Posttranslational Modifications in Type I Collagen from Different Tissues extracted from wild type and Prolyl 3-hydroxylase 1 Null Mice

Elena Pokidysheva‡§, Keith D. Zientek‡, Yoshihiro Ishikawa‡§, Kazunori Mizuno‡, Janice A. Vranka‡, Nathan T. Montgomery‡§, Douglas R. Keene§, Tatsuya Kawaguchi†, Kenji Okuyama† and Hans Peter Bächinger‡§*

‡Research Department, Shriners Hospitals for Children, Portland, OR 97239
§Department of Biochemistry and Molecular Biology, Oregon Health & Science University, Portland, OR 97239
†Department of Macromolecular Science, Graduate School of Science, Osaka University, Osaka 560-0043, Japan

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Address correspondence to: Hans Peter Bächinger, Ph.D., Shriners Hospital for Children, Research Department, 3101 SW Sam Jackson Park Road, Portland, OR 97239, USA
Phone: (503) 221-3433, FAX: (503) 221-3451, E-mail: hpb@shcc.org

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Background: 3-hydroxylation of proline residues in type I collagen is rare but important. Results: 3Hyp sites have been identified in both chains of mouse type I collagen in wild type and P3H1 null mice. Conclusion: The absence of 3Hyp does not alter the D-period of collagen fibrils, but alters the lateral growth of the fibrils. Significance: Type I collagen prolyl 3-hydroxylation is tissue specific.

SUMMARY
Type I collagen extracted from tendon, skin and bone of wild type and prolyl 3-hydroxylase 1 (P3H1) null mice show little difference compared to wild type. These results demonstrate that the A1 site in type I collagen is exclusively 3-hydroxylated by P3H1 and presumably this enzyme is required for the 3-hydroxylation of the A3 site of both α-chains in bone but not in tendon. The increase in glycosylation of hydroxylysine in P3H1 null mice in bone was found to be due to an increased occupancy of normally glycosylated sites. Despite the severe disorganization of collagen fibrils in adult tissues, the D-period of the fibrils is unchanged. Tendon fibrils of newborn P3H1 null mice are well organized with only a slight increase in diameter. The absence of 3-hydroxyproline and/or the increased glycosylation of hydroxylysine in type I collagen disturbs the lateral growth of the fibrils.

Collagens are the most abundant proteins in vertebrate animals. Procollagen biosynthesis occurs in the rough endoplasmic reticulum of cells and involves a large number of posttranslational modifications (1). Two modifications of proline residues are known: 4(R)-hydroxyproline and 3(S)-hydroxyproline. The role of 4(R)-hydroxyproline is well established, it stabilizes the collagen triple helix and the enzyme prolyl 4-hydroxylase is well
characterized (2-3). Although it has been known for a long time that type I collagen also contains 3-hydroxyproline residues (4), the enzyme activity was only partially characterized (5-7) and the enzymes were only recently identified (8). The prolyl 3-hydroxylases (P3Hs) belong to a family of three proteins P3H1, P3H2 and P3H3, which share a common carboxy-terminal dioxygenase domain, which is similar to lysyl hydroxylases and the α-subunit of prolyl 4-hydroxylase, and a unique amino-terminal domain with four cysteine repeats (CXXXC) (8). This domain is homologous to cartilage associated protein (CRTAP) and SC65. The function of this domain is not established, but it also contains tetra-tricopeptide repeat domains that are known to be important in protein-protein interactions (9-10).

P3H1 forms a complex with CRTAP and cyclophilin B (CypB) and this complex was shown to 3-hydroxylate proline residue 986 of the α1-chain of type I collagen (8,11). The P3H1•CRTAP•CypB complex is also a potent molecular chaperone for procollagen biosynthesis (12). The absence of P3H1 in mice leads to abnormalities in fibrillar collagen-rich tissues such as tendon, skin and bone (13). Type I collagen from these mice lack the 3-hydroxyproline at position 986, and show increased glycosylation of hydroxylysine residues (13). Recessive Osteogenesis Imperfecta cases have been reported to be caused by mutations in CRTAP, P3H1, or CypB (14-18). The CRTAP null mice and the CypB null mice also show severe Osteogenesis Imperfecta and both show the absence of 3-hydroxylation at position 986 (11,19).

A comprehensive analysis of the location of 3-hydroxyproline sites in fibrillar collagens from human and bovine tissues showed the existence of additional sites (20). Four sites were identified in fibrillar collagens and termed the A1 site (position 986), the A2 site (position 944), the A3 site (position 707) and the A4 site (position 470). The α1-chain of type I collagen was shown to only contain a 3-hydroxyproline at the A1 site, which was 96 to 99 % occupied. The α2-chain of type I collagen was 80 % 3-hydroxylated at the A3 site. The α1-chain of type II collagen was 95 to 99 % 3-hydroxylated at the A1 site and 10 to 87 % at the A2 site (20). The spatial arrangement of these sites, which are staggered by a D-period, suggested a fundamental role for these modifications in the supramolecular assembly by forming hydrogen bonds between adjacent collagen triple helices (20-21). In rat tail tendon, a 3Hyp-rich motif was found at the C-terminal end of the triple helix (22).

Glycosylation of hydroxylysine residues is a normal posttranslational modification observed in many collagens (23). The degree of glycosylation varies among different collagens (24-25). For type I collagen, the degree of glycosylation also varies between different tissues.

Here we report that type I collagen extracted from wild type and P3H1 null mice tissues show differences to the published data on human and bovine collagens and we show that another prolyl 3-hydroxylase beside P3H1 also acts on type I collagen. We demonstrate that the D-period of collagen fibrils is not affected by the lack of 3-hydroxyproline or increasing amounts of hydroxylysine glycosylation. In addition, we see only small differences in the fibril diameter of newborn mice tail tendons compared to the severe disorganization found in adult P3H1 null mice, implicating the lateral growth of fibrils in the phenotype.

**EXPERIMENTAL PROCEDURES**

**Collagen extraction from tissues** - Tendons and skin were taken from adult mice and pepsinized in 0.5 M acetic acid. Bone was first decalcified in 0.2 M EDTA in 50 mM Tris/HCl, pH 7.8 in a Pelco (Ted Pella, Inc., Redding, CA) 3450 microwave oven equipped with a 3430 variable power module. The high temperature limit set to 40 ºC and power set to 99 watts. The EDTA solution was changed by guest on August 17, 2017 http://www.jbc.org/ Downloaded from

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added to a final concentration of 2.5 M to preferentially precipitate type I and III collagen. The solution was incubated at 4 ºC for several hours and then centrifuged at 20,000 x g for 40 minutes to collect the collagen precipitates. The pellets were resolubilized in 0.1 M acetic acid, analyzed on SDS-PAGE gels and lyophilized for further digestion and analysis. For the skin extracts the type III collagen was precipitated with 1.6 M NaCl to separate it from type I collagen.

**Collagen digestion and MS analysis** - SDS–PAGE bands were subjected to in-gel digestion with trypsin using a protocol similar to that described by (26). Digest conditions were 13 ng/µL Promega trypsin in 100 mM ammonium bicarbonate at 37 ºC for 18 h. Identification of tryptic peptides was performed on a Q-TOF Micro mass spectrometer (Waters, Billerca, MA) equipped with an electrospray ionization source. Data were collected with the MassLynx (version 4.1) data acquisition software (Waters, Billerca, MA) and processed using Mascot Distiller (Matrix Software, London, UK). High performance liquid chromatography was performed with a nanoAcquity (Waters, Billerca, MA) system using a 75 µm x 100 mm 3 µm Atlantis dC18 column as the analytical column and a 180 µm x 20 mm 5 µm Symmetry C18 column as the trapping column. Chromatography mobile phases consisted of solvents A (0.1 % formic acid and 99.9 % water (v/v)) and B (0.1 % formic acid and 99.9 % acetonitrile (v/v)). Peptide samples were loaded onto the trapping column and equilibrated for 2 min in 99 % solvent A, followed by a 120-min gradient to 60% solvent A at a constant flow rate of 1 µL/min. Analysis was performed in survey scan mode. Tryptic peptides were identified from MS/MS spectra by a Mascot search against the NCBI nr database (peptide tolerance 1.0 Da, MS/MS tolerance 1.0 Da).

Quantification of glycosylated hydroxylysine residues in type I collagen was accomplished by initially identifying glycosylated peptides in the hydrazide extracted samples. These peptides were then located in raw MS data from non-extracted trypsin digests of type I collagen from P3H1 null and wild type mouse samples from skin, bone, and tail tendon. Extracted ion chromatograms were prepared for the peptides containing the hydroxylated lysine and unhydroxylated lysine of interest and the corresponding glycosylated peptides. Extracted ion chromatograms were integrated using the MassLynx software package. The average area ratios of the modified peptides for each lysine site are shown in Table 1. Verification of the identity of the peptides was achieved by examination of the MS/MS spectra for each peak of interest.

**N-Terminal sequencing** – 1 M Tris was added to solubilized collagen extracts to adjust the pH to 8 and the solutions were incubated at 50 ºC for 1 hour. The temperature was reduced to 37 ºC and sequencing grade trypsin was added. Collagen samples were incubated at 37 ºC for 18 hours. Tryptic digests of collagen were loaded onto Oasis HLB solid phase extraction cartridges, washed with 0.1 N acetic acid and eluted with 40 %/60 % acetonitrile/0.1 % trifluoroacetic acid. After vacuum evaporation, the tryptic digests of collagen were reconstituted in 0.1 % trifluoroacetic acid. Peptides were isolated by reverse phase HPLC on a 1090 Liquid Chromatograph (Agilent, Santa Clara, CA) with a 2.1 mm x 50 mm 5 µm Zorbax 300 SBC-18 analytical column. Fractions were collected in 0.5 minute intervals, dried by rotary evaporation, and reconstituted in 0.1 % formic acid in water before analysis by LC-MS/MS using the method described above. Peptides in each fraction were identified by Mascot search using the conditions described above. HPLC conditions on the 1090 liquid chromatograph were adjusted so that a single chain contained peptides identified as containing a 3-Hyp modification from a single isoform of type I collagen. Two fractions, one containing the α1 A3 3-Hyp site and one containing the α2 A3 3-Hyp site were analyzed by N-terminal sequencing. Peptide fractions containing the 3-Hyp site were applied to Applied Biosystems Pro sorb cartridges (Applied Biosystems. Foster City, CA, US) and coated with 3 µL of diluted Biobrene solution (20 µg/µL) (Applied Biosystems. Foster City, CA, US). Edman sequencing was performed on an Applied Biosystems Procise 492 Protein Sequencer (Applied Biosystems. Foster City, CA, US). Sequencing data for these peptide fractions were processed using SequencePro 2.1 software (Applied Biosystems. Foster City, CA, US). 3-Hyp and 4-Hyp PTH Standards were prepared by manual Edman
chemistry using 3-Hyp and 4-Hyp dried standards purchased from Fluka Chemie (Buchs, CH) and Aldrich (St. Louis, MO, US) respectively. These were then added to PTH-Amino Acid Standard Solution (Applied Biosystems. Foster City, CA, US) in appropriate concentrations for an injection standard.

Amino acid analysis - Acid hydrolysis was performed in 6 x 50 mm Pyrex culture tubes placed in Pico Tag reaction vessels fitted with a sealable cap (Eldex Laboratories, Inc, Napa, CA, USA). Samples were placed in culture tubes, dried in a Speed Vac (GMI, Inc. Albertsville, MN, USA), and then placed into a reaction vessel which contained 250 ml of 6 N HCl (Pierce, Rockford, IL, USA) containing 2% phenol (Sigma Aldrich, Milwaukee, WI, USA). The vessel was then purged with argon gas and vacuumed using the automated evacuation workstation Eldex Hydrolysis/Derivatization workstation (Eldex Laboratories, Inc., Napa, CA). Closing the valve on the Pico Tag cap maintained the vacuum during hydrolysis at 110 °C for 24 h. The hydrolyzed samples were then dried in a Savant Speed Vac. The dried samples were dissolved in 100 ml of 0.02 N HCl containing an internal standard (100 µM norvaline; Sigma). Appropriate further dilutions were made using the same dilution solvent for concentrated samples. Analysis was performed by ion exchange chromatography with postcolumn ninhydrin derivatization and visible detection (440 nm/570 nm) with a Hitachi L-8800A amino acid analyzer (Hitachi High Technologies America, Inc., San Jose, CA, USA) running EZChrom Elite software (Scientific Software, Inc., Pleasanton, CA, USA).

Mouse tissue preparation for RNA extraction - Bone, skin and tail tendon were isolated from wild type and P3H1 null one-month-old mice and stored at -20 °C in RNAlater stabilization solution (Qiagen). Mouse hind limb bones, femur and tibia, were cut from both ends and extensively washed with PBS to remove bone marrow.

Real time quantitative PCR - Total mouse RNA was extracted from bone, skin and tail tendon of wild type and P3H1 null mice using RNeasy Fibrous Tissue Mini Kit (Qiagen). RNA quality was validated by 260/280 absorption ratio and concentration was estimated. Each representative RNA was reverse transcribed using Superscript Reverse Transcriptase III (Invitrogen) according to the manufacturer’s instructions with random hexamers (Invitrogen). Resulting cDNA was used as the template in real-time quantitative PCR experiments with the P3H1, 2, or 3 or GAPDH gene specific oligonucleotides and the iQ SYBR Green Supermix according to the manufacturer’s instructions (Bio-Rad). The following primer pairs were used for each gene amplification:

P3H1: CCCCCAGCCTACACGTTCG
      TGCCCCCTGTTGACACCTTC;
P3H2: ATGCTAAACCGTGACTGCC
      CGCTCCAGTTCTCGGTAAAG;
P3H3: AGAGGCTCATTACCCATTTG
      GTTGGGTTTGCACAAAG;
GAPDH: CCACCAGAAGACTGTTGAT
       TTCAGCTTGGGATGACCTT

Quantitative PCR data were measured and analyzed using the iQ5 Multicolor Real-Time PCR Detection System and software version 2.0 (Bio-Rad). Each PCR reaction was 25 µL. Because SYBR Green binding is not sequence specific, careful validation of the primer pairs was undertaken to ensure that only the target gene sequence was amplified. To verify the specificity of each amplicon, a single gel band of the expected size was amplified for each primer pair and melting curve analysis confirmed the presence of a single PCR product. Fold differences in target genes were normalized to GAPDH expression in each tissue where indicated (ΔΔC_T method). Data analysis was performed using the iQ5 Optical System Software version 2.0.

Analysis of glycosylation - To verify the presence of sugar modifications, the glycosylated peptides were isolated using hydrazide chemistry (27). Galactose oxidase was bound to Sepharose 4B under basic conditions. Type I collagen from wild type and P3H1 null mouse tendon and bone was digested with trypsin at 37 °C for 18 hours. The lyophilized digests were reconstituted in 0.1 M sodium phosphate buffer, pH 7.2, containing 0.15 M NaCl. 115 units of catalase (from bovine liver) and peroxidase (from horseradish) were added in a Bio-Rad Poly-Prep Column and allowed to incubate overnight at 37 °C. The flow-through of the columns containing oxidized and unoxidized tryptic peptides of collagen was collected and adjusted to pH 5.5 with 1 M HCl. The hydrazide resin was
washed with 0.1 M sodium acetate, pH 5.5, containing 0.15 M NaCl and the flow-through was added to the resin in a Poly-Prep column. The solution was incubated overnight at 37 °C. The columns were then washed with 0.1 M sodium acetate, pH 5.5 containing 0.15 M NaCl, followed by 0.15 M NaCl in water, then methanol, and finally water. The washed hydrazide gel bound to the oxidized galactose containing peptides was incubated with 0.1% formic acid for 1 h at 37 °C. The flow-through containing the oxidized peptides was collected and reduced with 1 mM sodium borohydride for 1 h at room temperature under basic conditions. The solution was then lyophilized and the residue was analyzed by LC-MS using the method described above. Peptides containing glycosylation sites observed in the tryptic digests were also present in the hydrazide extracted samples.

**P3H1 null mice** - P3H1 null mice were purchased from Deltagen (San Mateo, CA). Directed knockouts were created in which exons 1-3 (nucleotides 15-817) of the mouse P3H1 or leprecan 1 gene (NCBI Reference sequence number: NM_019782.2) were deleted. Then a LacZ-Neo cassette was inserted into the area of the target gene that was deleted. P3H1 or leprecan gene inactivation was verified in mice by RNA preparation from tissues of null mice, followed by reverse transcription and PCR with primer sets spanning the length of the target gene. No P3H1 transcripts or gene expression was detected, as compared with normal levels of the target gene expressed in wild type animals. Mice were bred multiple generations into a C57B6 background prior to analysis to verify phenotype effects and data reported is from 8th to 10th generation (13).

**Electron microscopy analysis of P5 tendon** - Freshly obtained tendons were fixed in cacodylate buffered 1.5% glutaraldehyde/1.5%paraformaldehyde containing 0.05% tannic acid (w/v), then rinsed, exposed to 1% osmium tetroxide, then dehydrated in a grade series of ethanol to 100%. Fixed tissues were rinsed in propylene oxide and infiltrated and embedded in Spurr’s epoxy. 80-nm ultrathin sections were mounted on formvar coated single hole slot grids and stained in ethanolic uranyl acetate followed by Reynold’s lead citrate. Stained sections were examined using a FEI Tecnai G2 electron microscope operated at 120kV and photographed using either a FEI Eagle 2K camera or an AMT 2K camera. Magnifications were calibrated using a grating replica (Ted Pella catalog # 603).

**Small angle X-ray diffraction** - The small angle X-ray diffraction (SAXD) measurements were carried out in the BL40B2 beamline of SPring-8 (Hyogo, Japan). Collagen fibers were dissected from mouse tail and immersed in phosphate buffered saline. A picked collagen thread (diameter 0.3~0.8 mm) was clamped on a small metal holder and quickly set to the goniometer on the SAXD equipment with the collagen fiber axis vertical. The 2D SAXD patterns were recorded with RAXIS IV++ (Rigaku Co., Japan). The camera length and X-ray wavelength $\lambda$ were set to 1151 mm and 0.1 nm, respectively. The scattering intensity was accumulated for 30 to 120 sec and all collagen samples were wet well during measurement. 1D diffraction profiles were calculated by circular integrating the 2D data over 20 degree fan-shaped sectors including the low angle meridional diffraction pattern as the function of scattering vector $s$, where $s = 2\sin \theta / \lambda$ ($\theta$ = half of the scattering angle). Each higher order diffraction peak (from 2nd to 36th) of the D-stagger of collagen samples was fitted with Pseudo Voigt peak function to estimate full width of half maximum (FWHM).

**RESULTS**

The A1 site (Pro986) of the $\alpha1$-chain of type I collagen is 3-hydroxylated almost completely in tendon and bone and to a high degree in skin of wild type mice. Type I collagen extracted from the same tissues of the P3H1 null mice show a complete absence of 3-hydroxylation at this site (Fig. 1). A tissue specific 3-hydroxylation was found for the A3 site (Pro707) in the $\alpha1$-chain. In tendon and bone of wild type mice this site was about 50 % 3-hydroxylated, whereas in skin the 3-hydroxylated peptide was not detectable. The only difference found in tissues extracted from P3H1 null mice at this site was the lesser extent of 3-hydroxylation in bone (Fig. 2).

The $\alpha2$-chain of type I collagen lacked 3-hydroxylation of the A1 site in all tissues, because the corresponding sequence is Gly-Pro-Ala instead of the required Gly-Pro-Hyp. In tendon and bone of
wild type mice a substantial amount of 3-hydroxylation was observed at the A3 site (Pro707). Skin had very little 3-hydroxylation at this site. In tendon extracted from P3H1 null mice, no significant reduction in 3-hydroxylation was observed. However, in bone a significant decrease in 3-hydroxylation at this site was found (Fig. 3) in knockouts. Schematic representation of 3-hydroxylation sites in both chains of type I collagen is shown in Figure 4.

To investigate the tissue specific expression levels of the P3H1, P3H2 and P3H3 enzymes in bone, skin and tendon of mutant and wild type mice, quantitative real-time PCRs were performed. Fold expression relative to GAPDH of the three enzymes in wild type tissues is shown in Figure 5A. The most highly expressed enzyme type was P3H2. This finding is consistent with the previously published data for bone and skin (28). P3H1 was upregulated in bone compared to skin and tendon.

Fold expression of the P3H2 and P3H3 in the P3H1 null tissues relative to the wild type controls is represented in Figure 5B. A dramatic 40 fold reduction in the P3H2 expression level was observed in bone of the P3H1 null. In mutant skin, the P3H2 expression level was also reduced to about 1/5 of that of the control. However, in tendon of the P3H1 null mice expression of the P3H2 was 100% preserved as compared to wild types. Expression of the P3H3 was slightly upregulated in P3H1 null bone and tendon but downregulated in skin.

Several glycosylation sites were identified in mouse type I collagen. It was shown previously that type I collagen in P3H1 null mice has increased levels of hydroxylysine and glycosylated hydroxylysines (13). However, it was not clear if this increase is due to additional hydroxylation and glycosylation sites or due to an increased occupancy of existing modifications. Figure 6 shows a comparison of the posttranslational modification of Lys174 of the α1-chain of type I collagen of wild type and P3H1 null mice. There is a clear increase in the amount of galactosyl-hydroxylysine and the existence of glucosyl-galactosyl-hydroxylysine at this position in the P3H1 null mouse. The increase in these glycosylations is confirmed by the analysis of tryptic peptides of the α1-chain of tendon type I collagen using hydrazide extraction to enrich for glycopeptides. Figure 7 shows the relative occurrence of the galactosyl-hydroxylysine and the glucosyl-galactosyl-hydroxylysine at this position in wild types and P3H1 nulls. Table 1 lists the occupancies of hydroxylysine modifications for a number of locations and tissues.

Tendons from adult wild type and P3H1 null mice were analyzed by small angle x-ray scattering. Both X-ray diffraction patterns of wild type and P3H1 null mice tendon show the set of meridional reflections parallel to the fiber axis. These reflections provide information about the axial order of the collagen fibrils. From this experiment, the D-period of the fibrils was determined to be 66.16 nm and 66.11 nm, for wild type and P3H1 null respectively. Since there is no significant difference between them, it is suggested that the basic molecular packing of collagen molecules along the fibril axis is the same. Figure 8 shows the \( s^2 \) dependence of FWHMs for diffraction peaks of D-period of two mouse types. The line broadening of the diffraction peaks is due to not only the instrumental broadening but also to the small size of diffracting coherent domains, and to the lattice distortions generated by imperfections. Since the instrumental broadening does not change between different samples measured under the same condition, the larger FWHMs indicate that the coherent domain size is smaller and/or the lattice distortions are larger. When the line width is plotted against \( s^2 \), the value obtained by extrapolation to \( s = 0 \) can be interpreted in terms of the crystal size effect, whereas the slope of such a plot can be related to the extent of lattice distortion (29). We easily perceive that the two extrapolated values from line width are almost the same (slightly larger for P3H1 null), but the slope of P3H1 null mouse data is larger than that of wild type. The larger slope of P3H1 null mouse is caused by the larger D-stagger distortions. In other words, collagen fibrils of null mouse tendon formed more disordered packing compared to wild type tendon.

Tail tendons from newborn wild type and P3H1 null mice (P5) were analyzed by electron microscopy. The diameter of these fibrils is rather uniform and the P3H1 null mouse tendons have a slightly larger diameter (Fig. 9). There is no difference in the repeat period of the fibrils when measured from electron microscopy images (data not shown). The tendon fibrils of adult P3H1 null mice
show a complete disorganization with large fibrils splitting into smaller structures and the merging of smaller fibrils into larger ones.

**DISCUSSION**

The proline 3-hydroxylation patterns of the mouse α1- and α2-chains of type I collagen extracted from tendon, skin and bone were analyzed and are summarized in Figure 4. The A1 site (Pro986) of the α1-chain is fully 3-hydroxylated. The lack of 3-hydroxylation of proline residue 986 in type I collagen extracted from P3H1 null tissues clearly demonstrates that P3H1 is solely responsible for the modification of this site. The A3 site exhibited a tissue specific degree of 3-hydroxylation and variable disappearance in the absence of the P3H1. This places one of the other two enzymes, P3H2 or P3H3, in the position to fulfill the function of this site 3-hydroxylation. In a recent study (30) based on cell culture experiments, the suppression of P3H2 by siRNA was associated with the sharp decrease in 3-hydroxylation of the A3 site in the α-2 chain of type I collagen. Our quantitative real-time PCR results, which summarize tissue specific expression patterns of the prolyl 3-hydroxylase family members, lead to the same conclusion. As shown in Figure 5B, the expression level of P3H2 drops dramatically in the P3H1 null bone where 3Hyp at the A3 site is drastically reduced. At the same time, P3H2 is fully expressed in the P3H1 null tendon where the 3-hydroxylation level of the A3 site remains very similar to that of the wild type. In the P3H1 null bone, the expression of P3H3 is slightly upregulated perhaps in attempt to compensate for the loss of the P3H2. In P3H1 null skin both P3H2 and P3H3 expression levels are somewhat reduced. This, however, does not have an effect for modification of the A3 site since it already lacks 3-hydroxylation in the normal skin (Fig. 4). Thus, P3H2 plays an important role in modification of the A3 site in type I collagen.

The increase in glycosylation of hydroxylysine, observed quantitatively by base hydrolysis and amino acid analysis in P3H1 null mice (13), was analyzed at Lys174 of the α1-chain and Lys174 and Lys219 of the α2-chain in various tissues. In almost all cases a significant increase of glucosyl-galactosyl hydroxylysine was found. In many cases the level of galactosyl hydroxylysine is also increased. This analysis is consistent with the quantitation previously published. The major increase in glycosylation occurs in bone as compared to other tissues and it seems that this increase occurs at normally modified hydroxylysines.

Full occupancy of the A1 site in the α1 chain of type I collagen along with the Osteogenesis Imperfecta phenotypes in the absence of such modification is suggestive of its importance for collagen fibrillogenesis. The concept of fibril molecular packing in which the subunits are molecular dimers in register and staggered axially by D-periods was proposed (31). Recently this concept was proposed to be strengthened by interactions involving 3-Hyp (20), because of the D-periodic spacing of the 3-Hyp sites in collagen. Additionally, limited evidence was found for the role of 3-Hyp at 986 position in dimerization of collagen triple helices (21). According to this concept, one would expect at least somewhat disturbed D-period in the collagen fibrils where A1 site is not 3-hydroxylated. However, our data clearly demonstrate no difference in the D-spacing between wild type and P3H1 null collagen fibrils both by small angle X-ray diffraction and by electron microscopy. Moreover, EM images of P5 tendon demonstrate that collagen fibrils of the mutant mice lack any disorganization. In fact, these fibrils are perfectly normal despite a slight increase in the diameter. Both findings disagree with the previously described concept.

Tendon fibrils apparently form normally during embryonic development even in the absence of the 3Hyp at the A1 site. Their diameter is likely increased due to bulkier collagen chains carrying more sugars. It is during postnatal growth when the disorganization is observed. Adult tendon appears to be composed of extremely disorganized fibrils that are branching and fusing as seen by EM. During the postnatal maturation process, the lateral fusion of smaller fibrils (40 to 50 nm) into larger ones (< 200 nm) occurs. The conclusion is that the P3H1 null immature tendon fibrils fail to fuse properly.

The bone and tendon embryonic development shows distinct differences in P3H1 null mice. While tendon appears almost normal at birth, the degree of bone mineralization is already severely affected at this stage. Simultaneously, 3-
hydroxylation of the A3 site of type I collagen in
bone is strongly affected in the P3H1 null mice.
Thus, it is possible that the 3Hyp located at the
A3 site could play a role during initial stages of
fibrillogenesis. On the other hand type I
collagen is least posttranslationally modified in
bone compared to other tissues. The low
abundance of sugars might be a prerequisite for
the tight packing and mineralization of type I
collagen in bone. It is possible that the excess
glycosylation in bone results in more severe
disturbances than the lack of 3-hydroxyproline.
REFERENCES


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FOOTNOTES

Abbreviations: P3H1, prolyl 3-hydroxylase 1

FIGURE LEGENDS

FIGURE 1. Analysis of the A1 site (Pro986) of the α1-chain of type I collagen. The α1-chain of type I collagen extracted from tendon (A), skin (B) and bone (C) was analyzed by mass spectrometry. The occurrence of the tryptic peptides DGLNGLOGPIGPOGPR (mass = 1548 Da) and DGLNGLOGPIGZOGPR (mass = 1564 Da) are shown from wild type and P3H1 null mice. O is 4(R)Hyp and Z is 3(S)Hyp. Figure 1A was previously published in (13).

FIGURE 2. Analysis of the A3 site (Pro707) of the α1-chain of type I collagen. The α1-chain of type I collagen extracted from tendon (A), skin (B) and bone (C) was analyzed by mass spectrometry. The occurrence of the tryptic peptides VGPOGPSGNAGPOGPOGPVGK and VGZOGPSGNAGPOGPOGPVGK are shown from wild type and P3H1 null mice. MS/MS spectra of these peptides of the α1-chain of type I collagen are shown in D. The Y19+2H ion shows a mass difference of 8 Dalton, corresponding to the hydroxylation of Pro707 (A3 site). Sequencing identifies this hydroxylation as 3-hydroxyproline. N-terminal sequencing cycles for the A3 site are shown in (E). 3-Hyp residues are highlighted to confirm the presence in the cycle. BL is baseline, ST is standards and the cycle numbers are indicated. O is 4(R)Hyp and Z is 3(S)Hyp.

FIGURE 3. Analysis of the A3 site (Pro707) of the α2-chain of type I collagen. The α2-chain of type I collagen extracted from tendon (A), skin (B) and bone (C) was analyzed by mass spectrometry. The occurrence of the tryptic peptides TGPOGPSGIAGPOGPOGAAGK and TGZOGPSGIAGPOGPOGAAGK are shown from wild type and P3H1 null mice. MS/MS spectra of the TGZOGPSGIAGPOGPOGAAGK and TGPOGPSGIAGPOGPOGAAGK peptides of the α2-chain of type I collagen are shown in (D). The Y19+2H ion shows a mass difference of 8 Dalton, corresponding to the hydroxylation of Pro707 (A3 site). N-terminal sequencing identifies this hydroxylation as 3-hydroxyproline (E). 3-Hyp residues are highlighted to confirm the presence in the cycle. BL is baseline, ST is standards and the cycle numbers are indicated. O is 4(R)Hyp and Z is 3(S)Hyp.

FIGURE 4. Schematic representation of the A1 and A3 3-hydroxylation ratios of type I collagen from skin, tendon and bone of wild type and P3H1 null mice. White circles indicate no 3-hydroxylation and black circle indicates full 3-hydroxylation. Partial 3-hydroxylation is indicated by blackened quarters.

FIGURE 5. Graphical representation of real-time quantitative PCR analysis of adult mouse bone, skin and tendon. Data are represented as the fold change differences. Graph A shows relative target gene expression to GAPDH signal. Graph B shows fold change expression of P3H2 and P3H3 genes in P3H1 null relative to wild type control tissues (B). All experiments were performed in triplicate and mean values are represented on each graph. P3H1 gene expression is represented by the black boxes, P3H2 is in white and P3H3 is in gray.

FIGURE 6. MS spectra of trypsin digested mouse tendon type I collagen. The peptide GARGNDGAAGGAAGGPTGPTGPAGAVGAK(174)GEAPQGAR is shown for P3H1 null and wild type extracts from tendon. The relative abundance of glycosylated peptide in collagen extracted from P3H1 null mice is much higher (A). MS/MS data for the glycosylated peptide GARGNDGAGGAAGGPTGPTGPAGAVGAK(174)GEAPQGAR (B). Spectra labeled 1, 2, and 3 show the partially fragmented 3+ charge state precursor ions for the glycosylgalactosyl, galactosyl, and non-sugar containing peptides of K174 respectively. Spectra labeled 1a, 2a, and 2c show an...
expanded view of the previous peptides at the low mass region containing the same identifying fragments for all three peptides.

**FIGURE 7.** MS spectra of trypsin digested mouse tendon type I collagen purified by hydrazide extraction. Only the glycosylated variants of the peptide GARGNDGAVGAAGPPGPTGPTGPPGFPFGAVGAK(174)GEAGPQGAR are observed by this method. The relative abundance of the two forms of glycosylation from one extract can be compared, but not the ratio between two extracts.

**FIGURE 8.** \( s^2 \) dependence of FWHMs for diffraction peaks of D-stagger. The FWHMs of low angle meridional X-ray diffraction peaks were estimated by fitting with Pseudo Voigt peak function. The black closed circles (●) represent the FWHM from diffraction of wild type mouse tendon while the red open circles (○) represent that of P3H1 null mouse tendon.

**FIGURE 9.** Electron micrographs of tendon fibrils of wild type and P3H1 null mice. Tendon fibrils of wild type (A) and P3H1 null (B) mice at P5 in cross- and longitudinal sections and the distribution of measured diameters from the cross sections underneath. The length of the bars in the micrographs corresponds to 100 nm. Tendon fibrils of adult wild type (C) and P3H1 null mice (D) in cross- and longitudinal sections. The length of the bars corresponds to 500 nm. The diameter distribution from cross sections is given in E for P5 and in F for adult mice. Figures C,D are similar to fields published previously and F is taken from Figure 2E in (13).
Table 1

Relative abundance of lysine residues and posttranslationally modified lysine residues at position 174 of the α1-chain and α2-chain and position 219 of the α2-chain of type I collagen extracted from wild type and P3H1 null mice.

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<th>Mouse</th>
<th>Type</th>
<th>Site</th>
<th>R-K</th>
<th>R-K(OH)</th>
<th>R-K(OH)-R'</th>
<th>R-K(Gal)-R'</th>
<th>R-K(GluGal)-R'</th>
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<td>0.46</td>
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The numbers represent the relative abundance of each species at each position and tissue when wild type and P3H1 null extracted collagens are compared. R indicates preceding amino-terminal sequence and R’ indicates continuing C-terminal sequence.
Figure 1
Figure 2

(A) Graph showing posttranslational modifications in mouse type I collagen.
(B) Graph showing posttranslational modifications in mouse type I collagen.
(C) Graph showing posttranslational modifications in mouse type I collagen.

(D) Table detailing specific modifications and their respective changes.

(E) Graph showing absorbance over time (min) for different conditions.

Y2, Y6, Y9, Y12, Y14, Y16, Y18, M+2H, 3-Hyp, 3-Hyp.
Figure 4

Wild-type

Tendon

Skin

Bone

A3 A1

Null

A3 A1
Figure 5

A

Fold expression relative to GAPDH

Tendon  Skin  Bone

P3H1  P3H2  P3H3

B

Fold expression in P3H1 null vs wild type tissues

Tendon  Skin  Bone

Fold expression in P3H1 null vs wild type tissues
Figure 6

A

P3H1 Null

No glycosylation

+Gal

+GluGal

WT

No glycosylation

+Gal

B

1

2

3

1a

2a

3a
Figure 7
Figure 9
Posttranslational Modifications in Type I Collagen from Different Tissues extracted from wild type and Prolyl 3-hydroxylase 1 Null Mice
Elena Pokidysheva, Keith D. Zientek, Yoshihiro Ishikawa, Kazunori Mizuno, Janice A. Vranka, Nathan T. Montgomery, Douglas R.. Keene, Tatsuya Kawaguchi, Kenji Okuyama and Hans Peter Bachinger

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