Variable Bone Fragility Associated With an Amish COL1A2 Variant and a Knock-in Mouse Model

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ABSTRACT

Osteogenesis imperfecta (OI) is a heritable form of bone fragility typically associated with a dominant COL1A1 or COL1A2 mutation. Variable phenotype for OI patients with identical collagen mutations is well established, but phenotype variability is described using the qualitative Sillence classification. Patterning a new OI mouse model on a specific collagen mutation therefore has been hindered by the absence of an appropriate kindred with extensive quantitative phenotype data. We benefited from the large sibships of the Old Order Amish (OOA) to define a wide range of OI phenotypes in 64 individuals with the identical COL1A2 mutation. Stratification of carrier spine (L1–4) areal bone mineral density (aBMD) Z-scores demonstrated that 73% had moderate to severe disease (less than −2), 23% had mild disease (−1 to −2), and 4% were in the unaffected range (greater than −1). A line of knock-in mice was patterned on the OOA mutation. Bone phenotype was evaluated in four F1 lines of knock-in mice that each shared approximately 50% of their genetic background. Consistent with the human pedigree, these mice had reduced body mass, aBMD, and bone strength. Whole-bone fracture susceptibility was influenced by individual genomic factors that were reflected in size, shape, and possibly bone metabolic regulation. The results indicate that the G610C OI (Amish) knock-in mouse is a novel translational model to identify modifying genes that influence phenotype and for testing potential therapies for OI. © 2010 American Society for Bone and Mineral Research.

KEY WORDS: OSTEGENESIS IMPERFECTA; BONE; COLLAGEN; KNOCK-IN; RODENT

Introduction

Osteogenesis imperfecta (OI) is a heritable form of bone fragility with an overall spectrum of disease severity that ranges from a perinatal lethal form to mild disease that can remain clinically silent.(1–3) Multiple fractures is the principal clinical presentation that brings OI patients to medical attention. Additional phenotype traits can include bone deformity (e.g., long bone bowing), short stature, blue sclerae, dentinogenesis imperfecta (DI), and hearing loss. Sillence originally classified OI into types I to IV on the basis of clinical presentation, radiographic features, and mode of inheritance.(4,5) OI types I to IV typically are associated with heterozygous mutations in the genes (COL1A1 and COL1A2) that encode the α chains of type I procollagen molecules. Currently, more than 800 COL1A2 mutations have been identified in OI patients, with many examples of recurrent mutations in unrelated individuals.(3) However, there are no large pedigrees with quantitative phenotype data for OI patients with the identical mutation.

The predominant type I collagen isotype in the extracellular matrix (ECM) is a heterotrimeric molecule comprised of two α1(I) chains and one α2(I) chain. A homotrimeric isotype of type I collagen, comprised of three α1(I) chains, is also found in tissues in small amounts in the nonpathologic state. The homotrimeric isotype is associated with autosomal recessive connective tissue disorders in humans when no α2(I) chains or only nonfunctional α2(I) chains are synthesized.(6,7) The amino acid sequences of both α1(I) and α2(I) chains are homologous and have a characteristic [Gly-X-Y]338 repeat within their respective triple-helical domains. The small side chains of glycine residues
positioned at every third position of the repeat are sterically required for helix formation, and mutations in these glycine residues account for 80% of the known triple-helical mutations that cause moderately severe to lethal forms of OI.

Over the past two decades, several laboratories have developed murine OI models based on mutations in the proα1(I) chain of type I collagen patterned after those described in human OI to extend our understanding of OI pathobiology. All the existing mouse models have a moderate-to-severe bone phenotype with impaired viability. The two early models of note are the Col1a1 null allele Mov-13 mice (8–10) and the “protein suicide” or collagen minigene model (11–13). The homozygous Mov-13 mouse produces no type I collagen, and this is lethal. Heterozygous Mov-13 mice deposit reduced amounts of normal type I collagen in the ECM that is associated with an osteopenic phenotype and abnormal skeletal biomechanics. The minigene model construct expresses a version of the human COL1A1 gene that is missing the central 41 exons that encode a significant portion of the triple-helical domain. The shortened human proα1(I) chains associate with normal murine proα1(I) and proα2(I) chains, depleting the amount of normal type I collagen and altering ECM organization. Most recently, the Cre/lox recombination system was used to develop a Col1a1 G349C substitution to reproduce a human OI type IV (moderately severe) phenotype termed BrtIV(14–17). The BrtIV mouse represented a major advance in OI mouse models because it is the only Col1a1 model with a triple-helical glycine mutation. The OI phenotype pattern of proα2(I) glycine substitutions appears to be different from proα1(I) substitution. Generally, proα2(I) mutations are less severe. They also may represent an easier target for treatment because only 50% of collagen molecules are affected in COL1A2 heterozygotes versus 75% in COL1A1 heterozygotes. However, the only reported mouse model for OI with a Col1a2 mutation is osteogenesis imperfecta-murine (oim). Homozygous oim mice exhibit skeletal disease with clinical and biochemical features of Sillence type III (severe progressive) OI. The oim mutation and its biologic consequences (18–21) are strikingly similar to those found in a German child with type III OI (6,22,23). In both instances, a frame-shift mutation in the region of the COL1A2 gene that encodes the terminal portion of the proα2(I) C-propeptide predicts the synthesis of nonfunctional proα2(I) chains. Despite being the only naturally occurring OI mouse model, the oim mouse has a rarely observed type of mutation and an atypical biochemical phenotype because only proα1(I) homotrimeric molecules are formed.

The absence of a COL1A2 murine OI model with autosomal dominant inheritance, mild to moderate disease, and phenotype variation represents a major gap in our tools to understand OI pathobiology and to develop effective treatment. Defining the molecular basis of phenotype variation also has been limited by the absence of large numbers of OI patients carrying the identical causative mutation with quantitative phenotype data to use as a prototype for a new OI mouse model. Here we report the clinical and biochemical phenotype of a COL1A2 gene variant found among a Lancaster County, Pennsylvania, Old Order Amish (OOA) kindred. This COL1A2 variant is a G-to-T transversion at nucleotide 2098 that alters the gyl-610 codon (GGT) to a cysteine (TGT) codon (24). We also report the creation of the first glycine-substitution knock-in Col1a2 mouse that replicates the gene and protein phenotypes observed in the OOA kindred. The knock-in G610C OI (Amish) mouse represents a unique translational model for evaluating the molecular basis of phenotype variability and to test potential therapeutic strategies for OI.

Materials and Methods

Ethical considerations

All human subject research was reviewed and approved by the Institutional Review Board of the University of Maryland Baltimore (UMB). Informed consent was obtained for all study participants. All animal procedures were reviewed and approved by the UMB Institutional Animal Care and Use Committee.

Recruitment of family members and phenotype assessment

The study had three stages of informed consent: (1) mutation screening, (2) clinical assessment, and (3) skin-biopsy harvest. The screening enrollment age requirement of 6 years or older was waived if one parent was mutation-positive by sequence analysis. After obtaining informed consent (or assent from the children), buccal cells were harvested as a source of genomic DNA (gDNA) for genotyping. All gDNA- and cDNA-derived sequence data (see below) were analyzed, using Sequencher software (GeneCodes, Ann Arbor, MI, USA).

All volunteers verified by gDNA sequence analysis to carry a single copy of the T allele (OI group) and their control (G allele) nuclear family members (age ≥ 6 years) were invited to participate in the clinical assessment and skin-biopsy phases. The phenotype assessment consisted of a medical and family history, physical examination, blood draws for serum procollagen type I C-terminal propeptide (PICP) and CrossLaps measurements (markers of bone formation and resorption), and measurement of areal bone mineral density (aBMD). Lumbar spine (L1–4) and proximal femur (total and subregions) aBMD was measured by dual-energy X-ray absorptiometry (DXA) on a QDR 4500W instrument (Hologic, Bedford, MA, USA), and the results reviewed by a single clinician (EAS) and another author (DJM). The in vivo coefficient of variation for replicate aBMD measurements was 0.8% for the lumbar spine, 0.9% for total hip, and 1.5% for the femoral neck. The Hologic Pediatric Reference Curve Option (hip: 5 to 20 years; spine: 3 to 20 years) and Caucasian reference (age > 20 years) were used to obtain age-specific Z-scores. Since OI is frequently associated with short stature and bone mineral content (BMC) and aBMD are influenced by bone size, volumetric bone mass [bone mineral apparent density (BMAD)] also was estimated to minimize the influence of small bone size (25,26). A subset of the phenotype assessment volunteers also provided 3 mm dermal punch biopsies from the posterior upper arm from which fibroblasts were derived and stored at UMB and/or submitted to the Coriell Institute (OI families 1997, 1998, 2187, and 2229).
Generation of knock-in mice

Knock-in mice were created using embryonic stem cells and Cre/lox P technology. A 13 kb genomic clone containing the relevant segment of the murine Col1a2 gene was isolated from a λ phage library constructed with gDNA of the 129Sv/Ev Taconic mouse strain. PCR site-directed mutagenesis was used to change the targeted codon from GGT to TGT. The Col1a2 G610C-positive embryonic stem cells were injected into blastocysts of C57BL/6J (B6) mice. Founder mice that retain the neo targeting vector are termed neo- or G610C Neo mice. Progeny obtained by breeding a founder male with a female that expressed Cre recombinase (Jackson Laboratory, Bar Harbor, ME, USA, Stock Number 003724) are termed neo- or G610C OI mice. The G610C OI mouse line is available to the research community through the Jackson Laboratory Mouse Repository (Jax Stock Number: 007248).

Four F1 strains of G610C OI mice were used to determine the effect of genetic background on phenotype in 2-month-old male mice. Incipient congenic G610C OI B6 (–98% B6 genetic background) male breeders were generated by six generations of backcrosses to Jackson Laboratory B6 mice (Stock Number 000664). Experimental heterozygous B6 male breeders were crossed with A/J (Stock Number 000646), BALB/cByJ (Stock Number 001026), C3H/HeJ (Stock Number 000659), and FVB/NJ (Stock Number 001806) females purchased from the Jackson Laboratory. Progeny of these crosses are designated, respectively, as A.B6, Cby.B6, C3.B6, and FVB.B6. Experimental mice were housed at UMB in a single specific-pathogen-free room and were exposed to identical environmental conditions consisting of a 12 hour light/dark cycle, an ambient temperature of 23°C, and ad libitum access to water and laboratory mouse chow. Genotype was assigned using a PCR assay that can discriminate the three possible G610C OI mouse genotypes. The forward primer (TCC CTG CTT GCC CTA GTC CCA AAG ATC CTT) and the reverse primer (AAG GTA TAG ATC AGA CAG CTG GCA CAT CCA) will generate a 165 bp (wild type) or a 337 and a 165 bp (heterozygous) or a 337 bp ( homozygous) PCR product using G610C OI mouse gDNA. All animals were euthanized by CO2 asphyxiation.

Human and mouse collagen analysis

Pepsin-soluble collagen was purified from mouse tail tendons andhuman fibroblast culture medium for analysis by SDS-PAGE and differential scanning calorimetry (DSC). Purified collagen samples were labeled by fluorescent monoreactive Cy5 NHS ester (GE Healthcare Bio-Sciences Corp., Piscataway, NJ, USA) or double-labeled by Cy5 and fluorescent BODIPY-L-Cys (Molecular Probes, Invitrogen, Inc., Carlsbad, CA, USA).127 Labeled proteins were size-separated by SDS/PAGE on precast 3% to 8% gradient Tris-acetate minigels (Invitrogen). Following electrophoresis, the gels were briefly rinsed in distilled water and scanned on a Fuji FLA5000 fluorescence scanner (Fuji Medical Systems, Stamford, CT, USA) using a 473 nm excitation laser for BODIPY-L-Cys and a 635 nm laser for Cy5. For DSC, collagen was redissolved (1 to 2 mg/mL final collagen concentration) in 2 mM HCl (pH 2.7) or in 0.2 M sodium phosphate/0.5 M glycerol (pH 7.5) and then dialyzed extensively against the same buffer to remove excess salt. DSC thermograms128 of approximately 0.1 mg/mL of pepsin-treated collagen in 2 mM HCl (pH 2.7) and in the phosphate/glycerol buffer (pH 7.5) were recorded in a Nano II differential scanning calorimeter (Calorimetry Sciences Corp., Lindon, UT, USA) at several different scanning rates. The apparent melting temperature Tm was defined at the maximum of the melting peak after baseline subtraction.

Total RNA was extracted from fibroblast cultures, and cDNA was synthesized using RACE1 [5'-GAT GGA TCC TGC AGA AGC (T)17-3']. An aliquot of cDNA and forward exon primer and downstream reverse exon primer (selected to cross intron-exon) boundaries were used for second-strand synthesis and PCR amplification. PCR products of the expected size were confirmed by agarose gel size separation prior to purification for direct nucleotide sequencing.

Phenotype assessment of F1 G610C OI Mice

aBMD measurements

The aBMDs of right femurs were measured by DXA (GE-Lunar densitometer, PIxImus; software version; 2.10, GE Medical Systems, WI, USA). Femur measurements were obtained by placing an excised femur on a Delrin block. The percent coefficient of variation (CV) was 1.3% for BMC and 1.4% for BMD.

Confocal Raman microspectroscopy

Segments of cortical bone from the femur diaphysis were embedded in Cryo-gel (Instrumedics, Inc., St. Louis, MO, USA) and cut longitudinally normal to the bone surface on a Cryo-Cut One cryostat (Vibratome, Bannockburn, IL, USA). The 30 μm sections were examined with a Senterra confocal Raman microscope (Bruker Optics, Billerica, MA, USA). Raman spectra from 40 to 4500 cm⁻¹ were collected with 9 to 18 cm⁻¹ resolution using a circularly polarized 20 mW (9 mW at the sample), 532 nm laser beam, X40/0.95 NA objective, and a 50 μm confocal pinhole. The spectra were acquired at 12 points visually selected outside osteocytes along the periosteal to endosteal axis in the midplane of the sections. The accumulation time for each point was 1 minute. The mineral content of bone matrix (PO₄/CH), collagen content of bone matrix (amide III:CH), and mineral/collagen ratio (PO₄/amide III) were determined from the areas under the 1060 (PO₄) and 1240 (amide III) cm⁻¹ peaks. Relative collagen content from Pro:CH ratio (833 to 867 cm⁻¹) was consistent with that evaluated from the amide III:CH ratio but less accurate owing to low intensity of the Pro peak.

Femur µCT

Micro-computed tomography (µCT) was used to quantify femoral geometry, morphology, and mineralization (GE Medical Systems, London, ON, Canada). Femurs from F1 male 2-month-old mice were scanned over 200 degrees of rotation and reconstructed on 18 μm voxels. Cortical and trabecular analyses included measures of bone mineral content and density, cortical and trabecular thickness, and trabecular spacing. Cortical regions of interest (ROIs) consisted of cylindrical segments (radius 1.35 mm, height 3 mm) of the middiaphyses. Trabecular ROIs also consisted of cylindrical segments (radius 0.57 mm, height equal to 10% of total femur length) of the distal femoral metaphyses.
proximal to the growth plates. Separate thresholds were applied to the cortical and trabecular ROIs (2000 and 1200 HU, respectively).

Ex vivo mechanical testing

Femurs were mechanically tested at room temperature using a four-point bending apparatus (858 Mini-Bionix, MTS, Eden Prairie, MN, USA) in a manner described previously by Jepsen. With posterior femoral surfaces kept in tension, displacement was applied at 0.05 mm/s. The upper and lower tines of the four-point support were separated by 6.35 and 2.2 mm, respectively. Custom software was used to determine stiffness, yield, maximum load, and pre- and postyield fracture energy. Standard beam theory was used to calculate yield stress and Young’s modulus based on the mechanical testing data and femoral geometry as measured by μCT.

Statistical analysis

Statistical analysis was performed using InStat and Prism statistical software packages (Graphpad, La Jolla, CA, USA) or SAS statistical software (SAS Institute, Cary, NC, USA). A two-tailed p value of .05 or less for any single analysis was considered statistically significant. Outliers (defined as values exceeding ±2σ from the group mean) were removed from the μCT and four-point bending data sets.

Results

OOA kindred genotyping/pedigree structure

Putative founder couple identification

A 54-year-old woman with a history of fracture, low hip and spine aBMD, and a differential diagnosis of idiopathic osteoporosis or OI was identified in the Amish Family Osteoporosis Study (AFOS). A brother and a niece of the woman also were identified with a history of fracture and low aBMD. gDNA was extracted from blood of the proband’s brother for analysis of COL1A1 and COL1A2 genes. Nucleotide sequencing identified a G-to-T substitution that converts the triple-helical gly-610 codon (GGT) to cysteine (TGT) codon. The mutant T allele variant is a G-to-T transversion at nucleotide 2098 that alters the gly-610 codon (GGT) to a cysteine (TGT) codon. The mutant T allele has been mapped in 64 heterozygous descendants of the putative founder couple born approximately 150 years ago. Four of the founder couple’s offspring have an estimated 800 living descendants, whereas the other five offspring produced no progeny. (Control genotype = GG; OI genotype = GT.)

Fig. 1. OOA COL1A2 G610C putative founder couple pedigree. The COL1A2 variant is a G-to-T transversion at nucleotide 2098 that alters the gly-610 codon (GGT) to a cysteine (TGT) codon. The mutant T allele has been mapped in 64 heterozygous descendants of the putative founder couple born approximately 150 years ago. Four of the founder couple’s offspring have an estimated 800 living descendants, whereas the other five offspring produced no progeny. (Control genotype = GG; OI genotype = GT.)

an average of approximately 6 individuals. The overall sibship genotype distribution exhibited a normal Mendelian 1:1 ratio expected for offspring from one carrier parent and one wild-type parent. No homozygous individuals and no carrier × carrier couples were identified.

OOA kindred phenotype

Standing height

Standing height as a function of age (Fig. 2) suggested that OI family members tended to be shorter than their unaffected family members. Standing height was converted to Z-scores to better assess the effect of the T allele across the wide age range of study participants. OOA-specific standing height Z-scores were generated separately for women and men from height data obtained from 1687 females and 1416 males (age > 20 years) enrolled in non-OI research studies (unpublished UMB data). Since OOA-specific data were unavailable for study participants 6 to 20 years of age, standing height Z-scores were generated using the National Health and Nutrition Examination Survey (NHANES) data (https://web.emmes.com/study/ped/resources/htwtcalc.htm). Mean standing height Z-score was significantly less (p < .001) for carriers of the T allele as compared with control family members but with appreciable overlap in Z-scores between control and OI study participants (see Fig. 2).

Biomarker measurements

PICP values for OI family members were relatively constant as a function of age (see Fig. 2). However, PICP for control family
members younger than 20 years of age were markedly higher compared with OI family members. Genotype (p < .0001) and age (p < .0086) were found to make significant contributions in a multiple regression model using age, sex, and genotype as variables for PICP. In contrast, only age (p < .0001) was found to make a significant contribution to serum CrossLaps.

**BMD measurements**

Lumbar spine (L1–4) and femoral neck (FN) BMD by age shown for males and females suggested that the OI group has reduced BMD compared with family controls. Age, sex, genotype, and weight were used as predictor variables in multiple regression models for lumbar spine and FN BMD. Sex (p < .03) and genotype (p < .0001) made significant contributions to lumbar spine and FN BMD. Weight (p < .0001) also contributed to lumbar spine BMD, and age (p < .0001) contributed to FN BMD. BMD assessed as Z-scores for the spine and hip are shown in Fig. 3. Mean Z-score differences between the OI and control groups at the lumbar spine and the FN were significant (p < .0001). The mean Z-score at the lumbar spine was −2.63 ± 0.99 for the OI group and −0.24 ± 0.82 for the control group. Mean FN Z-scores were −1.18 ± 0.91 for the OI group and 0.26 ± 0.78 for the control group.

Production of the knock-in mice

Nucleotide gDNA sequence (GenBank Accession Identifier DQ377843) analysis demonstrated that F1 founder mice (male = 1, female = 2) were heterozygous for the knock-in mutation in Cola2 exon 33 (equivalent to COL1A2 exon 35). Intron 32 for the F1 founder knock-in allele [Col1a2^{tm1Mcbr}] Mouse Genome Informatics (MGI) ID 3805459] contains 2712 nucleotides, whereas the wild-type B6 allele contains 730 nucleotides. F1 founder mice cDNA sequence analysis confirmed expression of mutant and wild-type mRNA. Offspring carrying one or two copies of the knock-in Col1a2^{tm1Mcbr} allele from F1 founder breeding survived without any detectable lethality at birth or beyond weaning and were fertile.

Genomic DNA sequence analysis confirmed the Cre-mediated excision of the Floxed neo cassette in an F1 male mouse that was subsequently used to sire G610C OI mice progeny on a B6 background. Intron 33 in G610C OI mice is comprised of 852 nucleotides, and its MGI allele designation is Col1a2^{tm1Mcbr} (MGI Accession Identifier 3711122). It differs in size from the wild-type B6 sequence by the addition of 144 bp of residual targeting vector sequence that flanked the lox P sites and the loss of 22 bp of B6 sequence (GenBank Accession Identifier DQ377844). Heterozygous × wild-type breeding produced litters with both expected genotypes. However, no viable homozygous offspring were obtained from heterozygous matings. Genotype analysis of dead pups recovered within 24 hours of birth did demonstrate the presence of pups homozygous for the Col1a2^{tm1Mcbr} allele in the litters.

Human and mouse SDS-PAGE analysis

Double labeling with Cy5 and Bodipy-L-Cys revealed intense Cys staining of α2(I) chains from all mutant animals and proband collagens (Fig. 4A, B), indicating the presence of exposed reactive Cys-SH residues in α2(I) chains of type I collagen triple helices. Interestingly, SDS-PAGE also revealed the presence of reducible α2-Cys-S-S-Cys-α2 dimers in tendons of all mutant animals (see Fig. 4C, D). Since type I collagen triple helix contains only one α2...
chain, such dimers must form between two different molecules. The intermolecular dimers likely form prior to fibrillogenesis because in the fibrils the Cys residues of adjacent triple helices are axially separated by at least one D period. Dimer formation likely occurs in the Golgi stack by nonspecific side-by-side interactions of procollagen molecules. Surprisingly, the aberrant intermolecular dimers are incorporated into the tissues of mutant animals. From the intensities of the bands containing the S-S dimers and corresponding bands without the dimers, the estimated fraction of mutant chains involved in the dimers was approximately 1% in heterozygous neo+, approximately 1.5% in heterozygous neo–, and approximately 9% in homozygous neo+ animals (Table 1).

SDS-PAGE analysis also demonstrated an abnormally high ratio of α1(I):α2(I) fluorescence intensities in neo+ (Col1a2tm1Mcbr allele) animals, approximately 3:1 in heterozygous neo+ and approximately 8:1 in homozygous neo+ versus approximately 2:1 in wild-type and heterozygous neo– (Col1a2tm1.1Mcbr allele) animals. Most likely this was caused by insufficient synthesis of mutant α2(I) chains and formation of α1(I) homotrimers [e.g., owing to partial degradation and/or slower transcription, splicing, and/or translation of the neo+ (Col1a2tm1Mcbr) mRNA containing the intron 33 targeting vector]. Based on this assumption, we estimated the homotrimer content as approximately 23% and approximately 67% in tendons of heterozygous and homozygous neo+ animals correspondingly (see Table 1).

Human and mouse DSC analysis

Previous differential scanning calorimetry (DSC) studies showed that the denaturation temperature \( T_m \) of type I collagen homotrimers at acidic pH is approximately 2.5°C higher than that of normal heterotrimers.\(^{28,37} \) Thus, to test whether the high α1(I):α2(I) ratio was consistent with homotrimer formation, we performed DSC scans of collagen from all mouse tendons and human fibroblast cultures in 2 mM HCl (pH 2.7). In addition to the denaturation peak of normal type I heterotrimers, the DSC thermograms of collagen from neo+ animals (see Fig. 4E) did reveal a low-temperature peak consistent with mutant heterotrimers and a high-temperature peak. The high-temperature peak was identical to oim and artificial homotrimers measured at the same conditions.\(^{28,38} \) This peak was absent in collagen from neo– animals (see Fig. 4F) and from human proband fibroblasts (see Fig. 4G). Deconvolution of the DSC thermograms suggested approximately 24% and approximately 62% content of homotrimers in heterozygous and homozygous neo+ animals, respectively, in agreement with SDS-PAGE.

To evaluate the effect of the G610C substitution on the thermal stability of mutant collagen, we measured similar DSC thermograms in 0.2 M sodium phosphate and 0.5 M glycerol (pH 7.5), which can be used to evaluate the \( T_m \) under physiologic conditions by subtraction of 1.7°C. From the buffer-corrected thermograms (see Fig. 4H–J), we estimate that the incorporation of the mutant α2(I) chain results in a \( T_m \) reduction by 2.3°C in mouse and approximately 1°C in human collagens correspondingly.

**F1 G610C OI mouse phenotype**

**Breeding**

A total of 301 experimental mice were obtained from 48 litters. Genotype was determined for the mice, except for one B6.Cby female mouse. Mean litter sizes differed among the four F1 stains (one-way ANOVA \( p = .0039 \)). The mean litter sizes were 7.9 ± 1.9 (FVB.B6), 6.5 ± 2.5 (Cby.B6), 5.0 ± 2.2 (C3.B6), and 5.0 ± 1.5 (A.B6).
Goodness-of-fit testing indicated a significant deviation from expected genotype ratios looking at data from all litters ($p = .008$). However, no individual strains had a significant deviation from expected ratios.

**Body weight**

There was a curvilinear relation between body weight (measured longitudinally on days 21 to 60) and age (Fig. 5; age and age squared both $p < .0001$). Wild-type animals were heavier than OI animals (difference 1.77 g, SE 0.3279, $p < .0001$). The A.B6 mice had the lowest and the FVB.B6 mice had the greatest body weight. FVB.B6 mice were significantly heavier than A.B6 mice (difference 2.07 g, $p < .0001$). FVB.B6 mice were significantly heavier than Cby.B6 mice (difference 1.37 g, $p < .003$ (both $p$ values adjusted for multiple comparisons using Tukey-Kramer). There was no evidence of a strain-by-genotype interaction (strain × genotype $p > .8$).
**Table 1. Differential Scanning Calorimetry (DSC) and SDS-PAGE Analysis of Type I Collagen**

| Sample  | Genotype | DSC | SDS-PAGE |
|---------|----------|-----|----------|
|         |          | $\alpha_1\alpha_2$, % | $\alpha_1\alpha_2^m$, %$^a$ | $\alpha_1\beta$, % | Reactive Cys-SH | $\alpha_2^m$-S-S-$\alpha_2^m$, %$^a$ |
| Mouse   | GG       | 100 | 0        | 0        | 0 | 0 |
| Mouse neo+ | GT      | 59 ± 5$^b$ | 17 ± 1$^b$ | 24 ± 5$^b$ | 23 ± 5 | Yes | 1 ± 0.3 |
| Mouse neo+ | TT      | 0 | 38 ± 1$^b$ | 62 ± 1$^b$ | 67 ± 5 | Yes | 9 ± 3 |
| Mouse Neo− | GT      | 51 ± 1$^b$ | 49 ± 1$^b$ | 0 | 0 | Yes | 1.5 ± 0.3 |
| Human   | GG       | 100 | 0        | 0        | 0 | No | — |
| Human   | GT       | 50 ± 10$^b$ | 50 ± 10$^b$ | 0 | 0 | Yes | — |

$^a$The superscript m identifies mutant chains.

$^b$Estimated deconvolution error.

aBMD of G610C OI mouse femurs

G610C OI mice had lower BMC ($\Delta = 0.005 \text{ g, } p < .001$) and BMD ($\Delta = 0.005 \text{ g/cm}^2, p = .001$). Mean aBMD rank order of the G610C OI mice by maternal background strain for was A.B6 < Cby.B6 < FVB.B6 < C3.B6. Pairwise comparisons of the G610C OI mice using the Scheffe multiple-comparisons adjustment procedure resulted in a statistically significantly difference between A.B6 OI and C3.B6 OI ($\Delta = 0.007 \text{ g/cm}^2, p = .0061$). A strain-by-genotype interaction was not significant for either BMC or BMD ($p \approx .9$).

Confocal Raman microspectroscopy

Mineral content of bone matrix (PO4:CH) (Fig. 6) was found to be affected by genotype ($p < .0001$) and mouse strain ($p = .009$). A significant curvilinear relationship was observed between mineral content and bone location ($p = .0065$). The mineral-collagen ratio (PO4:amide III) also was found to have a significant genotype ($p < .0001$) effect and a curvilinear relationship between mineral content and location ($p = .047$) but no evidence of a strain effect ($p = .36$). The curvilinear relationships for PO4:CH and PO4:amide III had peak values located around the middle of the cortical bone. Only genotype was found to have a significant ($p < .0001$) effect for the collagen content of bone matrix (amide III:CH). Mouse strain-by-genotype interaction was not significant for any of the Raman data.

$\mu$CT

Table 2 shows mean values of selected trabecular and cortical bone parameters obtained from isolated femora. Cortical shape and femur length were qualitatively very similar for each
background strain control-OI pair (data not shown). Overall, the μCT trabecular and cortical data show that all G610C OI mice carrying one copy of the Col1a2tm1.1Mcbr allele had less bone as measured by bone volume fraction. The trabecular bone was distinguished by fewer and more widely spaced trabeculae. Except for the FVB.B6 mice, the mean cortical thickness and area were reduced by the presence of the Col1a2tm1.1Mcbr allele. The FVB.B6 control and OI groups had similar mean values for cortical thickness and area. All the OI femurs had reduced moment of inertia ($I_{yy}$) compared with age-matched genetic background controls. The cortical volumetric tissue mineral content (vTMC) was lower in all OI mice relative to their genetic background controls, except for the FVB.B6 OI group, which was the same as the FVB.B6 control group. The cortical volumetric tissue mineral density (vTMD) was elevated in all mice carrying Col1a2tm1.1Mcbr allele relative to their genetic background controls. The trabecular vTMD was statistically the same for each genetic background pair.

Four-point bending biomechanics

Isolated femurs were loaded to failure in a four-point bending testing apparatus to assess the impact of a single copy of the Col1a2tm1.1Mcbr allele and the role of genetic background on the structural phenotype (Table 3). Specifically, femurs from mice with one copy of the variant allele were significantly weaker and more brittle than femurs from control mice. In addition, a genotype-by-genetic-background (strain) interaction was significant for failure load, stiffness, postyield ultimate displacement, and energy to failure. The presence of the Col1a2tm1.1Mcbr allele resulted in a striking reduction in failure load for all F1 OI mice compared with their strain-specific controls. The mean failure load range was greater for control mice (24 N for A.B6 mice to 35 N for C3.B6) compared with OI mice (15 N for A.B6 mice to 18 N for C3.B6 mice). The impact of the Col1a2tm1.1Mcbr allele on the magnitude of the intrastrain control-OI difference in stiffness was variable. The greatest intrastrain OI-control mean difference was found in the C3.B6 group, where the Col1a2tm1.1Mcbr allele was associated with a 71% reduction in mean stiffness. The OI mice in the other three groups had mean stiffness values of 82% (A.B6), 91% (Cby.B6), and 98% (FVB.B6) of their intrastrain controls. Mean values for postyield ultimate displacement for OI mice were 45% (Cby.B6), 52% (C3.B6), and 59% (FVB.B6) of strain-matched control values. However, the mean value for A.B6 OI mice was 95% of the control value. Failure energy ranged from 0.78 (Cby.B6) to 1.14 N-mm (C3.B6) for the OI mice and from 3.11 (A.B6) to 6.3 N-mm (FVB.B6) for controls.

Ultimate stress and Young’s modulus had statistically significant genotype and strain differences, but the genotype-by-genetic-background interaction was not significant (see Table 3). Mean ultimate stress was reduced for all OI groups relative to their genetic background controls and ranged from 67% (Cby.B6 OI mice) to 81% (C3.B6 OI mice). Young’s modulus was increased across all OI groups carrying the Col1a2tm1.1Mcbr allele relative to their genetic background controls. The magnitude of the mean Young’s modulus was not different between OI-control pairs but varied by genetic background of the strain.

Fig. 6. F1 G610C OI mouse Raman microspectroscopy. The collagen content of the OI (heterozygous for the Col1a2tm1.1Mcbr allele) femurs was reduced relative to wild-type (WT) control femurs, and the collagen was hypermineralized. A representative Raman spectrum illustrates the peaks used for data analysis (upper left), and comparison of chemical-specific ratios in WT (black bars) and G610C OI (gray bars) femurs determined from the Raman spectra are shown in the balance of the figure.
| Mean                          | A.B6  | Cby.B6 | FVB.B6 | C3.B6 | ANCOVA p value |
|------------------------------|-------|--------|--------|-------|----------------|
| SD                           |       |        |        |       | Interaction    |
| Trabecular BV/TV (%)         | 0.082 | 0.200  | 0.148  | 0.172 | <.0001         |
|                              | 0.026 | 0.076  | 0.020  | 0.068 | <.0001         |
|                              |       | 0.046  | 0.014  | 0.026 |                |
| Trabecular number            | 2.923 | 4.793  | 4.501  | 4.051 | <.0001         |
|                              | 0.781 | 1.114  | 0.434  | 1.197 | <.0001         |
|                              |       | 0.859  | 0.473  | 0.713 |                |
| Trabecular vBMD              | 469.400 | 526.000 | 459.900 | 533.300 | <.0001 |
|                              | 10.860 | 33.330 | 7.096  | 18.550 |                |
| Cortical bone vol. fraction (%) | 0.12   | 0.15   | 0.14   | 0.16   | 0.0309 |
|                              | 0.0076 | 0.019  | 0.0083 | 0.016  |                |
|                              |       | 0.014  | 0.011  | 0.014  |                |
| Cortical thickness (mm)      | 0.18   | 0.20   | 0.18   | 0.23   | 0.0094 |
|                              | 0.0084 | 0.019  | 0.0067 | 0.015  |                |
|                              |       | 0.016  | 0.011  | 0.014  |                |
| Cortical area (mm²)          | 0.66   | 0.81   | 0.76   | 0.904  | 0.0154 |
|                              | 0.044  | 0.102  | 0.047  | 0.086  |                |
|                              |       | 0.079  | 0.061  | 0.077  |                |
| Cortical Iₚₓ (mm⁴)           | 0.082  | 0.13   | 0.12   | 0.13   | <.0001         |
|                              | 0.0090 | 0.027  | 0.014  | 0.022  | <.0001         |
|                              |       | 0.016  | 0.015  | 0.021  |                |
| Cortical vTMC (mg)           | 2.00   | 2.55   | 2.28   | 2.84   | 0.0124 |
|                              | 0.14   | 0.38   | 0.15   | 0.29   |                |
|                              |       | 0.28   | 0.20   | 0.30   |                |
| Cortical vTMD (mg/mL)        | 999.48 | 1019.20 | 969.32 | 1048.21 | 0.0052 |
|                              | 9.51   | 22.83  | 13.26  | 14.21  |                |
|                              |       | 21.07  | 22.46  | 26.27  |                |
The ascertainment of 64 related individuals harboring the mutant T allele represents the largest reported collection of OI patients with an identical collagen mutation. Amino acid substitutions at invariant glycine residues typically result in clinically apparent phenotypes and make ascertainment more likely. While the original brother-sister-niece proband cluster presented with some of the most severe clinical signs and symptoms of OI, our overall clinical data support a Sillence type I (mild) to type IV (moderate) phenotype range. Pedigree analysis suggests that the T allele was a de novo mutation that occurred in the putative founder couple born circa 1860. The rapid OOA population growth from approximately 600 individuals in 1860 to about 31,000 children and adults today,(39) combined with stochastic events, endogamy, and large sibships, likely contributed to the mutation's expansion within the Lancaster Settlement.

Fracture history is the most relevant phenotypic trait used to make a clinical diagnosis of OI and to classify disease severity. However, since radiographic and medical record documentation of fractures were unavailable in the OOA, the use of DXA measurements of aBMD were used as a continuous quantitative trait to assess phenotypic severity that reflected the Sillence OI type clinical spectrum. Prior work suggests that mean aBMD for children and adults with OI types III and IV is significantly reduced compared with controls, but a significant number of OI type I patients had aBMD values within the reference range.(40–42) Non-OOA OI patients often have L1–4 spine Z-scores in the −2 to −4 range for type IV OI patients (JCM clinical experience). These stratification ranges demonstrate that the T allele was associated with a wide range of disease severity (Table 4) in the spine and a milder phenotype in the hip.

Table 3. Structural and Predicted Material Parameters Derived From Four-Point Bending

| Mean | A.B6 | Cby.B6 | FVB.B6 | C3.B6 | ANCOVA p value |
|------|------|--------|--------|-------|----------------|
| SD   | WT   | OI     | WT     | OI    | WT            |
| Failure load (N) | 24.24 | 14.86 | 28.76 | 15.20 | 34.51         |
|       | 1.92  | 2.68   | 6.02  | 2.35  | 2.78          |
| Stiffness (N/mm) | 187.48 | 154.51 | 190.78 | 174.06 | 194.81 |
|       | 26.11 | 13.85  | 37.66 | 28.092 | 28.18 |
| Postyield displacement (mm) | 0.0037 | 0.0036 | 0.0058 | 0.0026 | 0.0063 |
|       | 0.00053 | 0.00013 | 0.0018 | 0.00016 | 0.00010 |
| Ultimate stress (MPa) | 425.00 | 334.04 | 415.82 | 276.80 | 377.12 |
|       | 53.42 | 61.37  | 40.79 | 47.44  | 30.10 |
| Young's modulus | 8.025 | 9.53   | 6.124 | 7.75   | 5.88 |
|       | 1.63  | 1.24   | 1.14  | 1.04   | 0.99  |
| Yield energy | 0.69  | 0.67   | 1.44  | 0.602  | 0.72  |
|       | 0.17  | 0.19   | 0.50  | 0.13   | 0.086 |
| Failure energy (N-mm) | 3.11  | 0.88   | 5.66  | 0.78   | 6.30  |
|       | 0.56  | 0.35   | 1.24  | 0.25   | 1.41  |

Table 4. DXA Z-Score Stratification

| Sillence classification | Total number | Control (GG) | OI (GT) |
|-------------------------|--------------|--------------|---------|
| Normal                  |              |              |         |
| −2 to −4                |              |              |         |
| −4                      |              |              |         |
| Type I                  |              |              |         |
| Type II                 |              |              |         |
| Type III                |              |              |         |
| Type IV                 |              |              |         |

- Spine
- Femoral neck

AMISH OI MUTATION

Pedigree analysis suggests that the T allele was a de novo mutation that occurred in the putative founder couple born circa 1860. The rapid OOA population growth from approximately 600 individuals in 1860 to about 3,500 children and adults today, likely contributed to the mutation's expansion within the Lancaster Settlement.

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| −2 to −4                |              |              |         |
| −4                      |              |              |         |
| Type I                  |              |              |         |
| Type II                 |              |              |         |
| Type III                |              |              |         |
| Type IV                 |              |              |         |

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|       | 0.56  | 0.35   | 1.24  | 0.25   | 1.41  |
fracture, have a shorter final standing height in the range of puberty and adolescence, some 90% of skeletal mass is attained, and peak stature normally is attained by approximately 20 years of age. The cross-sectional serum biomarker data suggest impaired skeletal accretion and linear bone growth in T allele carriers; PICP levels were relatively flat across the entire age spectrum for T allele carriers (see Fig. 3). In normal, healthy children, PICP levels reflect collagen biosynthesis associated with skeletal growth and are known to change significantly with age and pubertal development stages. In contrast to the PICP estimates of age and sex patterns for both the OI and control groups. This suggests an uncoupling between bone formation and resorption for the OI group during a major period of bone mass accretion that favors a net decrease in bone mass.

The OOA kindred and the G610C OI mouse share the classic OI features of dominant inheritance and a triple-helical glycine substitution. The biomedical significance of this kindred is its clinical and quantitative prototypic varied disease severity that is further enhanced by its rich resource of phenotype data, DNA, and fibroblast cell lines. However, repeated access to kindred bone samples and other tissues for experimental uses are unlikely, thus necessitating creation of the G610C OI mouse to aid future studies. Establishing the validity of the G610C OI mouse as the first α2(II) glycine substitution model required the following elements: (1) duplication of the codon change, (2) expression of mutant proα2(II) transcript, (3) deposition of mutant protein in the ECM, and (4) confirmation of a bone phenotype.

Our initial evaluation of founder (neo+) mice and their progeny carrying the Col1a2tm1 mice documented that the G610 Codon was altered as planned and the mutant transcript was expressed. Surprisingly, these mice secreted a significant fraction of type I collagen homotrimer in addition to G610C-containing heterotrimeric collagen. The formation of homotrimers appeared related to intron 32 targeting vector retention and fibroblast cell lines. However, repeated access to kindred bone samples and other tissues for experimental uses are unlikely, thus necessitating creation of the G610C OI mouse to aid future studies. Establishing the validity of the G610C OI mouse as the first α2(II) glycine substitution model required the following elements: (1) duplication of the codon change, (2) expression of mutant proα2(II) transcript, (3) deposition of mutant protein in the ECM, and (4) confirmation of a bone phenotype.

Given that phenotype variation was a significant clinical feature observed among the OOA kindred, the impact of the genetic background on phenotype severity was tested in G610C OI mice carrying the Col1a2tm1 mice. Heterozygous x wild-type mouse crosses resulted in good reproductive fitness that matched the reproductive fitness of the OOA kindred and yielded F1 strains with approximately 50% interstrain genetic background identity. Interestingly, reproductive fitness of heterozygous x heterozygous B6 matings was poor because trial matings failed to produce offspring homozygous for the Col1a2tm1.1Mcbr allele that survived beyond the perinatal period.

Growth curves (body weight) from weaning to 2 months of age demonstrated impaired weight gain for all four F1 mouse strains carrying the Col1a2tm1.1Mcbr allele relative to their controls. This is a likely generalizable effect in mice caused by type I collagen gene mutations because it has been reported for the Mov13, oim, and BrlIV OI models. Body weight is a complex quantitative trait, and different inbred mice are exemplified by characteristic patterns of weight gain with time (MGI Database). It is likely that the different growth curves reflect the maternal contribution to the genetic background of the F1 mice because inbred mouse strains exhibit characteristic patterns of weight gain with age.

Since the degree and nature of bone matrix mineralization may be an important determinant of resistance to fracture, whole-femoral mineral mass in F1 mice was assessed by bBMD and µCT measurements. PIXImus measurements (Fig. 7) demonstrated that all strains of mice carrying the Col1a2tm1.1Mcbr allele had a reduction in aBMD compared with controls, and this is consistent with DXA data from the OOA kindred. However, µCT measurement of increased cortical vTMD (see Table 3) better reflects the tissue effects of the variant Col1a2 allele. This apparent contradiction is explained by the 2D nature of aBMD estimated by DXA versus the 3D data obtained by µCT. The µCT data suggest the G610C OI mice have reduced total tissue mineral content but also have reduced total bone mass that results in greater bone mineralization compared with control mice. Hypermineralization at the level of collagen fibers was confirmed by Raman microspectroscopy and is consistent with higher mean mineralization density in OI patient bone samples measured by backscattered electron imaging.

Structural integrity of femoral cortical bone was assessed using four-point bending, which is an ex vivo surrogate in mice for increased long bone fracture risk associated with OI. The structural (failure load) and material property (ultimate stress) measurements of femur strength at 2 months of age demonstrated that a single copy of the Col1a2tm1.1Mcbr allele reduced mechanical integrity regardless of genetic background. Key
contributors to the structural soundness of bone include geometric parameters (i.e., size, shape) and inherent material properties such as the molecular integrity, quantity and organization of type I collagen, and degree and nature of collagen mineralization. While the mineral content of bone contributes substantially to its stiffness, collagen tends to dominate the properties of the postyield region of the load-deformation curve. It would appear from the yield energy-total energy ratio that once the bone starts to break, the presence of the mutated collagen offers less resistance to complete failure in G610C OI mice. The failure of the abnormal collagen heterotrimer to “toughen” the bone is likely due to a structural and/or organizational defect caused by the mutation and/or a general reduction in the amount of collagen per unit volume of bone. Our human and mouse findings of reduced DSC denaturation temperature Tm suggest that the global molecular (structural) stability of the mutant collagen heterotrimer is weakened. This reduction in Tm is commonly associated with OI collagen mutations, although there is no simple relationship between the magnitude of Tm decrease and OI disease severity. Reduced bone matrix collagen content (measured by Raman microspectroscopy) also appears to contribute to loss of femoral structural integrity in G610C OI mice. The variations in the murine bone biomechanical properties across the several experimental strains used in this study (genotype-by-strain interactions) provide additional insight about skeletal complications among human patients. The effect of the mutation on the estimated bone tissue properties was the same across all strains, each demonstrating significant and similar reductions in inherent tissue integrity. However, the magnitude of structural property alterations (such as failure load) appears to be dependent on genetic background (strain). Taken together, these results suggest that the effect of the mutation on basic, inherent bone tissue integrity is similar in all OOA kindred patients (reduced modulus and increased brittleness). However, the consequence of the defect on whole-bone fracture susceptibility is influenced by individual genomic factors that are reflected in size, shape, and possibly bone metabolic regulation.

Mouse models have the advantage over human samples of tight control of genetic background and the ability to circumvent or minimize the role of environment on phenotype. The F1 G610C OI screening crosses were successful in demonstrating that genetic background variation affected bone geometry and structural property phenotype. We speculate that additional backcrosses will further magnify interstrain phenotype differences and enhance the probability of identifying major genetic modifiers using standard and novel backcross and intercross genetic mapping tools with proven success in the detection of human disease gene modifiers. Bone trait modifiers identified in the G610C OI mouse likely will include Col1a1 or Col1a2 mutation-specific modifiers, some that are generalizable for many OI mutation loci, and others that overlap with genes associated with common forms of osteoporosis. Quantitative trait locus (QTL) investigation in the G610C OI mouse also likely will confirm QTLs found in prior osteoporosis studies, thus narrowing the critical QTL regions. The OOA OI kindred and AFOS data then can be used to confirm both sets of genes—those common to osteoporosis and those unique to OI.

In summary, the OOA kindred is the largest known OI kindred with an identical type I collagen defect and extensive phenotype data. The triple-helical glycine substitution and dominant inheritance pattern are prototypic for most reported cases of OI. While all the affected individuals share the same mutation, they varied widely in phenotype, as quantified by aBMD Z-scores. The G610C OI mouse, patterned on the OOA family, duplicates its gene and protein profile. The magnitude of structural property alterations owing to the mutation was dependent on genetic background (strain). The bone geometry and structural property gene × genetic background interactions suggest that noncollagen genes can modify the G610C OI mouse OI phenotype. Overall, these results demonstrate that G610C OI mice represent a unique translational OI model to define the complex genetic architecture controlling phenotype and for testing of potential therapeutic approaches and treatment strategies for OI.

Disclosures

The authors state that they have no conflicts of interest.

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