temperature, 80 °C. To ensure optimal fragmentation, the collision energies were optimized over the range 20–30 V.

Data Analysis—Data analysis was performed using the MassLynx software, version 4.1 (Waters), including the embedded deconvolution algorithms MaxEnt 1 and 3. The glycoform distribution of C-telo-derived cross-linked peptides was determined from the LC/MS data of chymotryptic digests. Quantitative glycosylation analyses (percentage of free, galactosyl (G), and glucosylgalactosyl (GG)) of cross-linked peptides were performed as described for non-cross-linked peptides by integrating the extracted ion chromatograms (EICs) generated post-data acquisition (24). Deconvolution of MS/MS spectra of the precursor ion of m/z 677.281 (5+), assigned to peptide α2-(928–963). The following modifications are consistent with the fragment ion spectrum: Hyl-933, Hyp-920, Hyp-942, Hyp-954, and deamidated Asn-936 (indicated as N→D). Site α2-Hyl-933 is involved in intermolecular cross-linking. Within the non-cross-linked structures containing residue α2-Hyl-933 (i.e. spectrum shown above), no glycosylation was observed in bovine bone.
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RESULTS

Glycosylation of Non-cross-linked Peptides—To identify glycosylation sites in type I collagen, a tryptic digest of reduced bone collagen was analyzed by nanoLC/MS/MS on a Waters nanoACQUITY UPLC-Q-Tof Premier mass spectrometer (24, 39). Five glycosylation sites were characterized, as follows: α1-/α2-87, observed in glycopeptides α1-/α2-(76–90); α1-/α2-174 in glycopeptides α1-(145–183) and α2-(145–192), respectively; and α2-219 in glycopeptide α2-(193–237). The distribution of Lys, Hyl, and G- and GG-Hyl at each site was determined as described previously (24), and the results are summarized in Table 1. The highest relative extent of glycosylation was found for residues α1-/α2-87 (i.e., 68% G and 13% GG form for α1-87, and 63.1% G and 18.5% GG form for α2-87, respectively). Site α2-174 was found mostly in the form of G (56%) with a minor amount of GG (2.9%) and with a considerable level of non-hydroxylated/non-glycosylated Lys (29.3%). Site α2-219 was found to contain 20% (Lys), 42.6% (Hyl), 29.9% (G-Hyl), and 7.5% GG-Hyl. Site α1-174 was found minimally modified by glycosylation, with only 1.5% G-Hyl and noise level GG-Hyl, with 49.6% non-glycosylated/non-hydroxylated Lys. Residues α1-/α2-87, the major glycosylation sites (24), are also major helical cross-linking sites, whereas the other three glycosylated residues are not involved in cross-linking.

Two additional helical Hyl residues involved in cross-linking are α1-930 and α2-933. The tryptic peptide containing α1-930 spans the residues 928 GI-Hyl 930 GHR 933. This peptide was not

FIGURE 2. A, molecular sieve elution profile of a tryptic digest of NaB₃H₄-reduced mature bone collagen. Eight peaks were collected, as follows: F1–F3 (fluorescent), R1 and R2 (radioactive content), and P1–P3 (EC-reactive species, absorbance at 572 nm). B and C, reversed phase purifications of peaks F1 and R1, resulting in peak 1 (fluorescent) and peaks 2 and 3 (radioactive), respectively. D–F, ion exchange purification of peaks 1, 2, and 3, resulting in fluorescent peak a and radioactive peaks b and c, respectively.
TABLE 2
Quantitative cross-link analysis (in mol/mol of collagen) of immature and mature cross-links in bovine bone type I collagen and estimated values of the C-telo-derived cross-links

| Cross-link | Immature | Mature |
|-----------|----------|--------|
| PLNL      | 1.58     | 2.07   |
| HLNL      | 0.50     | 0.67   |
| Pyr       | 0.28     | 0.32   |
| d-Pyr     | 0.04     | 0.05   |

The Mₐ of immature cross-linked peptides was determined by accounting for the incorporation of two ³H atoms, as a result of reduction with NaB₃H₄.

TABLE 3
Theoretical molecular weight (Mₐ) determination of collagen cross-linked peptides in bone

| Equation | Immature | Mature |
|----------|----------|--------|
| Mₐ(DHNL) = Mₐ(peptide 1) + Mₐ(peptide 2) - 16 | 1 | 1 |
| Mₐ(HLNL) = Mₐ(DHNL) - 16 | 2 | 2 |
| Mₐ(Pyr) = Mₐ(peptide 1) + Mₐ(peptide 2) + Mₐ(peptide 3) - 57 | 3 | 3 |
| Mₐ(d-Pyr) = Mₐ(Pyr) - 16 | 4 | 4 |
| Mₐ(pyrrole) = Mₐ(Pyr) - 15 | 5 | 5 |

FIGURE 3. MS spectra of pyridinoline cross-linked tryptic peptides α₁(993–22°C) × α₁(993–22°C) × α₁(76–90) (A) (inset shows the masses obtained by deconvoluting the m/z range 1000–2600); DHLNL α₁(993–22°C) × α₁(76–90) (B); and DHLNL α₁(993–22°C) × α₂(76–90) (C). Glycosylation is indicated by G and GG. These spectra were obtained by flow injection analysis of fractions purified by multistage chromatography.

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observed by LC/MS/MS, most likely due to its short length. Residue α₂-933 was observed in peptide α₂(928–963) and assigned based on the MS/MS of m/z 677.281 (5+) (supplemental Fig. S1). The following modified residues are consistent with this fragment ion spectrum: Hyp-930, -942, and -954, Hyl-933, and deamidated Asn-936 (Fig. 1). Residue α₂-933 was observed non-glycosylated. Both Asn-936 and the deamidated form were observed in peptide α₂(928–963), in an approximate Asp/Asn ratio of 3:1. This is relevant for the analysis of cross-linked peptides containing α₂(928–963) (see below).

Isolation, Molecular Characterization, and Glycosylation of Cross-linked Peptides—Using a NaB₃H₄-reduced collagen trypptic digest as starting material, a multistep chromatographic approach was employed to obtain highly purified tryptic cross-linked peptides. Three sequential chromatographic steps were performed: 1) molecular sieve; 2) reversed phase; and 3) ion exchange. After each step, the fractions containing immature and mature cross-linked peptide were detected based on radioactivity and fluorescence, respectively (Fig. 2), whereas Prl cross-links were detected by reaction with EC (see “Experimental Procedures”) (40). The chromatographic profiles are shown in Fig. 2, with the major fluorescent and radioactive peaks designated as F₁–F₃ and R₁ and R₂, respectively. Cross-link analysis performed after molecular sieve separation was in agreement with the profiles shown in Fig. 2 (i.e. Pyr/d-Pyr were found in fractions F₁–F₃, whereas DHLNL/HLNL were found in R₁ and R₂). Quantitative cross-link analysis on bone collagen (Table 2) indicated that DHLNLs are the most abundant (1.58 mol/mol of collagen), followed by HLNL (0.50 mol/mol) and Pyr (0.28 mol/mol), whereas d-Pyr was found in low amounts (0.04). Based on these numbers, the molecular loci identified in F₁–F₃ and R₁ and R₂ (i.e. R₁ and F₁ contain the C-telo derived cross-links; see below), and direct cross-link analyses of these

TABLE 4
Glycoform distribution (free, G, and GG)

| Glycosylated form (%) | Free | G- | GG- |
|-----------------------|------|----|-----|
| DHLNL (α₁C × α₁B)    | 63 (1)| 40 (3)| 25 (2)|
| DHLNL (α₁C × α₂B)    | 3 (5) | 48 (5)| 49 (6)|
| Pyr (α₁C × α₁C × α₁B)| 6 (3) | 81 (5)| 13 (2)|
| Pyr (α₁C × α₁C × α₂B)| 17 (1)| 55 (3)| 28 (4)|
| Prl (α₁C × α₁C × α₁B) | ND* | 80 (4)| 20 (4)|
| Prl (α₁C × α₁C × α₂B) | ND Detected at low abundance/not quantifiable |

% relative abundance

| Free | G- | GG- |
|------|----|-----|
| α₁C × α₁C × α₁B | 85 (3) | 15 (3)|

* ND, not detected.

** Identified, but due to the low abundance, the relative glycoforms were not estimated (see "Results").
fractions, we estimated that within DHLNLs, the C-telo-derived (α1C/α1C-87) cross-link was 4-fold higher than that of the N-telo-derived (α1C/α1C) (R9251 4.3 versus 0.3), whereas the levels of C- and N-telo-derived HLNLs were comparable. Pyr was almost equally located between C- and N-telo, whereas d-Pyr was enriched in the N-telo (80% of total d-Pyr). The relative small fluorescence of the N-telo Pyr/d-Pyr (F2 and F3) in Fig. 2 is probably due to the fluorescence quenching (30). From F1 and R1 fractions, after the final chromatographic step, one fluorescent (a) and two radioactive peaks (b and c) were collected (Fig. 2, D–F) and were analyzed by flow injection nano-ESI-MS.

**Flow Injection Analysis-Mass Spectrometry of Peaks a, b, and c**—The theoretical molecular weights of collagen cross-linked peptides were determined using the equations listed in Table 3. The raw mass spectrum of the cross-linked species contained in peak a is shown in Fig. 3A. The charge state envelope containing the ions of m/z 2532.62 (4+), 2026.27 (5+), 1688.75 (6+), 1447.65 (7+), 1266.81 (8+), and 1126.14 (9+) was deconvoluted to the average mass of 10,126.4 Da (Fig. 3A, inset). This value is consistent with the theoretical average molecular weight of G-Pyr cross-linked tryptic peptides α1-(993–22C) × α1-(76–90). The peptide identities were confirmed by MS/MS (not shown). Ion clusters of species with a molecular mass of 162 Da lower or higher than G-Pyr, corresponding to Pyr and GG-Pyr, respectively, were observed with very low relative abundance, indicating that G is the predominant glycoform of the C-telo-derived Pyr cross-link.

The deconvoluted mass spectra of the DHLNL cross-linked peptides α1-(993–22C) and α1-(76–90) observed in peaks b and c are shown in Fig. 3B and C, and Supplemental Fig. S2, respectively. The ratio of the former to the latter was 3:1 (Fig. 2C), which is consistent with our previous report (36). In contrast to Pyr, both the G and GG glycoforms of α1α2-87-containing DHLNLs represent major species. As shown in Fig. 3, B and C, extended molecular het-
The occurrence of the latter was confirmed in the fractions b and c (i.e. when the fractions were hydrolyzed and directly analyzed for cross-links, HLNL was present as a minor species in both fractions). These low abundance species are difficult to assign because their MS/MS lack informative fragment ions. Nevertheless, the species of molecular mass 5791.96 Da (Fig. 3C) probably represent the HLNL (α1-16C × α2-87). This would be in agreement with a previous study suggesting higher...
abundance of HLNL (α1-16C × α2-87) compared with HLNL (α1-16C × α1-87) (41).

Quantitative Determination of Glycosylation in the C-telopeptide Containing Immature and Mature Cross-linked Peptides—For LC/MS-based quantitative glycosylation analysis of the C-telo cross-linked peptides, the multistage chromatographic approach was modified as follows. 1) the NaB3H4-reduced bone tryptic digest was separated by molecular sieve chromatography as described above. Fractions were collected based on fluorescence (F1–F3) and radioactivity (R1 and R2). 2) the tryptic peptide fractions F1–F3, R1, and R2 were digested with chymotrypsin to reduce the size of the cross-linked peptides. 3) the resulting double digests were analyzed by nanoLC/MS/MS.

The major chymotryptic product was formed by cleavage in the C-telopeptide between Tyr and Asp, resulting in the α1C sequence DLSFLPQPPQEKAHGDGR, whereas the helical peptides (residues 76–90) remained undigested. All mature cross-link species, including C-telo Pyr/Prl (α1C × α1C × α1b/α2b) were identified in the chymotryptic digests of F1/P1 by nanoLC/MS/MS. Prl (α1C × α1C × α2b) was identified as G-Prl, but, due to the low abundance, its relative glycoforms (free, G-, and GG-) were not quantitatively estimated (Table 4A). The average molecular mass of G-Pyr (α1C × α1C × α1b) was the most abundant glycoform (81%), whereas GG-Pyr (m/z 963.634 (6+)) and non-glycosylated Pyr (m/z 909.607 (6+)) had markedly lower abundances (13 and 6%, respectively; Table 4A). The G-Prl (α1C × α1C × α1b) species, having a 15 Da lower molecular mass than G-Pyr, was observed as the ion of m/z 934.292 (6+). Furthermore, G- and GG-Pyr (α1C × α1C × α2b) were observed in chymotryptic fraction F1/P1 as the ions of m/z 947.969 (6+) (supplemental Fig. S3A) and 974.951 (6+), respectively. The MS/MS of the ion of m/z 947.969 (6+) is shown in Fig. 4B.

The C-telo DHNLNs were well characterized in both tryptic and chymotryptic forms. The MS/MS spectra of the G-DHNLN α1C × α1b/α2b protonated molecules (Fig. 4, C and D, and supplemental Fig. S3, C and D) confirm their structural assignment. In MS/MS, no neutral loss of G was observed, suggesting increased stability of the glycosidic bond of cross-linked peptides. Lower abundance glycosylated ions G- and GG-DHNLNs were assigned based on the 16 Da lower mass compared with the corresponding DHNLN glycoforms.

To determine the distribution of the free, G, and GG forms in immature/mature C-telo-containing cross-linked species, glycoform EICs were generated from the LC/MS of F1- and R1-chymotryptic digests and were normalized to the most abundant species within a cross-link type (Fig. 5). Quantitative analyses of technical triplicates are shown in Table 4. Within mature cross-links Pyr/Prl α1C × α1C × α1b/α2b, the G glycoform is the most abundant, whereas GG is minimal. The relative amounts of α1C free Pyr are higher in α2b compared with α1b. In contrast to mature cross-links, both the G and GG glycoforms of the DHNLN cross-links α1C × α1b/α2b are highly abundant, with minute amounts of free DHNLNs. The glycosylation patterns of HLNL α1C × α1b/α2b are comparable with those of DHNLNs. However, quantitative analyses of HLNLs and of α2b-Prl glycoforms were not performed because of poor signal/noise ratio.

The relative abundance of the C-telo/helical residue 87-based Pyr (i.e. the α1b/α2b ratio) was estimated from the LC/MS data of chymotryptic F1. The EICs of α1b and α2b G-Pyr (Fig. 6A) suggest that α1b-Pyr is far more abundant (~85%) than α2b-Pyr (~15%). This might explain why the α2b-Pyr was not observed previously by flow injection analysis of peaks a, b, and c. Moreover, we determined the relative abundance of Pyr/Prl containing α1-87 to be ~4:1 (Fig. 6B and Table 4B).

Characterization of N-telopeptide Containing Immature and Mature Cross-links—By flow injection analysis, the molecular mass of free (non-glycosylated) DHNLN α1N × α2b, determined as 5943.18 Da, is consistent with the structure of the reduced cross-link α1N-(15N–9) × α2b-(928–963) containing pyro-Gln in α1N peptide and three Hyp residues and deamidated Asn-936 in the α2b peptide. This assignment was confirmed from the MS/MS of the ion of m/z 991.530 (6+) (Fig. 7A and supplemental Fig. S4, A and B). The α1N × α2b was observed completely non-glycosylated, consistent with the profile of the non-cross-linked peptide α2-(928–963) (see above).

After molecular sieve separation, the free DHNLN α1N × α1b, consisting of tryptic peptides α1N-(15N–9) × α1b-(928–933), was identified in fraction R2 as the ions of m/z 811.878 (4+) and 649.703 (5+) (data not shown). The MS/MS spectrum of the precursor ion of m/z 807.866 (4+) (Fig. 7B and supplemental Fig. S4, A and B), assigned to reduced non-glycosylated HLNLN (α1N × α1b), confirms the identity of peptides α1N-(15N–9) and α1b-(928–933). The cross-links α1N × α1b were...
also found completely non-glycosylated. The N-telopeptide cross-link \( \alpha_2 \times \alpha_1 \) was not observed.

The mature N-telo cross-linked tryptic peptides were observed in fractions F2/P2 and F3/P3 after molecular sieve chromatography, with the major species assigned to non-glycosylated Pyr/Prl \( \alpha_1 \times \alpha_2 \times \alpha_1 \) in fraction F2/P2 and non-glycosylated Pyr \( \alpha_1 \times \alpha_2 \times \alpha_1 \) and \( \alpha_2 \times \alpha_1 \times \alpha_2 \) in fraction F3/P3. The average molecular mass of 7678.70 Da (Fig. 7C) is consistent with the calculated mass of \( \alpha_1 \times (1N–9) \times \alpha_2 \times (928–963) \), containing pyro-Gln in the \( \alpha_1 \times \alpha_2 \times \alpha_1 \) peptides and deamidated Asn-936 and three Hyp residues in the \( \alpha_2 \) peptide. Additional species at +16 Da most likely represent the Prl (−15 Da), whereas +16 Da might be due to Met oxidation or additional Pro hydroxylation. Peptide identities were established from the MS/MS of \( m/z 1097.83 \) (Fig. 7C and supplemental Fig. S4C). Furthermore, incomplete hydroxylation of Hyp-954 in \( \alpha_2 \times (928–963) \) is suggested by the fragment ion of \( m/z 878.49 \), which is 16 Da lower than \( y_{10h} \) \( m/z 894.48 \). These features and the fragment ion clusters at \( m/z 6781.37, 6782.18, \) and 6797.09 suggest increased molecular heterogeneity of Pyr/Prl \( \alpha_1 \times \alpha_2 \times \alpha_1 \times \alpha_2 \). Hence, the average mass of 7678.70 most likely arises from overlapping isotopic envelopes of Pyr/Prl \( \alpha_1 \times \alpha_2 \times \alpha_1 \times \alpha_2 \). Qualitatively, these data suggest that the Prl/Pyr ratio is higher in the N- than in C-telopeptide (Fig. 6). Most importantly, all immature and mature N-telo cross-links were found non-glycosylated.

**Glycosylation in the C-telopeptide-containing Immature and Mature Cross-linked Peptides from Type II Collagen**—Following the initial molecular sieve chromatography of reduced tryptic digests, the cross-link-containing fractions were characterized by nanoLC/MS as described above. The C-telo (type II collagen)-derived DHLNL \( \alpha_1(1N–9) \times \alpha_2(928–963) \) (top) and deconvoluted MS spectrum of cross-linked species \( \alpha_1(1N–9) \times \alpha_2(1N–9) \times \alpha_2(928–963) \), confirming the identity of the indicated cross-linked peptides.
Pyr, G-Pyr, and GG-Pyr (Fig. 8). Non-glycosylated Hyl is markedly lower abundance (i.e., 1%; Fig. 8). This is consistent with the above bone type I collagen data where it is one of the major forms in its precursor DHLNL. This is in contrast to collagen secreted from mouse calvaria-osteoblastic cells, where G-Hyl with lower relative amounts of GG-Hyl was found as GG-Hyl, and G- and GG-glycosylation between bovine and mouse could be related to species-specific variations in the collagen sequence (GFKGVK in mouse versus GFKGIR in bovine and other species), suggesting a certain substrate specificity for collagen galactosyltransferases (24).

DISCUSSION

Collagen glycosylation, consisting of the O-glycosides in the form of G- and GG-Hyl, is a key modification involved in collagen cross-linking, fibrillogenesis, mineralization, and collagen-protein interactions (1, 21, 24, 29, 43, 44). Type I collagen, the main organic component in bone, is one of the minimally glycosylated members in the collagen family (45). Even among type I collagen in various tissues, such as skin type I collagen, the extent of glycosylation in bone type I collagen is low (45, 46). Despite numerous studies indicating the functional importance of this modification in type I collagen biosynthesis, the type and distribution of Hyl glycosides and their involvement in cross-linking have not been characterized in a comprehensive manner. In this study, by employing a wide range of analytical methods, we performed residue-specific quantitative glycosylation analysis in both non-cross-linked and cross-linked peptide species from bovine bone type I collagen.

Five Hyl residues were found glycosylated in bovine bone type I collagen (i.e., α1-α2-87, α1-α2-174, and α2-219). Among the non-cross-linked peptides, the major glycosylation occurs at the helical cross-linking site, α1-α2-87, and is present mainly as G-Hyl with lower relative amounts of GG-Hyl. This is in contrast to collagen secreted from mouse calvaria-derived MC3T3-E1 osteoblastic cells, where α1-87 is mostly found as GG-Hyl, and α2-87 is >90% non-glycosylated Hyl (24). Differences in the α2-87 glycosylation between bovine and mouse could be related to species-specific variations in the α2 collagen sequence (GFKGVK in mouse versus GFKGIR in bovine and other species), suggesting a certain substrate specificity for collagen galactosyltransferases (24).

The C- and N-telo immature and mature cross-links were characterized in the form of tryptic/chymotryptic cross-linked peptides. In agreement with previous studies of bone collagen (24, 36), cross-link analysis suggested that the ~80% of DHLNL was derived from the C-telo site. This might represent a feature conserved among species, because a similar distribution was found in mouse osteoblasts as well (24). Cross-linked peptides containing DHLNL α1-16C × α1/α2-87 and HLN α1-16C × α1-87 were found mostly glycosylated in the form of both G and GG glycoforms having similar abundances. In contrast, the N-telo DHLNL α1-9N × α1-930, α1-9N × α2-933, and α2-5N × α2-933 were found non-glycosylated (30).
Glycosylation and Collagen Cross-linking

In contrast to the immature cross-links, the mature Pyr/Prl species ($\alpha1-16\times1-16\times1/2-87$) were found glycosylated primarily as G-Pyr, with minute amounts of the free and GG glycoforms. In this study, we identified the previously undescribed $\alpha2-87$-containing Pyr and Prl peptide. In comparison with the $\alpha1-87$-involved Pyr and Prl, this is a minute species representing ~15% of the former. We previously reported the residue $\alpha1-87$ to be the preferential cross-linking site over $\alpha2-87$ (~3:1) for the formation of C-telo-derived divalent cross-links in bone. Thus, the low abundance of the mature trivalent cross-links is not surprising. This is probably due to the specific molecular packing in the fibril (35), which could be important for collagen mineralization in an orderly fashion (36).

It is generally accepted that two residues of immature cross-links, DHLNL/HLNL, mature into one residue of trivalent cross-link, Pyr/d-Pyr, with aging (29, 43). However, our in vitro incubation study and the report by Saito et al. (47) showed that the decreased level of immature cross-links is disproportionally higher than the increase of Pyr (24). Most likely, the spontaneous non-enzymatic maturation is controlled by the microenvironment, such as presence of mineral (30, 48), the glycosylation state of the immature cross-links (35), and the presence of collagen-binding proteoglycans around the cross-linking sites. The data in the current study showing abundance of both G- and GG-DHLNL/HLNL but the predominance of G-Pyr and G-Prl forms suggests that GG divalent cross-links may not favor maturation into trivalent cross-links. This was supported by the data on type II collagen. In this heavily hydroxylated and glycosylated collagen type (49), we also found that ~50% of the Hyl-87 involving DHLNL is in the form of GG-DHLNL, whereas GG-Pyr at the same locus is minimal. Eyre et al. (32) also reported the lack of glycosylated Pyr at this locus (residue 87) in type II collagen. Possibly, the bulky disaccharide structure sterically hinders or delays the condensation reaction to form mature cross-links. Along with this conjecture, it is interesting to note that, for the N-telo-derived cross-links in which no glycosylated forms were found, immature cross-links were significantly lower, and mature cross-links were higher than the heavily glycosylated C-telo-derived cross-links. Possibly, without glycosylation of the helical Hyl, maturation of the N-telo-derived cross-links is accelerated. Clearly, further studies are warranted to determine the fate of glycosylated immature cross-links by employing, for instance, an in vitro maturation study (24) using type I and other types of collagen.

Recently, trivalent Prl cross-links having an identical chain topology with Pyr were identified (30, 50, 51). These were predominantly observed at the N-telo-to-helix site (30, 50, 51), concentrated at the loci $\alpha1-89\times\alpha2-5N\times\alpha2-933$ (30, 50) and $\alpha1-9N\times\alpha1-930/\alpha2-933$ (51). Here, we characterized the Prl species $\alpha1-16\times\alpha1-16\times\alpha1-87/\alpha2-87$ and $\alpha1-9N\times\alpha2-5N\times\alpha1-930/\alpha2-933$. Prl has a glycosylation pattern similar to that of Pyr (i.e. C-telo Prl contains mainly G, whereas N-telo Prl are non-glycosylated). This indicates there is no preference in the glycosylation pattern between Pyr and Prl cross-links.

In conclusion, this study provides a detailed molecular characterization of glycosylation and its involvement in intermolecular cross-linking in bovine bone type I collagen. Specific molecular loci and the differential glycosylation pattern between the immature and mature cross-links suggest that glycosylation might regulate the cross-link maturation in fibrillar collagen.

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