Mutations in **fam20b** and **xylt1** Reveal That Cartilage Matrix Controls Timing of Endochondral Ossification by Inhibiting Chondrocyte Maturation

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**Abstract**

Differentiating cells interact with their extracellular environment over time. Chondrocytes embed themselves in a proteoglycan (PG)-rich matrix, then undergo a developmental transition, termed “maturation,” when they express *ihh* to induce bone in the overlying tissue, the perichondrium. Here, we ask whether PGs regulate interactions between chondrocytes and perichondrium, using zebrafish mutants to reveal that cartilage PGs inhibit chondrocyte maturation, which ultimately dictates the timing of perichondral bone development. In a mutagenesis screen, we isolated a class of mutants with decreased cartilage matrix and increased perichondral bone. Positional cloning identified lesions in two genes, *fam20b* and *xylosyltransferase1* (*xylt1*), both of which encode PG synthesis enzymes. Mutants failed to produce wild-type levels of chondroitin sulfate PGs, which are normally abundant in cartilage matrix, and initiated perichondral bone formation earlier than their wild-type siblings. Primary chondrocyte defects might induce the bone phenotype secondarily, because mutant chondrocytes precociously initiated maturation, showing increased and early expression of such markers as *runx2b*, *collagen type 10a1*, and *ihh* co-orthologs, and *ihha* mutation suppressed early perichondral bone in PG mutants. Ultrastructural analyses demonstrated aberrant matrix organization and also early cellular features of chondrocyte hypertrophy in mutants. Refining previous *in vitro* reports, which demonstrated that *fam20b* and *xylt1* were involved in PG synthesis, our *in vivo* analyses reveal that these genes function in cartilage matrix production and ultimately regulate the timing of skeletal development.

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**Introduction**

Vertebrate bone is produced by two major developmental processes, intramembranous ossification (forming dermal bones) and endochondral ossification (forming chondral bones). The latter process has many stages that must be coordinated in space and time. First, some cells in the mesenchymal condensation differentiate as chondrocytes, which secrete a cartilage extracellular matrix rich in proteoglycans (PGs), such as chondroitin sulfate PGs [1,2]. A thin layer of cells, the perichondrium, surrounds the developing cartilage. Later, a subset of chondrocytes undergoes a developmental transition, expressing markers of chondrocyte maturation, including *indian hedgehog* (*ihh*) and *collagen type 10a1* (*col10a1*) [3]. Meanwhile, cells of the perichondrium that overlie the maturing chondrocytes differentiate as bone-forming osteoblasts.

Tissue interactions between cartilage and perichondrium ensure that these early events in endochondral ossification are coordinated in space and time [4–6]. Critically, the growth factor Ihh produced by maturing chondrocytes induces perichondral bone, and both mouse *Ihh* and zebrafish *ihha* mutants have delayed endochondral ossification [7,8]. Although specific signaling pathways and transcription factors are involved [9–11], the full ontology of genes that regulate the timing of endochondral ossification is still unknown, but may include genes involved in extracellular matrix production.

Mutations disrupting PG synthesis commonly affect skeletal tissues and can change the timing of skeletal development. Hereditary Multiple Exostoses is caused by mutations in *Exostosin (Exo)* genes, which encode enzymes that synthesize heparan sulfate PGs [12,13]. Mouse and zebrafish models of *Ex* loss-of-function reveal defects in endochondral ossification associated with delays in chondrocyte maturation, which were attributed to altered Ihh signaling [14–16]. Mutations affecting various components of PG synthesis, from sugar precursor production enzymes to sulfation enzymes, affect dramatically the shape and composition of skeletal
tissues, and also can delay endochondral ossification [14,17–20]. Interestingly, acceleration or inhibition of developmental timing may be a function of the class of PG. Evidence above indicates that mutations in HSPG synthesis delay endochondral ossification, while mutations in chondroitin sulfate PG (CSPG) synthesis may accelerate this bone-forming process. Aggrecan is the major chondroitin sulfate PG (CSPG) in cartilage, and Aggrecan (Acan) mutant chicks exhibit increased levels of Col10 expression [21], which these authors interpreted as evidence that endochondral ossification had initiated earlier. Here we perform direct investigation of developmental timing in order to test the hypothesis that loss of CSPGs accelerates endochondral ossification.

In a forward genetic screen of skeletal tissues in developing zebrafish larvae, we isolated a class of mutants with increased chondral bone and decreased cartilage matrix at 6 days post-fertilization (dpf). Positional cloning identified mutations in fam20b and xylt1, two critical genes for PG synthesis that are associated with human disease [22–27]. Biochemical and histochemical analyses revealed that mutants in both genes have defects in CSPG production. Rescue experiments, gene expression studies, ultrastructural, and mutational analyses all argue that PG mutant chondrocytes undergo precocious maturation, thus inducing early perichondral bone. Our analyses of in vivo models for loss of function in fam20b or xylt1 refine previous in vitro reports [23,28] by demonstrating an important role for these PG synthesis genes in cartilage matrix production in intact, developing animals. In closing, we discuss the phenotype of genes in cartilage matrix production in intact, developing animals.

Results

Two mutant loci with increased bone matrix and decreased cartilage matrix phenotypes

Screening of larval skeletal phenotypes in mutagenized zebrafish revealed a class of mutant with normal gross anatomy at 6 dpf, but with specific defects in the degree of cartilage and bone tissue formation (Figure 1A, 1B). We identified four independently derived lines (b1125, b1127, b1128, and b1189), in which homozygous mutants showed increased ossification of many bones, as judged by Alizarin red staining (Figure 1C–1H). In addition, these mutants had decreased Alcian blue staining of cartilage matrix in all cartilaginous elements (Figure 1C–1H), which reflected defects in the production or secretion of cartilage extracellular matrix. Cartilage and bone phenotypes were both present in 100% of mutants examined, although there was variation in the degree to which Alizarin red staining increased. Complementation crosses, scored for these skeletal phenotypes, suggested mutations at two genetic loci among these four mutants. The b1125 and b1127 mutations failed a genetic complementation test, producing 51/212 (24%) mutant larvae when heterozygotes were crossed to one another. Likewise, b1128 and b1189 failed to complement, producing 60/208 (29%) mutant larvae when heterozygotes were crossed. All other pair-wise complementation crosses produced larvae with wild-type skeletal development. The b1127;b1128 double mutants showed Alizarin red staining that was similar to each single mutant, although the loss of Alcian blue staining seemed more severe (Figure 1F, quantified below), suggesting that both of these loci drive production of cartilage matrix. Other than these changes to skeletal tissues, overall mutant cranial morphology appeared smaller at 6 dpf (Figure 1A, 1B). In particular, mutant chondral bones appeared shorter than those in wild-type siblings. All homozygous mutants can grow to viable adults, allowing us to investigate whether morphological phenotypes were more apparent at later stages. Compared to wild-type siblings, mutant adults displayed foreshortened upper and lower jaws, hypoplastic midface, and bulging eyes (Figure 1I, 1J). Imaging Alizarin red-stained head skeletons with optical projection tomography (OPT) revealed that morphological defects in adult mutant heads were accompanied by altered craniofacial skeletal morphology (Figure 1K, 1L), predominantly an apparent loss of anterior neurocranial growth.

Identification of fam20b and xylt1 lesions in skeletal matrix mutants

Genetic mapping confirmed that the two complementation groups of skeletal mutants mapped to independent loci, and sequencing nearby candidates revealed molecular lesions in fam20b and xylt1 that underlie the skeletal phenotype. RAD mapping and subsequent simple sequence repeat (SSR) mapping identified a genetic interval of 0.2 cM on LG20 containing the b1127 mutation (Figure 2A; [29]; see Materials and Methods). The lesion in fam20b [b1127] mutants (991T>C) disrupted a highly conserved cysteine residue (C331R; Figure 2B, 2C; [30]). For the b1125 allele, cDNA sequencing revealed a 1162C>T mutation in the seventh coding exon of fam20b, which would alter amino acid 388 from Gln to STOP, thereby truncating the last 22 amino acids, including a highly conserved Gys residue at aa389 (Figure 2B, 2C; [30]). fam20b [b1125] transcripts were down-regulated at 55 hours post-fertilization (hpf), while fam20b [b1125] transcripts were present at wild-type levels (data not shown), which is consistent with nonsense-mediated RNA degradation [31]. Therefore, mapping and sequence data suggested that mutations within fam20b caused the b1125 and b1127 phenotypes.

As the first steps in identifying molecular lesions in xylt1, RAD mapping and subsequent SSR mapping defined a genetic interval of 0.7 cM on LG3 containing the b1128 mutation (Figure 2D; [29]). Sequencing of xylt1 from b1128 cDNA and gDNA revealed a splice donor mutation (2103G>A) in exon 9 that would produce a frameshifted and truncated C-terminal portion of the zebrafish Xylt1 protein from at least amino acid 702 (out of 919, Figure 2E). For the b1189 allele, cDNA sequencing revealed a 1600T>G mutation in the seventh coding exon of xylt1, which altered
Figure 1. Craniofacial skeletons of mutant zebrafish larvae exhibit increased bone matrix and decreased cartilage matrix. A–H, Alcian blue/Alizarin red-stained 6 dpf larvae, entire heads viewed laterally (A,B; eyes removed) or flat-mounted, dissected pharyngeal skeletons viewed ventrally (C–H). I–L, lateral and dorsal views of entire 8-month-old heads (I,J), and corresponding lateral view images of Alizarin red fluorescence taken by optical projection tomography (OPT; K,L). Overall gross anatomy of the head and craniofacial skeleton are similar in wild types (A) and mutants (B), although staining of cartilage and bone appeared altered in mutants. Compared to wild types (C), fam20b<sup>b1125</sup> (D), fam20b<sup>b1127</sup> (E), xylt1<sup>b1128</sup> (F), and xylt1<sup>b1189</sup> (G) mutants had increased Alizarin red staining of bone (arrows) and decreased Alcian blue staining of cartilage. There was no sided-ness to the phenotype, for defective cartilage and bone appeared symmetrically on left and right sides. Double mutant fam20b<sup>b1127</sup>;xylt1<sup>b1128</sup> larvae (H) had Alizarin red staining (arrow) similar to that seen in single mutants, but more severe loss of Alcian blue staining. Relative to wild-type siblings (I), fam20b<sup>b1127</sup> mutant adults (J) showed foreshortened upper and lower jaws, hypoplastic midface, and bulging eyes.
highly-conserved amino acid 354 from Ser to Ala (Figure 2E, 2F). yxl1b1120 transcripts were down-regulated at 53 hpf, while yxl1b1120 transcripts were present at wild-type levels (data not shown), which is in agreement with nonsense-mediated RNA degradation [31]. These data suggested that the b1120 and b1109 phenotypes were caused by mutations in yxl1.

Expressing wild-type fam20b rescues b1125 and b1127

We used rescue experiments to test our conclusion that lesions in fam20b caused the skeletal phenotypes of b1125 and b1127. Mutant embryos were injected with Tol2 expression plasmids driving expression of wild-type fam20b, and then were assayed for skeletal phenotypes. Wild-type fam20b cDNA rescued fam20b b1125 (n = 10/15) and fam20b b1127 (n = 3/8) skeletal phenotypes when driven by the ubiquitous beta-actin2 promoter (Figure 3A-3D, data not shown; [32]). There was variation in the degree to which the entire skeleton, or even an entire skeletal element, was rescued, which would be expected from mosaicism inherent to the transient injection protocol. We observed in these rescues a correlation between chondrocytes surrounded by faint Alcian blue staining and overlying bone (n = 11 skeletal elements). That is, in adjacent patches of chondrocytes surrounded by faint Alcian blue staining and overlying bone (n = 0/11; Figure S3). Experiments injecting fam20b (n = 0/23) or xylt1b1128 (n = 0/12) cDNA did not show differences in skeletal phenotypes at 6 dpf (Figure 3D, dashed lines), a finding that will be of significance for future studies. Apart from their up-regulated expression in skeletal tissues, fam20b showed diffuse ubiquitous expression in brain and craniofacial mesenchyme, and yxl1 transcripts were expressed in discrete domains of the brain, such as the developing forebrain (Figure 4B, 4C, 4G, 4H). In summary, fam20b and yxl1 are expressed in developing chondrocytes, but not in cells of the perichondrium, even at stages when we know bone-forming cells have differentiated within this tissue. Our findings are thus consistent with the notion that mutations in these genes act directly in chondrocytes to produce the mutant cartilage phenotype, but only indirectly in causing elevated perichondral bone.

fam20b and xylt1 mutants exhibit partial loss of cartilage PGs

Quantifying spectrometrically lysates from at least three Alcian blue-stained clutches for fam20b or yxl1 single mutants, or fam20bxylt1 double mutants (see Materials and Methods; [35]), we found statistically significant changes in Alcian blue levels between all genotypes tested (Figure 5A-5D, 5F; ANOVA p<0.0001). Levels of Alcian blue in fam20b and xylt1 mutants were 50±3.0% and 57±2.0% of their wild-type siblings, respectively, while fam20bxylt1 double mutants further reduced Alcian blue levels to 39±2.6% of their wild-type siblings (Figure 5F). Recently, we showed that a null mutation in UDP-xylose synthase1 (uxs1) abolished zygotic production of UDP-xylose [18], a sugar that ultimately is the substrate for both Fam20b and Xylt1 during PG synthesis (Figure 5G; [22,23]). We used the uxs1 mutant to estimate whether fam20b and xylt1 mutants were null alleles. Relative to fam20b and xylt1 single and double mutants, cartilages from uxs1 mutants showed even less Alcian blue staining (Figure 5E), reducing levels to 22±1.5% of those in wild-type siblings (Figure 5F). Therefore, quantitative comparisons with uxs1 mutants suggested that the fam20b b1125 and xylt1b1120 alleles did not completely eliminate cartilage PG production; even fam20bxylt1 double mutants were less severe than uxs1 mutants.

To show by an independent method that fam20b and xylt1 mutants had cartilage PG defects that were less severe than uxs1 mutants, we analyzed GAG disaccharide levels using biochemical methods. Both heparan sulfate PGs (HSPGs) and chondroitin sulfate PGs (CSPGs) are xylose-dependent, so both of these classes of PG may be affected in fam20b and xylt1 mutants. HPLC analyses on lysates of whole 5 dpf larvae demonstrated that fam20b and xylt1 mutants had decreased levels of CSPG disaccharides, which are predominant in cartilage [36], and again, the losses were less severe than seen in uxs1 mutants (Figure S3). HSPG disaccharide levels were not affected consistently among these mutants; fam20b and uxs1 mutants showed decreases, whereas xylt1 mutants did not (Figure S3).
Figure 2. Mapping the mutants reveals lesions in \textit{fam20b} and \textit{xylt1}. A, SSR markers mapped \textit{b1127} to a 0.2 cM interval on LG20 around \texttt{z20582}, showing 0 cross-overs/641 meioses. B, cDNA sequencing of \textit{fam20b} revealed a T991C mutation in the sixth exon; genotype assays confirmed perfect linkage of this mutation to mutant phenotype (0 cross-overs/862 meioses). Similarly, \textit{fam20b} had a C1162T mutation in the seventh exon. C, The \textit{fam20b} mutation changed a highly-conserved Cys to Arg at aa331 (numbering based upon zebrafish protein), while \textit{fam20b} had a C1162T mutation in the seventh exon. D, SSR markers mapped \textit{b1128} to a 0.7 cM interval on LG3. E, cDNA sequencing of \textit{xylt1} revealed a T1600G mutation in the seventh exon; \textit{xylt1} contained a G2103A splice donor mutation in the ninth exon, which forced usage of cryptic splice donors, typically causing a tetranucleotide insertion. Genotype assays confirmed perfect linkage of these mutations to mutant phenotypes (\textit{xylt1}: 0 cross-overs/632 meioses; \textit{xylt1}: 0 cross-overs/738 meioses). F, \textit{xylt1} mutation changed a highly-conserved Ser to Ala at aa702. Lines in A, B, D, and E are not to scale. Abbreviations: \textit{D.re} = \textit{Danio rerio}; \textit{G.ga} = \textit{Gallus gallus}; \textit{H.sa} = \textit{Homo sapiens}; \textit{X.la} = \textit{Xenopus laevis}. doi:10.1371/journal.pgen.1002246.g002
Figure 3. Wild-type fam20b expression rescues the fam20b<sup>81127</sup> mutant phenotype. A–E, Flat-mounted, dissected pharyngeal skeletons of Alcian blue/Alizarin red-stained 6 dpf larvae. Both sides of the pharyngeal skeletons are shown because the mosaic nature of the rescue experiment produced asymmetric phenotypes in injected larvae. Compared to uninjected wild-type (A) and mutant (B) controls, fam20b<sup>81127</sup> mutant larvae that were injected with wild-type fam20b cDNA under the control of beta-actin2 promoter (C) showed skeletal elements with rescued cartilage matrix staining by Alcian blue and decreased bone matrix staining (asterisk) by Alizarin red. D, Higher magnification of boxed region in C illustrates patches of light Alizarin red staining surrounded by heavy Alcian blue staining (arrow, dashed lines) adjacent to patches of dark Alcian blue staining produced asymmetric phenotypes in injected larvae. Compared to uninjected wild-type (A) and mutant (B) controls, fam20b<sup>81127</sup> mutants embryos similarly injected with fam20b<sup>81127</sup> did not rescue the mutant skeletal phenotype. Abbreviations: UIC = uninjected control.

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Consistent with the biochemical data, immunodetection assays revealed losses of both CSPGs and HSPGs specifically around developing chondrocytes in fam20b mutants (Figure 6A–6F). No loss of CSPGs in xylt1 mutant larvae were detected (data not shown), suggesting that the CSPG antibody may recognize epitopes still present in xylt1 mutants. Together, these biochemical and immunohistochemical data revealed cartilage PG defects in fam20b and xylt1 mutants. Furthermore, the quantitative comparisons with xus1 mutant embryos suggested either that fam20b<sup>81127</sup> and xylt1<sup>11224</sup> are not null alleles, or that redundant genes compensate for their loss of function (see Discussion).

Since both Fam20b and Xylt1 depend upon UDP-xlose, and hence xus1 function, in order to promote GAG synthesis (Figure 5G), loss of xus1 function should mask the skeletal phenotypes of fam20b and xylt1 embryos. Consistent with fam20b and xylt1 being downstream of xus1, skeletal phenotypes of fam20b;xus1 and xylt1;xus1 larvae appeared identical to xus1 single mutants, including severe loss of Alcian blue staining and undetectable levels of perichondral bone, as predicted (Figure 7A–7H). Despite the strong xus1 mutant phenotype, elimination of just one copy of xus1 did not sensitize zebrafish skeletons to the loss of fam20b or xylt1. Trans-heterozygous larvae displayed normal skeletal phenotypes, and no enhancement of the homozgyous fam20b and xylt1 mutant phenotypes was observed when larvae also were heterozygous for the xus1 mutation (data not shown). These epistasis experiments provided genetic support that fam20b and xylt1 function in the PG synthesis pathway in vivo.

Mutants have increased bone due to precocious differentiation of perichondral osteoblasts

We find that two parameters of bone matrix production along the endochondral ossification pathway are increased in both fam20b and xylt1 mutants, which is in contrast to decreased production of cartilage matrix. First, the frequency of chondral bone Alcian red staining at 6 dpf was increased (see Figure 1C–1H for example of material scored), whereas dermal bone staining did not consistently show statistically significant effects (Figure 8A, 8B). Second, the amount of staining in a given chondral bone (e.g., the ceratohyal) was also increased significantly in fam20b and xylt1 mutants (Figure 8C–8G). The increased bone resided on the inner surface of the perichondrium (see Figure 9H, 9I), as expected for chondral, rather than dermal, bones. Therefore, fam20b and xylt1 mutants showed significant increases in the amount of perichondral bone.

The observed increase in perichondral bone might result from three possible scenarios: 1) osteoblasts differentiate at the correct time, but exhibit enhanced secretory activity; 2) osteoblasts differentiate at the correct time, but a larger pool of pre-osteoblasts exists in mutant perichondria; or 3) osteoblasts differentiate early. Examination of skeletal phenotypes and cellular and molecular markers of osteoblasts at time points prior to 6 dpf demonstrated that the increase in mutant perichondral bone resulted from early initiation of osteogenesis. While neither mutants nor wild types showed signs of perichondral bone at 3 dpf (Figure 9A–9C), Alcian red staining was observed in chondral bones of fam20b and xylt1 mutants, but not wild types, at 4.5 dpf (Figure 9D–9F, arrows). Bone formation was not accelerated in time further in fam20b;xylt1 double mutants, as perichondral bone was not detected at 3 dpf, but was prominent by 4.5 dpf (data not shown). To address the cellular basis for early perichondral bone, fam20b and xylt1 mutants were bred into transgenic zebrafish expressing EGFP under a promoter that is restricted to osteoblasts (Tg[tg7:EGFP]<sup>b1212</sup>; [37]). Transgenic mutants showed GFP expression and Alizarin red fluorescence in the perichondrium of chondral bones by 4.5 dpf, whereas their wild-type siblings showed neither of these markers at this time point (Figure 9G, 9H, arrows). Further support for the temporal shift in bone development came from similar analyses at 6 dpf, when wild types demonstrated levels of GFP and Alizarin red staining that were comparable to those observed in the perichondrium of mutant chondral bones at 4.5 dpf (data not shown). Expression of molecular markers of osteoblasts, such as col10a1 and runx2b, was increased in the perichondrium of mutant chondral bones at 4 dpf, compared to that seen in their wild-type siblings (Figure 9J–9L, arrows, data not shown). In total, these data argue that the cellular basis of excessive bone in fam20b and xylt1 mutants is the precocious differentiation of secretory osteoblasts in mutant perichondria.

Early chondrocyte maturation promotes precocious perichondral bone in PG mutants

Our expression data (Figure 4C, 4F), biochemical data (Figure 5F, Figure S3), and rescue experiments (Figure 3C, 3D) suggest that the premature perichondral bone in fam20b and xylt1 mutants is caused by chondrocyte defects. Indian hedgehog (Ihh) is an osteo-inductive signal expressed by chondrocytes as they mature, and is required for perichondral bone formation [7,8,38]. Therefore, we hypothesized that Ihh and other markers of chondrocyte maturation would appear earlier in fam20b and xylt1 mutants. The transcription factor Runx2 positively regulates chondrocyte maturation through transcriptional activation of such genes as col10a1 and ihh [39–41], so our hypothesis predicted premature up-regulation of runx2 genes and their downstream targets. In support, transcript levels for runx2b were up-regulated in mutant chondrocytes at 3 dpf compared to wild-type chondrocytes (Figure 10A–10C), and col10a1, ihha, and ihhb expression was up-
regulated in mutant chondrocytes earlier than in chondrocytes of wild-type siblings (Figure 10D–10L). Expression of sox9a and col2a1a was high in both xylt1 mutant and wild-type chondrocytes at 3 dpf (data not shown). On the other hand, both transcripts were down-regulated in mature chondrocytes by 6 dpf (data not shown), suggesting that some markers of chondrocyte gene expression were not affected in PG mutants. Importantly, the initiation of chondrocyte differentiation was not accelerated overall in mutants, as Alcian blue staining of pharyngeal cartilages was absent at 48 hpf and present at 60 hpf both in wild types and in PG mutants (Figure S2C). Therefore, the timing of chondrocyte differentiation was accelerated only after mutant cells had begun to immerse themselves in PG-rich extracellular matrix. The relative timing of these events is consistent with our ability to rescue the fam20b mutant phenotype with induction of wild-type fam20b at 55 hpf (see Figure S1).

Given that PG mutant chondrocytes expressed molecular markers of maturation prematurely, we performed ultrastructural analyses to assess whether PG mutant chondrocytes showed early cellular features of chondrocyte hypertrophy. Examined by transmission electron microscopy at 84 hpf, wild-type chondrocytes in the central portion of the ceratohyal contained abundant rough ER, Golgi complexes, and mitochondria, indicative of their high biosynthetic and secretory activity (Figure 11A; [12]). By comparison, xylt1 mutant chondrocytes in the central portion of the ceratohyal at 84 hpf displayed ultrastructural hallmarks of hypertrophic differentiation, including increased cell size, reduction in number of biosynthetic organelles, and cytoplasmic clearing (Figure 11B). Although chondrocytes along almost the entire ceratohyal showed premature hypertrophy in mutants, the most pronounced changes in cellular morphology were in the center of this cartilage element, consistent with the timing and location of molecular changes in gene expression (see Figure 10). Near the end of the xylt1 mutant ceratohyal, we observed chondrocytes that appeared like those in the center of the wild-type ceratohyal (data not shown). Furthermore, we observed differences between wild type and mutants in cartilage ECM ultrastructure. Wild-type cartilage ECM contained well-dispersed matrix proteins within fibrillar collagen, forming a well-defined ECM network (Figure 11C). Both territorial (pericellular) and interterritorial (outer) matrix showed uniform organization. In contrast, xylt1 mutant cartilage ECM was electron dense, tightly packed, and highly fibrillar (Figure 11D). As a result, xylt1 mutant chondrocytes were closer to each other than the cells in WT cartilage (Figure 11B). Furthermore, territorial matrix lateral to xylt1 mutant chondrocytes contained more poorly defined, electron dense precipitates 20–50 nm in diameter than seen in wild-type matrix (* in Figure 11C, 11D). In addition to verifying matrix defects, our ultrastructural analyses demonstrated that xylt1 mutant chondrocytes displayed cellular hallmarks of hypertrophic differentiation prior to those of wild-type siblings.

Given these accelerated molecular and cellular features of chondrocyte maturation in PG mutants, we used molecular and genetic means to demonstrate that chondrocyte Ihh expression signaled prematurely to induce perichondrial bone. Expression of ptc2, a downstream marker of Hh signaling, was increased in ceratohyal perichondrium of fam20b and xylt1 mutants at 3 dpf (Figure 10M–10O), suggesting increased Ihh signaling in PG mutants. No differences in perichondrial ptc1, gli2, and gli3 expression were observed between xylt1 or fam20b mutants and wild types (Figure 10P–10R, data not shown), perhaps reflecting the notion that potential transcriptional targets of Hh signaling are not employed in every cell type [43]. Because suha mutant zebrafish have delayed perichondral ossification [7], we could test
the functional significance of early ihha expression by creating fam20b;ihha and xylt1;ihha double mutants. Perichondral bone was suppressed in fam20b;ihha and xylt1;ihha double mutants (Figure 12A–12H), showing that ihha is required for early perichondral bone in fam20b and xylt1 mutants. While epistatic to the bone phenotypes, the ihha mutation did not alter cartilage matrix reduction of fam20b and xylt1 mutants, consistent with the interpretation that ihha acts downstream of the cartilage matrix defects. In summary, these data argue that cartilage PG defects in fam20b and xylt1 mutants primarily accelerated the timing of chondrocyte maturation and ihha expression, which then secondarily triggered early perichondral bone formation.

**Discussion**

Investigation of the vertebrate skeletal system has revealed substantial insight into the structural and functional roles of proteoglycans (PGs; [14–16,18–20]). Here, we provide experimental data linking functions of long-studied (Xylt1) and recently-identified (Fam20b) members of the PG synthesis pathway to skeletal development in vivo. Xylosyltransferases (Xylts) have been known for 40 years to initiate glycosaminoglycan side chain outgrowth onto protein cores of PGs by transferring xylose to serine residues (Figure 5G; [22,44]). Based on tissue culture and biochemical assays, Xylts were shown to function in the synthesis of xylosyltransferases.
of both heparan sulfate PGs (HSPGs) and chondroitin sulfate PGs (CSPGs; [28,45–47]). Xylt1 expression increased during the course of chondrogenic differentiation in vitro [48], and high Xylt serum activity has been linked to osteoarthritis [49]. Our expression, biochemical, and mutational analyses argue strongly for a predominant role of Xylt1 in CSPG production, specifically in cartilage matrix, thus providing an in vivo context for decades of in vitro studies.

In contrast to Xylts, Fam20 molecules were only recently identified, and our studies reveal in vivo functions of Fam20b [30]. Knock-down and over-expression studies in cell lines indicated that mouse Fam20c could promote odontoblast differentiation from mesenchymal stem cells [50], although the molecular mechanism for this role of Fam20c remains unclear. Renewed focus on Fam20 molecules arose from the revelation that humans with a skeletal disease called Raine syndrome have mutations in FAM20C [26,27]. Subsequently, the paralogous protein Fam20b was shown in vitro to phosphorylate xylose on a nascent glycosaminoglycan side chain (Figure 5G), which appears to increase the likelihood that disaccharide repeats will be added by Exostosins and Chondroitin synthases [23]. Our expression and biochemical data show that fam20b functions similar to xylt1 in cartilage PG production in vivo, and our analyses of xylt1, fam20b, and uxs1 zebrafish mutants highlight the importance of xylosylation in skeletal development. Furthermore, all of these fam20b and xylt1 mutants are homozygous viable, providing the only current vertebrate models in which to study how Fam20 and Xylt molecules affect the variety of human disease-related physiological processes for which they have been implicated [24–27,49]. In particular, we note that humans with Raine syndrome display similar facial dysmorphies and osteosclerosis as seen in fam20b mutant zebrafish, potentiating these fish as an informative model by which to understand the etiology of Raine syndrome.

The mutations in fam20b and xylt1 reported here help identify functional domains of the enzymes these genes encode. While Fam20b domains previously have not been probed experimental-
ly, the conservation of amino acid residues among Fam20 family members suggests functional sites [30]. Both fam20bb1127 and fam20bb1127 disrupt cysteine residues that are highly conserved among vertebrates and may play a role in Fam20b molecular structure. Future in vitro enzyme assays will confirm whether such mutations abolish Fam20b kinase activity. Functional analyses of

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**Figure 7. The uxs1 mutation is epistatic to cartilage and bone phenotypes of fam20b and xylt1 mutants.** A–H, Alcian blue/Alizarin red-stained 6 dpf ceratohyals. Compared to wild types (A,E), cartilage matrix of fam20b14127 (B) and xylt114128 (F) mutants stained less with Alcian blue, but the perichondria of fam20b and xylt1 mutants stained more with Alizarin red (arrows) than in wild types. Similar to uxs1 mutants (C,G), uxs1;fam20b14127 (D) and uxs1;xylt114128 (H) double mutants showed a greater decrease in Alcian blue staining of cartilage matrix than the decrease seen in fam20b and xylt1 mutants; also, Alizarin red staining (arrows) in uxs1 single mutant and uxs1;fam20b and uxs1;xylt1 double mutant perichondria was at wild-type levels. Abbreviations: bsr = branchiostegal ray.

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**Figure 8. fam20b and xylt1 mutants have increased perichondral bone.** At 6 dpf, Alizarin red staining (see Figure 1C–1G) was visible more often in skeletal elements of fam20b14127 (A) and xylt114128 (B) mutants than in wild types. Statistically significant (*, p<0.05) increases, however, were observed in more chondral bones (pq, hm, ch) than dermal bones (ept, mx). Quantitative analyses of the sum of bone areas in left and right ceratohyals (ch(l) and ch(r), dashed outlines) in whole-mount, ventral images of live Alizarin red fluorescence (C–E) revealed statistically significant (*, p<0.05) increases in fam20b14127 (F) and xylt114128 (G) mutants, compared to wild-type siblings at 6 dpf. Abbreviations: A = anterior; ch = ceratohyal; ept = entopterygoid; hm = hyomandibular; l = left; mx = maxilla; pq = palatoquadrate; P = posterior; r = right.

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Xylt1 domains [28,33,51] did not target residues affected in our mutants, so xylt1b1128 and xylt1b1129 will provide new insights into Xylt1 function. The replacement of serine with alanine in xylt1b1128 may alter minimally protein biochemical properties. As we show, this serine residue is conserved in Xylt1’s across vertebrates, suggesting that it might serve as a target for phosphorylation that ultimately impacts enzyme performance. The xylt1b1129 mutation alters some amino acids and then truncates a domain that is associated with protein-protein interactions [88645 superfamily; www.ensembl.org]; thus, this alteration might change substrate recognition or inhibit molecular interactions with other glycosyltransferases in the proposed multi-enzyme complex [22,52]. Future experiments designed to investigate whether skeletal defects of PG synthesis mutants arose merely from quantitative reduction in PG levels or rather also qualitative differences in protein substrate recognition will provide unique insight into the functional roles of PGs in vivo.

Our biochemical analyses demonstrate that the mutations in fam20b and xylt1 reported here do not abolish cartilage PG synthesis completely. Mutations in both fam20b and xylt1 reduce cartilage PG synthesis to similar extents; fam20b;xylt1 double mutants are quantitatively more severe than either mutant alone; and xsl1 mutants, which represent a complete loss of zygotic xylose-dependent PG production [10], exhibit an even more dramatic reduction in cartilage PG synthesis than fam20b;xylt1 double mutants. Therefore, the alleles reported here might not be null, and/or redundant genes partially mask their loss. Five total zebrafish fam20 genes have been reported [fam20a, fam20b, fam20c1, fam20c2, and fam20c3; [30]], and zebrafish have two xylt genes (xylt1, orthologous to human XLYT1, and xylt2, orthologous to human XLYT2; http://useast.ensembl.org/Danio rerio/). Expression and functional analyses of the full set of zebrafish fam20 and xylt genes would test for redundant activities that might compensate for the loss of fam20b and xylt1. We do not expect xylt2 to compensate for a loss of xylt1 function during zebrafish skeletogenesis, because mice deficient for Xylt2 exhibit polycystic kidney disease, but do not have skeletal defects [53].

Our finding that mutations in genes encoding CSPG synthesis enzymes accelerate endochondral ossification reveals that CSPGs can negatively regulate skeletogenic timing. This finding suggests to us that CSPGs might serve as therapeutics for skeletal defects that result from precocious developmental timing, such as craniosynostoses [54,55]. Since various mutants affecting HSPG synthesis do not have increased perichondral bone [14–16], the gain in perichondral bone we describe here must be a CSPG-specific effect. In addition to this qualitative difference between HSPGs and CSPGs, PG levels may be interpreted quantitatively by developing skeletal cells. Homozygous uxs1 mutants demonstrate loss of Alcian blue staining in cartilage in the absence of zygotic CSPGs, but uxs1 mutants have delayed, rather than accelerated, bone formation [17,18]. These data suggest that intermediate levels of CSPGs are required for the unique osteogenic acceleration observed in fam20b and xylt1 mutants. In fact, degradation of CSPGs may be a normal, required stage of endochondral ossification, for lighter Alcian blue staining is associated with more mature cartilage matrix [see Figure 5; [56,57]]. Another explanation for the lack of accelerated bone formation in uxs1 mutants is that these fish fail to produce both CSPGs and HSPGs. The gain in bone formation that loss of CSPGs imparts during endochondral ossification may depend upon HSPG function. This assertion is not supported by our data, however, as fam20b mutants exhibit increased perichondral bone while suffering loss of HSPG production in chondrocytes.

Together, our data argue that defective cartilage PG synthesis alters expression of transcription factors that determine the rates of

Figure 9. Precocious bone formation and osteoblast differentiation in perichondria of fam20b and xylt1 mutants. A–F, Alcian blue/Alizarin red-stained ceratohyals. G–I, live Alizarin red fluorescence of fam20b1127;Tg(sp7:EGFP)b1212 and xylt1b1128;Tg(sp7:EGFP)b1212 larvae. J–O, whole-mount in situ hybridization of 4 dpf ceratohyals. No signs of Alizarin red staining were observed at 3 dpf in ceratohyals of wild types (A), or fam20b1127 (B) or xylt1b1128 (C) mutants. By 4.5 dpf, Alizarin red staining was still absent from ceratohyals of wild-type larvae (D, arrow), although it was detected in ceratohyals of fam20b1127 (E, arrow) and xylt1b1128 (F, arrow) mutants. Both GFP expression and Alizarin red staining were at background levels in the perichondrium of the ceratohyal in 4.5 dpf wild types (G, arrow), but were obvious in the fam20b1127 and xylt1b1128 mutant perichondria (H, arrow). At 4 dpf, col10a1 expression was not detected in the ceratohyal perichondrium of wild types (J, arrow), but was expressed highly in fam20b1127 (K, arrow) and xylt1b1128 (L, arrow) mutant perichondria. Abbreviations: bsr = branchiostegal ray.

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chondrocyte maturation, thus changing the timing of perichondral bone formation. Increased expression of runx2b in fam20b and xylt1 mutant chondrocytes likely drives early expression of chondrocyte maturation markers [39–41], such as ihh co-orthologs, which induce early perichondral bone [7,8,38]. As such, fam20b mutants may provide a completely new etiology for Raine syndrome: defects in chondrocyte differentiation underlie the increased perichondral bone (i.e., osteosclerosis) and skeletal dysmorphies observed in these humans [26,27]. Premature cellular hallmarks of chondrocyte hypertrophy, including cytoplasmic clearing and loss of rough ER, Golgi complex, and mitochondria, accompany molecular features of early chondrocyte maturation in xylt1 mutants. This premature terminal differentiation is also accompanied by changes in cartilage ECM, which contains well-defined, but more tightly packed collagen fibrils, and that might result from failure of GAGs to properly incorporate into the matrix.

The general hypothesis emerging from our work that CSPGs negatively regulate chondrocyte maturation is consistent with a recent, detailed study of the chick Aggrecan mutant [21]. Future work will aim to decipher the mechanism by which mutations in the cartilage PG synthesis pathway impact the timing of chondrocyte differentiation. Recent studies, for example, highlight a novel role for CSPGs in modulating growth factor signaling in developing cartilage [21,58]. Here, we demonstrate that mutations in the CSPG synthesis pathway can accelerate developmental timing, thus expanding the ontology of genes regulating the rate of skeletogenesis.

Materials and Methods
Zebrafish lines
All fish lines were maintained and embryos raised according to established protocols [59] with IACUC approval. We obtained the b1125, b1127, b1128, and b1189 mutant alleles through mutagenesis with N-ethyl-N-nitrosourea (ENU) in an AB background [59]. ihhaahu2131 fish were obtained from P. Ingham.

Histological stains
Embryos were fixed in 2% PFA in PBS for 1 hr., washed in 100 mM Tris pH 7.5/10 mM MgCl2 for 10 min., stained in 0.04% Alcian blue/10 mM MgCl2/70% EtOH pH 7.5 overnight, taken through graded EtOH series (80% EtOH/100 mM Tris

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Figure 10. Early molecular markers of chondrocyte maturation in fam20b and xylt1 mutants. A–R, horizontal section in situ hybridization of developing ceratohyals. The levels of runx2b transcripts appeared higher in chondrocytes (c) and perichondrium (pc) of the ceratohyal in fam20bmutants (B) and xylt1mutants (C) mutants, than in wild types (A) at 72 hpf. Expression of col10a1 was abundant in chondrocytes (c) and perichondrium (pc) of the ceratohyal in fam20bmutants (E) and xylt1mutants (F) mutants, but was not detectable in wild types (D) at 83 hpf. Levels of ihha and ihhb transcripts were up-regulated in chondrocytes (c) of the ceratohyal in fam20bmutants (H,K) and xylt1mutants (I,L) mutants, but were not detectable in wild types (G,J) at 72 hpf. Transcript levels for ptch2 were increased in the perichondrium of fam20bmutants (N) and xylt1mutants (O) mutants, compared to wild types (M) at 72 hpf, whereas no obvious differences in levels of ptch1 expression in perichondria were apparent (P–R). Abbreviations: c = chondrocytes; pc = perichondrium.

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pH 7.5/10 mM MgCl2; 50% EtOH/100 mM Tris pH 7.5; 25% EtOH/100 mM Tris pH 7.5), bleached in 3% H2O2/0.5% KOH for 10 min. with lids open, washed twice in 25% glycerol/0.1% KOH for 10 min. each, stained in 0.01% Alizarin red/25% glycerol/0.1% KOH pH 7.5 for 30 min., and de-stained with two washes of 50% glycerol/0.1% KOH. Larvae were also incubated in 0.003% Alizarin red in Embryo Medium to visualize live mineralized bone.

Figure 11. Ultrastructural evidence of premature chondrocyte hypertrophy and aberrant matrix production in xylt1 mutants. A–D, transmission electron micrographs of 84 hpf ceratohyals. At 5600X magnification, chondrocytes in the central region of the wild-type ceratohyal displayed abundant rough ER, Golgi, and mitochondria in the cytoplasm (A). Similar views of xylt1 mutant chondrocytes at the same age and comparable positions within the developing ceratohyal showed cytoplasmic clearing and tremendous reduction in biosynthetic organelles (B). Mutant chondrocytes also appeared to be separated by less extracellular matrix. While wild-type (C) and mutant (D) extracellular matrix demonstrated fibrillar collagens at 31000X magnification, mutant matrix showed tighter fibril packing and also contained more amorphous electron-dense structures (*) in pericellular matrix than seen in wild types. Abbreviations: cyt = cytoplasm; ecm = extracellular matrix; nuc = nucleus. Scale bars: A,B = 2 μm; C,D = 500 nm.

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Figure 12. The ihha mutation is epistatic to bone, but not cartilage, phenotypes of fam20b and xylt1 mutants. A–H, Alcian blue/Alizarin red-stained 6 dpf ceratohyals. Alcian blue staining was comparable between wild-type (A,E) and ihha mutant (B,F) cartilages, but was decreased in fam20b<sup>b1127</sup> (C) and xylt1<sup>b1128</sup> (G) single mutant, and in fam20b<sup>b1127</sup>;ihha (D) and xylt1<sup>b1128</sup>;ihha (H) double mutant, cartilages. Alizarin red staining of fam20b<sup>b1127</sup> (C) and xylt1<sup>b1128</sup> (D) chondral bones was abundant, while no such staining was observed in wild types (A,E), ihha mutants (B,F), or fam20b<sup>b1127</sup>;ihha (D) and xylt1<sup>b1128</sup>;ihha double mutants (H). Abbreviations: bsr = branchiostegal ray.

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Mapping, cloning, and genotyping PG mutants

All oligonucleotide sequences appear in Figure S4. Sequence alignments were made using MultiAlin [60]. RAD mapping localized b1127 to LG20 [29]. Subsequent simple sequence repeat (SSR) mapping identified a genetic interval of 0.2 cM between SSRs defined by primers A+B and C+D on scaffold 2914 of Zv7 (http://uswest.ensembl.org/Danio_rerio/Info/Index) containing the b1127 mutation (Figure 2A). In this interval, the SSR marker z20582 showed zero cross-over with the mutant phenotype in 862 meioses. RNA isolated (TRI Reagent; Ambion Inc.) from live 5 dpf Alizarin red-stained zebrafish larvae that were screened for increased perichondral bone were made into cDNAs (First Strand Synthesis kit; Invitrogen Corp.). Full sequence of fam20b cDNA was determined by overlapping PCR fragments, generated from primers I+J and K+L. Four genes (angtl1, salgs2, blal1, and fam20b) around z20582 were sequenced, and of these, mutant-specific coding sequence changes were identified only in the sixth coding exon of fam20b (Figure 2B, data not shown). PCR-based genotyping assays showed perfect correspondence between the fam20b(1125) skeletal phenotype and this mutation (0 cross-overs/614 meioses; Figure 2B). The second allele in the fam20b complementation group, b1125, also mapped tightly to fam20b. The map cross contained three cross-overs between fam20b(1125) and z20582 among 602 meioses (0.5 cM). b1127 was genotyped by digesting PCR product from primers V+W with HaeIII, which only cuts mutant sequence. b1125 was genotyped with the marker z10805.

RAD mapping localized b1128 to LG3 [29]. Subsequent SSR mapping defined a genetic interval of 0.7 cM between SSRs defined by primers E+F and G+H on scaffold 380 of Zv7 containing the mutation (Figure 2D). We focused on a predicted gene (LOC560951; www.ensembl.org) within this interval with homology to xylosyltransferase1 (xylt1). Comparison of the Ensembl-predicted protein to Xylt1 of other vertebrates suggested that the annotated version of Xylt1 for zebrafish was lacking about 50 amino acids at the N-terminal portion of the protein. Therefore, we used 5’RACE on a 3 dpf cDNA library, using the nested primers M and N, along with universal primers (Clontech Laboratories, Inc.), to reveal an unannotated xylt1 exon in zebrafish over 50 kb upstream of the annotated version. PCR and sequencing analyses confirmed the gene structure of zebrafish xylt1, which consists of 11 exons, similar to human XYLTI (Figure 2E). Full sequence of xylt1 cDNA generated from primers O+N, P+Q, R+S, and T+U was submitted to GenBank (Accession: HQ692884). Sequencing of xylt1 from b1128 cDNA and gDNA revealed a splice donor mutation (G2103A) in exon 9 (Figure 2E). PCR-based genotype assays showed perfect correspondence between the xylt1(1128) phenotype and this splice site mutation (0 cross-overs/738 meioses; Figure 2E). Sequencing from 5 dpf xylt1(1128) mutant cDNAs showed that in the absence of the wild-type splice donor site, cryptic sites were used (6/6 clones). Although the most common of these (5/6 clones) resulted in a tetranucleotide insertion (Figure 2E, 2F), all mutant cDNAs would produce a frameshifted and truncated C-terminal portion of the zebrafish Xylt1 protein from at least amino acid 702 (out of 919). The second allele in the xylt1 complementation group, b1189, also had a unique mutation in xylt1. cDNA sequencing revealed a T1600G mutation in the seventh coding exon of xylt1(1129), which altered highly-conserved amino acid 534 from Ser to Ala (Figure 2E, 2F). PCR-based genotype assays showed perfect correspondence between this mutation and the xylt1(1129) phenotype (0 cross-overs/632 meioses; Figure 2E). b1128 was genotyped by digesting PCR product from primers X+Y with Ddel, which only cuts wild-type sequence. b1189 was genotyped by digesting PCR product from primers Z+AA with Hhal, which only cuts mutant sequence.

fam20b rescue

Full-length fam20b was amplified from mutant and wild-type cDNAs with primers BB+CC and inserted into pDONR211 by BP recombination (Invitrogen Corp.). LR recombination reactions inserted fam20b under the control of beta-actin2 or hsp70 promoters with GFP fused in-frame in the destination vector pDestTo2CG2, which also contained cmvl2-GFP [32]. mRNA from pcS2FA-transposase (http://chien.neuro.utah.edu/tol2kitwiki) was made (mMessage mMMachine; Ambion Inc.). Approximately 3 nl of 100 ng/μl of fam20b expression plasmid, 70 ng/μl transposase RNA, and 0.2% phenol red were injected into one to four-cell embryos. Embryos containing green hearts were screened, and for heat shock activation, 55 hpf embryos were incubated at 40°C for 20 min.

Expression analyses

Whole-mount and section RNA in situ hybridization were carried out as described [61,62]. Probes used were runx2b, ibha, ihhh, ptc1, ptc2, col10a1 [18]. Nomenclature for ptc genes reflects a recent update, whereby previously termed ptc1 is now ptc2 and ptc2 is now ptc1 (www.zfin.org). Probes for fam20b and xylt1 were created by cloning PCR amplicons from primers K+L and R+S, respectively, into pCR4-TOPO (Invitrogen Corp), digesting with NotI, and transcribing with T3 Polymerase. Primers DD+L, and P+EE were used for fam20b and xylt1 RT-PCR analyses, respectively. Immunostaining was carried out as previously reported [18].

Alcian blue quantitation

Since Alcian blue binds sulfated glycosaminoglycans (GAGs; [63]), defects in cartilage PG production should be reflected by Alcian blue levels. As an indication of the levels of sulfated glycosaminoglycans in cell and tissue lysates, we modified the protocol of Bjornsson et al. (1998) as follows: larvae underwent all the stages except the Alizarin red staining of the Alcian blue, Alizarin red procedure outlined above, and tails were removed for genotyping. Samples were lysed (DEasy kit; QIAGEN Inc.), spun briefly, and read spectrophotometrically at 620 nm. All analyses were performed in triplicate. Error bars represent standard error of the means.

HPLC analyses

Glycosaminoglycan (GAG) biochemistry was carried out by the UCSD Glycotechnology Core Resource, according to protocols Pre006, Mis002, Mod004, Pro006, and Pro007 (http://glycotech.ucsd.edu/seq.html). Some disaccharide species did not appear in the majority of samples, due to their overall low levels, and were not included in analyses.

Quantitation of perichondral bone

Digital images of flat-mounted, ventral views of cartilaginous in live Alizarin red-stained 6 dpf larvae were obtained under equivalent microscope settings, and subsequently analyzed with ImageJ (http://rsbweb.nih.gov/ij/). Standard threshold values were applied to each image to define mineralized portions of cartilaginous perichondria. From eight samples each for genotyped mutants and wild types, mineralized area from the two cartilaginous was determined by the Analyze Particles function. Error bars represent standard error of the means.
Proteoglycans Regulate Skeletal Timing

OPT imaging

Fish were fixed overnight in 4% PFA, washed for an hour in 1% KOH, bleached in 3% H2O2/0.5% KOH for 40 min, with lids open, washed in 1% KOH, stained overnight in 0.003% Alizarin red in 1% KOH, and de-stained in 1% KOH. After eyes were removed, heads were embedded in agarose, washed twice in methanol, and cleared in benzyl alcohol/benzyl benzoate (2:1). Images were captured using Bioptics OPT Scanner 3001 M (MRC Technology).

Transmission electron microscopy

Samples were processed as previously described [42], with the following modification. Zebrafish embryos were fixed in 2.5% glutaraldehyde in 0.1 M sodium cacodylate at room temperature for 1 hr and then at 4°C for at least 24 hr. They were then rinsed in 0.1 M sodium cacodylate and post-fixed with 1% osmium tetroxide in 0.1 M sodium cacodylate for 1 hr. Following additional rinsing, specimens were dehydrated stepwise in ethanol and then propylene oxide, infiltrated with resin step-wise, and then embedded in resin for 48 hr at 60°C. Ultra-thin 50 nm sections were collected on a Leica Ultracut Microtome and analyzed on a Phillips CM-12 Transmission Electron Microscope provided by VUMC Cell Imaging Shared Resource.

Supporting Information

Figure S1 Exogenous wild-type fam20b expression does not alter skeletal phenotypes of wild types or yxl+/+, but can rescue cartilage of fam20b++ mutants with a single induction at 55 hpf. A–E, dissected, flat-mounted pharyngeal skeletons of Alcian blue/Alizarin red-stained (A,B) and Alcian blue-stained (C–E) larvae. Injection of larvae with wild-type fam20b under control of the beta-actin2 promoter did not alter skeletal phenotypes of wild types (A) or yxl+/+ mutants (B). Injection of larvae with wild-type fam20b under control of the hsp70 promoter rescued cartilage matrix production in fam20b++ mutants when heat shocked at 55 hpf (E), compared to injected fam20b++ embryos that did not undergo heat shock (D), while heat shock had no effect on un.injected control larvae (C). Abbreviations: HS = heat shock; UIC = uninjected control. (TIF)

Figure S2 Cartilage proteoglycan secretion initiates at the same time in yxl mutants and wild types. Alcian blue matrix was not apparent in pharyngeal arch 1 or 2 of wild-type larvae (A). Injection of yxl++ mutants (B) at 48 hpf. Staining in the trabeculae of the neurocranium is evident in both wild types and yxl++ mutants. Alcian blue staining was obvious in pharyngeal arches of both wild types (C) and yxl++ mutants (D) at 60 hpf. Abbreviations: ch = ceratohyal; pa1 = pharyngeal arch 1; pa2 = pharyngeal arch 2; pq = palatoquadrate; t = trabeculae. (TIF)

Figure S3 Loss of proteoglycans in fam20b, yxl, and uxl mutants. A–C, HPLC quantitation of disaccharides in proteoglycans from whole larval lysates of fam20b++, yxl++ and uxl mutants at 5 dpf. A, Chondroitin sulfate levels were reduced in all mutants compared to wild-type siblings. B, Heparan sulfate levels were unaffected largely in yxl mutants, but were decreased in fam20b and uxl mutants, compared to wild-type siblings. C, Instead of plotting values as a percentage of wild-type, this graph illustrates picomoles of each disaccharide species in fam20b++, yxl++ and their wild-type siblings, demonstrating that the relatively high level of UA-2S-GlcNAc in fam20b++ mutants (* in B) is due to the low levels of this disaccharide that typically were detectable in these samples (* in C). (TIF)

Figure S4 Oligonucleotide sequences used in this study. (DOC)

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Author Contributions

Conceived and designed the experiments: BFE EWS JHP CBK. Performed the experiments: BFE DSL Y-LY MES. Analyzed the data: BFE Y-LY MES DSL EWS JHP CBK. Contributed reagents/materials/analysis tools: BFE Y-LY MES EWS CBK. Wrote the paper: BFE.

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