The type I collagenopathies are a group of heterogeneous connective tissue disorders, that are caused by mutations in the genes encoding type I collagen and include specific forms of osteogenesis imperfecta (OI) and the Ehlers-Danlos syndrome (EDS). These disorders present with a broad disease spectrum and large clinical variability of which the underlying genetic basis is still poorly understood. In this study, we systematically analyzed skeletal phenotypes in a large set of type I collagen mutants, representing different genetic forms of human OI, and a zebrafish model resembling human EDS, which harbors a number of soft connective tissue defects, typical of EDS. Furthermore, we provide insight into how zebrafish and human type I collagen are compositionally and functionally related, which is relevant in the interpretation of human type I collagen-related disease models. Our studies reveal a high degree of intergenotype variability in phenotypic expressivity that closely correlates with associated OI severity. Furthermore, we demonstrate the potential for select mutations to give rise to phenotypic variability, mirroring the clinical variability associated with human disease pathology. Therefore, our work suggests the future potential for zebrafish to aid in identifying unknown genetic modifiers and mechanisms underlying the phenotypic variability in OI and related disorders. This will improve diagnostic strategies and enable the discovery of new targetable pathways for pharmacological intervention.

Significance

Type I collagenopathies are a heterogenous group of connective tissue disorders, caused by genetic defects in type I collagen. Inherent to these disorders is a large clinical variability, of which the underlying molecular basis remains undefined. By systematically analyzing skeletal phenotypes in a large set of type I collagen zebrafish mutants, we show that zebrafish models are able to both genocopy and phenocopy different forms of human type I collagenopathies, arguing for a similar pathogenetic basis. This study illustrates the future potential of zebrafish as a tool to further dissect the molecular basis of phenotypic variability in human type I collagenopathies, to improve diagnostic strategies as well as promote the discovery of new targetable pathways for pharmacological intervention of these disorders.
residue for a bulky, polar, or charged amino acid, disrupting the highly conserved Gly-X-Y triplet sequence. Following the Silence et al. (4) classification, type IV is described as being moderate, type III as severe or progressively deforming, and type II as causing perinatal lethality. Besides the classic types of OI, mutations in noncollagenous genes have been reported to mostly cause autosomal recessive forms of OI. These genes generally encode for proteins that interact with type I collagen, acting as key players in processes such as collagen synthesis, collagen folding and post-translational modification, intracellular trafficking of collagen, or collagen fibril cross-linking (5).

Inherent to the pathology of OI is a large clinical variability, with phenotypes ranging from nearly asymptomatic with a mild predisposition to fractures and a normal lifespan, to forms associated with multiple fractures and severe deformities already present in utero that may cause perinatal lethality (7). In the classic dominant forms of OI types II–IV, some general principles have emerged for genotype–phenotype correlations (3, 8). Mutations in COL1A1 are generally associated with a more severe phenotype than mutations in COL1A2. Furthermore, the position of mutations along the α-chains can modulate the outcome. Nevertheless, numerous exceptions to these guidelines have already been demonstrated (9). Of note, extensive phenotypic variability resulting from identical mutations has been described in recent years to be common in both dominant and recessive forms of OI (10–12). This renders prediction on the clinical outcome of certain mutations in OI particularly challenging. Dissecting the underlying basis of this phenotypic variability is crucial to enable a more profound understanding of molecular mechanisms in OI, to enable the discovery of new targetable pathways for intervention, and to identify novel biomarkers for diagnostics and monitoring of pharmacological treatments.

To study human skeletal dysplasias, zebrafish (Danio rerio) models are increasingly being used as a valuable complementation or predecessors to the traditional murine models (13, 14). Besides their unique attributes, such as the rapid development, large offspring numbers, and ease and speed in generating mutant lines, zebrafish bone mutants tend to survive into adulthood far more easily than corresponding mouse models, making them also available for the study of later stages of skeletal development and maturation (13, 15). Furthermore, due to the high conservation of key developmental programs in osteogenesis between teleosts and mammals, functional gene and pathway analysis in zebrafish can yield relevant insights into human bone disease (16). Detailed reports have already documented similarities and differences between teleost and mammalian bone biology that are relevant for modeling human bone disease (13, 14, 17, 18). One of these differences is the composition of type I collagen in zebrafish, which harbors not one but two orthologs of the human COL1A1 gene, namely col1ala and col1alb, encoding the α1- and α3-chain, respectively. Although some characteristics of zebrafish type I collagen have already been addressed (19), more insight into the functional implications of this additional teleost-specific α-chain is needed.

In recent years, different zebrafish mutants have already been reported to accurately model certain genetic types of human OI (20–23). However, these studies were focused on a detailed analysis of single mutants, modeling certain subtypes of OI, while the greatest strength of the zebrafish lies in the ease of accommodating the parallel analysis of multiple mutant models. Moreover, recent advances in μCT-based methods enable detailed and rapid skeletal phenotyping of zebrafish mutants, which often have been resistant to such methodologies (24). Advances in processing and analysis have now allowed analysis of hundreds of morphological and densitometric traits in large sets of zebrafish skeletal mutants (25). In this work, we applied systematic collagen analysis with skeletal phenomics to characterize a large set of zebrafish with mutations in type I collagen genes, as seen in patients with different forms of classical OI and EDS, to assess to which extent key features of human type I collagenopathies are recapitulated.

**Results**

**Zebrafish with Different Mutations in Type I Collagen Show Variable Skeletal Phenotypes.** We systematically analyzed skeletal phenotypes in a large set of zebrafish models carrying different mutations in the zebrafish type I collagen-encoding genes col1ala, col1alb, and col1a2 (Table 1). As a reference, we also included two knockout mutants, bmp1a−/− and plod2−/−, representing severe recessive forms of OI with defects in type I collagen processing and cross-linking, respectively (22). Upon μCT scanning, we observed a large diversity of skeletal phenotypes throughout the whole set of mutants. Severe morphological abnormalities included callus formation, bowing and kinking of the ribs, malformation of the vertebral column, short stature, and craniofacial abnormalities (Fig. 1, Table 2, and SI Appendix, Fig. S1), indicating compromised bone quality. Next, we performed μCT-based phenomic profiling (25) (SI Appendix, Fig. S2) to quantify 200 different descriptors of bone morphology and mineralization in the axial skeleton of each animal. Specifically, we segmented each of 20 vertebrae into three skeletal elements—the neural arch (Neur), centrum (Cent), and haemal arch/ribs (Haem)—and for each skeletal element, we computed four primary parameters: tissue mineral density (TMD, mg/cm3), bone volume (Vol, μm3), and thickness (Th, μm), and also centroid length (Cent.L, μm). In total, we analyzed 28,000 phenotypic data points derived from 140 different animals. To explore inter-genotype variability for each trait, means were calculated within each genotype (mutant and control) and normalized to the SD of their respective control population [z-score = (mean value − mean value control)/SD control]. In general, we observed a large diversity of skeletal phenotypes throughout the whole set of mutants (Fig. 2). We found that phenotypic severity (as measured by the absolute value of the z-score) tended to be highest for mutants associated with collagen processing (plod2, bmp1a) and qualitative collagen defects (e.g., col1ala−/−, col1alaβ1−/−, col1alaβ2/−+), and lower for mutants associated with quantitative collagen defects (e.g., col1alas−/−, col1albs−/−, col1a2−/−). Remarkably, some mutants with qualitative defects (and collagen processing defects) exhibited a pronounced enrichment of positive z-scores for some traits (e.g., Cent. TMD). We interpret the different degrees of phenotypic severity in mutants associated with qualitative or quantitative collagen defect, as well as the variability in direction of effect (i.e., positive or negative effect on bone mass or mineralization) across genotypes, to reflect the clinical variability observed in human type I collagenopathies.

**Heterozygous Loss of both α1(I) and α3(I) in Zebrafish Is Remiscent of Mild OI Type I.** Human patients with a heterozygous COL1A1-null allele (COL1A1 haploinsufficiency) develop OI type I, characterized by short or normal stature, mild bone fragility, relatively few fractures, and minimal limb deformities. In zebrafish two orthologs of the human COL1A1 exist, namely col1ala encoding α1(I) and col1alb encoding α3(I). Two zebrafish knockout mutants with a premature stop-codon mutation in either col1ala or col1alb show absence of α1(I) and α3(I), respectively, in the vertebral bone (Fig. 3A and SI Appendix, Fig. S3). However, both qualitative and quantitative assessment of μCT scans of the vertebral column of heterozygous col1alaα1−/− or col1albα3−/− zebrafish mutants revealed no skeletal abnormalities (Fig. 1 and SI Appendix, Figs. S4 and S5), which is most likely related to functional redundancy between both paralogs. Hence, we generated a double-heterozygous knockout mutant (col1alaα1−/−col1albα3−/−), which displays a mild skeletal phenotype, with a low frequency of spontaneous fractures (calluses
in the ribs), scoliosis and localized compression, fusions, and mild malformation of the vertebral bodies in some of the mutant fish (Figs. 1 and 4 and Table 2). Quantitative measures of bone in this mutant revealed localized reduction of centrum volume, length, and TMD (SI Appendix, Fig. S6). However, these results were not found to be statistically significant throughout the entire

Table 1. List of mutant zebrafish alleles analyzed in this study

| Gene     | Allele  | Effect on protein level | Effect on homologous human protein |
|----------|---------|-------------------------|------------------------------------|
| **Type I collagen knockout alleles (quantitative defect)** |           |                         |                                    |
| col1a1a  | sa1748  | p.(Gly1179X)             | p.(Gly736Asp)                      |
| col1a1b  | sa12931 | p.(Cys68X)               | p.(Gly288Y)                        |
| col1a2   | sa17981 | p.(Ala154Cysfs*23)       |                                    |
| **Type I collagen amino acid substitutions (qualitative defect)** |           |                         |                                    |
| col1a1a  | chihuahua (chi) | p.(Gly736Asp) | α1(I) p.(Gly574Asp) |
| col1a1a  | microwaved (med) | p.(Gly888Y)ys | α1(I) p.(Gly726Arg) |
| col1a1a  | dmh13  | p.(Gly1093A)             | α1(I) p.(Gly931A) |
| col1a1a  | dmh14  | p.(Gly1144A)             | α1(I) p.(Gly981A) |
| col1a1b  | dmh29  | p.(Gly1123A)             | α1(I) p.(Gly958A) |
| col1a2   | dmh15  | p.(Gly882A)              | α2(I) p.(Gly802A) |
| **Knockout alleles in genes involved in Type I collagen processing** |           |                         |                                    |
| bmp1a    | sa2416  | p.(Arg522X)              |                                    |
| plod2    | sa1768  | p.(Tyr679X)              |                                    |

Mutant alleles are listed, along with the effect of the mutation on protein level (p-notation). For alleles causing an amino acid substitution, the corresponding effect on the homologous human protein is listed in the p-notation (35).

![Fig. 1. μCT images of different zebrafish models with affected type I collagen. For μCT scanning, we included several mutants with a nonsense or splice mutation in col1a1a, col1a1b, or col1a2, generating a knockout of these genes (col1a1a−/− not viable). These mutants are grouped together and indicated as “type I collagen knockout (KO) alleles—quantitative defect” (shown in orange). Another set of mutants carries mutations resulting in substitutions (indicated as “type I collagen amino acid (AA) substitutions—qualitative defect, shown in blue) of a Gly residue in α1(I) (col1a1a dmh13/+, col1a1a dmh14/+, col1a1b dmh29/+), α2(I) (col1a2 dmh15/+, or α3(I) (col1a1b dmh29/+)”. The microwaved (col1a1a dmh13/med/med) mutant carries a homozygous Glu substitution in α1(I). Finally, we included two mutant models with a knockout mutation in the bmp1a and plod2 genes (“KO alleles in genes involved in type I collagen processing,” shown in green). Representative fish from each mutant genotype are shown. Callus formation in ribs (arrowheads), local compressions of the vertebral column (brackets), and kyphosis (arrow), are indicated (also listed in Table 2).
length of the vertebral column, most likely due to the intragenotype phenotypic variability and incomplete penetrance, which is further demonstrated by the much larger spread of values for the different skeletal parameters in the group of mutant fish compared with control fish (Fig. 4B).

A Complete Loss of Zebrafish α1(I), but Not α3(I), Causes Lethality in Early Larval Stages. We next assessed the effect of a complete (homozygous) loss of α1(I) or α3(I) on survival and on skeletal integrity. In progeny resulting from an in-cross of col1a1a−/−; col1a1b−/− mutant fish, all possible genotypes were shown to be present at 7 dpf postfertilization (dpf), while from 15 dpf on some genotypes were underrepresented according to Mendelian predictions, or even lost (Fig. S4). Eventually, all genotypes containing a homozygous knockout of col1a1a [loss of α1(I)] were found to be lethal by the age of 3 mo. Homozygous mutant col1a1b−/− mutants [loss of α3(I)] were present by the age of 3 mo, however at reduced numbers if combined with heterozygous loss of col1a1a (col1a1a−/−; col1a1b−/−). Fish with a complete loss of α3(I), but intact α1(I) did not show significant alterations of bone morphology (SI Appendix, Fig. S7), although some fish showed evidence of fractures (Fig. 1 and Table 2) and increased TMD throughout the vertebral column, indicating an impaired bone quality.

Because the knockout of col1a1a compromises viability from 7 dpf on, the general morphology of col1a1a−/− mutant larvae was assessed at this stage. These larvae lacked the presence of an inflated swim bladder, a structure that is essential for survival once larvae are required to actively feed (Fig. 5B). Furthermore, the distal margins of the pectoral fins, and the finfold, were shown to be ruffled in these mutants (Fig. 5 B and C). These fin structures are structurally supported by fin actinotrichia, which are prefin-ray structures composed of collagen fibers (26). Upon detailed examination of the finfold, these actinotrichia were shown to contain type I collagen only. Type II collagen is associated with the cardiac valvular type of EDS, a known to be pathogenic, while a complete absence of COL1A2 is associated with the cardiac valvular type of EDS, a connective tissue disorder characterized by skin fragility, joint hypermobility, early-onset, severe and progressive cardiac valvular defects, and mild osteopenia with increased risk of fractures in some patients (6, 29).

Complete Loss of α2(I) in Zebrafish Leads to Soft Connective Tissues Abnormalities, Reminiscent of the Human EDS. We studied a zebrafish mutant with a splice mutation in col1a2, resulting in the absence of α2(I) (Fig. 3B). In humans, heterozygosity for COLLA2-null mutations (COLLA2 haploinsufficiency) is not known to be pathogenic, while a complete absence of α2(I) chains is associated with the cardiac valvular type of EDS, a connective tissue disorder characterized by skin fragility, joint hypermobility, early-onset, severe and progressive cardiac valvular defects, and mild osteopenia with increased risk of fractures in some patients (6, 29).

Table 2. Phenotypical features observed upon μCT-scanning of adult mutant fish included in this study

| Zebrafish mutant | Associated human disorder + unique clinical features | No. of animals with rib fracture callus | No. of vertebral fusions | VC | No. of animals over/undermineralized (TMD, P < 0.05) | Kyphosis (K) and/or scoliosis (S) | Decreased bodylength–SL (P < 0.0001) |
|------------------|-----------------------------------------------------|----------------------------------------|----------------------------|---|-------------------------------------------------|-----------------------------|---------------------------------|
| Type I collagen knockout alleles (quantitative defect) | | | | | | | |
| col1a1a−/−       | OI type I, Mild. Normal                             | 0/5                                    | 0/5                        | / | /                                               | /                          | /                               |
| col1a1b−/−       | or short stature, little                           | 0/5                                    | 0/5                        | / | /                                               | /                          | /                               |
| col1a1α−/--;col1a1b−/− | or no deformities                                   | 1/5                                    | 2/5                        | / | S                                               | /                          | /                               |
| col1a1b−/−       | 2/5                                                 | 0/5                                    | /                          | / | /                                               | /                          | /                               |
| col1a2−/−       | No disease                                         | 0/5                                    | 0/5                        | / | /                                               | /                          | /                               |
| col1a2−/−       | EDS, cardiac valvular type                          | 0/5                                    | 0/5                        | / | K                                               | Y                          | Y                               |
| Type I collagen amino acid substitutions (qualitative defect) | | | | | | | |
| col1a1aH10R      | OI types II–IV, moderate                           | 5/5                                    | 5/5                        | / | /                                               | K + S                      | Y                               |
| col1a1aMED       | to severe/lethal.                                   | 1/6                                    | 0/6                        | Undermineralized | S | /                                               | /                          | /                               |
| col1a1a(nmnl)12αR| Short stature and bone deformities.                 | 1/6                                    | 4/6                        | / | S                                               | Y                          | Y                               |
| col1a1a(nmnl)14αR| Overmineralization                                 | 0/5                                    | 1/5                        | / | /                                               | /                          | /                               |
| col1a2(nmnl)28αR| (36)                                                | 5/6                                    | 6/6                        | Overmineralized | K + S | Y                                               | Y                          | Y                               |
| Knockout alleles in genes involved in type I collagen processing | | | | | | | |
| bmp1a−/−        | OI type XII, severe. Marked overmineralization     | 0/5                                    | 0/5                        | Overmineralized | K + S | Y                                               | Y                          | Y                               |
| pld2−/−         | Overlap B5–O1 moderate/severe                      | 5/5                                    | 5/5                        | Overmineralized | K + S | Y                                               | Y                          | Y                               |

For each mutant model, the specific disorder and unique clinical features associated with a similar mutation in human patients is listed (7, 20). In addition, per genotype the ratio (no.) of animals that had at least one rib fracture callus (third column, never observed in any wild-type control animals), or fusion of vertebral bodies in the vertebral column (fourth column, only rarely detected in wild-type control animals) are given. TMD in the VC was quantified in FishCuT software and was indicated as “undermineralized” if significantly decreased or as “overmineralized” if significantly increased according to the global test (P < 0.005). The presence of kyphosis (K) and or scoliosis (S) was also evaluated on 3D-scan images. Finally, the presence of a significant decrease (P < 0.001) in the mean body length, [i.e., the standard length (SL, measured from the tip of the snout to basis of the caudal fin)] of mutant genotypes is indicated (see SI Appendix, Fig. S20 for boxplots of SL measurements per genotype, compared with their wild-type siblings). Other abbreviations: B5, Bruck syndrome; VC, vertebral column.
Given the joint involvement in human patients with a complete loss of α2(I) (6), we explored the presence of similar defects in col1a2−/−mutant fish. Histological analysis of the intervertebral ligament (IVL) showed a normal elastin layer, but reduced notochord sheath layer and a complete loss of type I collagen fibrous ligament, causing local distortion and dislocation of the IVL and intervertebral space in some parts of the vertebral column (Fig. 6D), which most likely underlies the kyphosis identified in this mutant and resembling the joint dislocations seen in human patients (6, 30, 31). The skin of col1a2−/−mutant fish was also more fragile and easily damaged, compared with the skin of wild-type control fish. Histological analysis of adult skin showed that the dermis, which is composed of two collagen fibril layers in zebrafish, is half the thickness in mutant fish compared with dermis of wild-type siblings (Fig. 6B). Biomechanical load-to-failure strength of soft connective tissues was analyzed by determining the ultimate tensile strength of tissue specimens from five col1a2−/−mutant and five control fish (Fig. 6C). Tissues lacking α2(I) ruptured at a significantly lower peak load (0.027 ± 0.0079 N), compared with tissue samples from wild-type siblings (0.081 ± 0.011 N, P < 0.0001), indicating significantly diminished strength of soft connective tissue in col1a2−/−mutant fish. Histological sections of the adult zebrafish heart from col1a2−/−mutants showed normal morphological appearances of the cardiac valves and col1a2−/−larvae displayed normal cardiac function and blood flow (SI Appendix, Fig. S10).

Genotype/Phenotype Relations in Mutants with Impaired Quality of Type I Collagen. We next studied the skeleton of a set of mutants carrying a point mutation in either of the type I collagen-encoding genes, leading to impaired type I collagen (Fig. 1, Table 1, and SI Appendix, Figs. S11–S17). In human patients these types of mutations lead to OI types II–IV, which have moderate to severe or even lethal phenotypes with short stature, bone deformities, and hypermineralized bone (32, 33).

The col1a2dmh15/+ mutant represents the most detrimental phenotype of all analyzed mutants (Fig. 1 and SI Appendix, Fig. S16), with heavily distorted, misshapen, and overmineralized axial and cranial skeletons. This is in concordance with human data because Gly substitutions in the homologous region of α2(I) are associated with lethal OI in human patients due to the presence of severe skeletal malformations early in fetal life.

Fig. 2. Inter- and intragenotype phenotypic variability demonstrated by quantitative μCT analysis in FishCuT. For a set of parameters that quantify morphology and mineralization in the vertebral column of the different mutant populations, z-scores are given for each mutant genotype [z-score = (mean value mutant − mean value control)/SD control]. On the y axis, mutants associated with a collagen processing defect are indicated in green, mutants with a qualitative defect in type I collagen are indicated in dark blue, and mutants with a quantitative defect in type I collagen are indicated in orange. Blue bars indicate values that were statistically significantly altered for that mutant genotype, compared with its control as analyzed by the global test (see SI Appendix, Figs. S4–S19). Note the large variability in the effect of each mutation on the different parameters.
development, in addition to other extraskeletal anomalies (Fig. 7) (34). The \textit{collala}^{chi/+} mutant, reported earlier (21, 35), carries the same mutation that was identified in a human patient with severe and heavily deforming OI type III (34), and also displays a moderate to severe phenotype with skeletal malformation, including shorter vertebral bodies, kyphoscoliosis, and evidence of fractures (Fig. 1, Table 2, and SI Appendix, Fig. S13). The alleles \textit{dmh13}, \textit{dmh14}, and \textit{dmh29} all cluster in the major ligand binding region 3 of type I collagen (Fig. 7), and while this region represents a hotspot for ligand interactions, mutations in this region are associated with variable phenotypic outcomes, ranging from moderate OI type IV to perinatal lethality (OI type II) (3, 34). This variability of phenotypic severity was also observed upon comparing the three different zebrafish mutants, with \textit{collala}^{chim} mutant fish displaying the most severe skeletal phenotype with kyphoscoliosis and shorter, thicker, and overmineralized vertebral bodies and frequent rib fractures (Fig. 1, Table 2, and SI Appendix, Figs. S14, S15, and S17). The \textit{collala med} allele, reported earlier by Asharani et al. (20), was shown to have no effect in heterozygous state but caused pronounced effects on TMD, volume, and thickness of the vertebral bodies when present in homozygous state (SI Appendix, Figs. S11 and S12). However, as no comparable mutations have been identified in human patients, this \textit{collala}^{med/med} mutant is likely less relevant for human OI.

To detect the presence of collagen overmodification, which is typically caused by collagen I glycine substitutions in human OI patients, collagen from adult bone was extracted by heat denaturation and subjected to SDS/PAGE. While migration of the \(\alpha_1\) and \(\alpha_3\) bands of each sample was performed and mass-spectrometry analysis was used to determine the relative amount of \(\alpha_1(0)\) in each sample. The expected \(\alpha_3(0)/\alpha_1(0)\) ratios (red lines in diagram) were approximated in samples from individual mutants (blue bars), illustrating that the reported nonsense mutations in \textit{collala} and \textit{collalb} result in loss of the \(\alpha_1(0)\) and \(\alpha_3(0)\) chain, respectively. Accordingly, in \textit{collalb}^{−/−} mutants, no tryptic peptides of \(\alpha_3(0)\) could be detected, confirming decay of mutant \textit{collalb} mRNA transcripts. (B) SDS/PAGE analysis of bone collagen extracted from individual adult spines of \textit{col1a2}^{−/−} and \textit{collalb}^{−/−} mutant fish and control siblings. For each genotype, three biological replicates were taken into account. \textit{Collalb}^{−/−} mutant fish show a complete absence of \textit{collalb} encoded \(\alpha_2(0)\) (red arrowhead). Note also the absence of the \(\beta_{1,2}\) dimers (black arrowhead) in \textit{collalb}^{−/−} mutant fish.

**Discussion**

In this work, we present a skeletal phenomic analysis of a set of zebrafish with mutations in type I collagen that model different
type I collagenopathies, including the classic types of OI and a specific subtype of EDS. By systematically analyzing the skeletal phenotypes in this set of mutants, we illustrate the high ability of zebrafish mutants to reproduce the genetic and phenotypic features of type I collagenopathies.

OI in human patients is characterized by bone fragility, leading to a higher susceptibility to fractures, and in addition, misshapen bones and skeletal malformation can be observed (7). The presence of these disease features and the extent to which they manifest is largely associated with the specific subtype of OI (7). As such, quantitative defects in type I collagen, underlying OI type I, tend to cause mild skeletal defects in human patients with an increased susceptibility to fractures but minimal bone deformity. Zebrafish mutants with similar quantitative type I collagenopathies, including the classic types of OI and a specific subtype of EDS. By systematically analyzing the skeletal phenotypes in this set of mutants, we illustrate the high ability of zebrafish mutants to reproduce the genetic and phenotypic features of type I collagenopathies.

Fig. 4. col1a1a\(^{-/-}\);col1a1b\(^{-/-}\) mutant fish exhibit intragroup variability in phenotypic expressivity. (A) Different col1a1a\(^{-/-}\);col1a1b\(^{-/-}\) mutant siblings were analyzed using \(\mu\)CT scanning at 5 mo of age (mutant 1 to mutant 5). Upon comparison of the different mutant siblings, both incomplete penetrance and a variable expressivity of features indicative of compromised bone quality was observed, which include fractures (arrowheads) and local compressions in the vertebral column causing a distorted shape of the vertebral bodies (brackets). (B) Intragroup variability for col1a1a\(^{-/-}\);col1a1b\(^{-/-}\) mutant fish is demonstrated for volume and TMD of the vertebral centra, with values in the mutant group showing a much larger spread than compared with the control group. The different colors in one graph represent different individual fish of the same group (mutant siblings or control siblings).

Fig. 5. In-depth analysis of mutants with partial or complete loss of col1a1a and/or col1a1b. (A) Segregation analysis of the different mutant genotypes generated from an in-cross of col1a1a\(^{-/-}\);col1a1b\(^{-/-}\) double-heterozygous mutant fish. Preny were genotyped at three different time points (100 fish per time point): 7 dpf, 15 dpf, and adult stage (3 mo). The relative abundance of each genotype is given for each time point as a percentage. “Expected” indicates the theoretical abundance of each genotype, following Mendelian segregation. All combinations with a complete loss of col1a1a (col1a1a\(^{-/-}\), indicated in red shades), are less viable and completely lost when progeny reached adulthood. Genotypes that contain a complete loss of col1a1b (col1a1b\(^{-/-}\), indicated in green shades) are still present at adult stage. Black indicates the parental genotype, white indicates wild-type fish, and gray represents the other remaining genotypes. (B) Phenotype of col1a1a\(^{-/-}\) mutant larvae at 7 dpf, compared with wild-type siblings. Col1a1a\(^{-/-}\) mutant larvae present with a noninflated swimbladder (asterisk) and a ruffled finfold (arrowhead). (C) Ventral view and detail of the pectoral fins showing frilly distal margins (arrowhead) in the pectoral fins of col1a1a\(^{-/-}\) mutant larvae at 7 dpf. (D) Detail of the col1a1a\(^{-/-}\) mutant larvae finfold at 7 dpf, showing an absence of fin actinotrichia fibers, which can be clearly observed in the finfold of wild-type siblings (arrow).
In-depth analysis of \textit{col1a2}^+/+ and \textit{col1a2}^−/− mutant fish. (A) Visual phenotype of \textit{col1a2}^+/+ control and \textit{col1a2}^−/− mutant fish and wild-type siblings. Both \textit{col1a2}^+/+ control, and \textit{col1a2}^−/− mutant fish display a normal morphology, while \textit{col1a2}^−/− mutant are dysmorphic and display kyphosis (arrowhead). The typical stripe pattern in the skin is disturbed in \textit{col1a2}^−/− mutant fish, but normal in \textit{col1a2}^+/+ controls and \textit{col1a2}^−/− mutant fish. (B) Masson’s Trichrome staining of histological sections of the skin of an adult \textit{col1a2}^+/+ mutant fish and a \textit{col1a2}^−/− control sibling. Note the much thinner dermis (asterisk), composed of layers of collagen fibrils (stained blue) in \textit{col1a2}^−/− mutant fish compared with \textit{col1a2}^+/+ controls. (C) Average maximum tensile strength, measured by biomechanical testing of skin flaps dissected from five adult \textit{col1a2}^−/− mutant fish and five \textit{col1a2}^+/+ control fish, illustrating strongly decreased strength of soft connective tissues in mutant fish ($P < 0.0001$). (D) Histological analysis of the vertebral column of adult fish, sagittal sections stained with H&E. Shown at the left are two adjoining vertebral body endplates (Vep, i.e., the region where two adjacent vertebral bodies meet and are connected by the IVL) in a \textit{col1a2}^+/+ control and a \textit{col1a2}^−/− mutant adult fish. The black rectangles indicate the region which is shown at a higher magnification in the image next to it on the right, depicting the IVL in detail. In control fish, the IVL is typically composed of three distinct layers or tissues: the notochord sheet (Ns, mainly type II collagen), the outer elastin layer (OEL), and a layer of type I collagen fiber bundles (cl). In the \textit{col1a2}^−/− mutants, IVL composition is abnormal: the OEL was still present, the Ns layer could be observed but was extremely reduced, but no layer of type I collagen fibers (cl) could be observed. \textit{Col1a2}^−/− mutant bone (B) in the vertebral bodies near the Vep showed a loss of typical Sharpey fibers (Sf), while the notochord cells (Nc) displayed a loss of vacuoles, in addition to scarring (Sc). As shown in the two rightmost pictures, we observed fusions of adjacent vertebral bodies (black arrowhead) and dislocation of the IVL and Vep (red arrow) in some of the vertebrae, likely resulting in kyphosis seen in \textit{col1a2}^−/− mutant fish. Other abbreviations: Ep, epidermis; Mu, muscle fibers; Ob, osteoblasts; Scl, Scale.

Defects (\textit{coll1a1a}^{+/−};\textit{coll1a1b}^{+/−} mutants) also presented with increased fracture risk, apparent from the presence of spontaneous fractures in the ribs, while skeletal deformities were minimal with only localized and mild compressions of the vertebral column in some of the mutant fish. Conversely, qualitative type I collagen defects in zebrafish evoked a more pronounced effect on the skeleton, with frequent fractures, kyphoscoliosis and malformation of the vertebral bodies, and misshapen ribs and fin bones. These phenotypes correspond to the moderately severe, progressive deforming or lethal clinical phenotypes of OI types II–IV. Importantly, the more severely affected mutants of this group (\textit{dmh15} and \textit{dmh29}) also displayed overmineralization in the vertebral column, a feature often seen in human OI, and especially associated with the more severe cases in human patients (32, 33). Additionally, biochemical analysis argued for type I collagen overmodification, one of the typical features in human OI, in some of the mutants with a qualitative defect. Moreover, the variability of the excessive posttranslational modification throughout this set of mutants corresponds to the situation in human patients with OI types II–IV, where variable collagen overmodification has been related to the position of the mutation in the type I collagen helix (34, 36). Overall, the genotype–phenotype analysis we performed shows the validity of the included zebrafish mutants as models for different forms of human OI.

A hallmark feature of OI in humans is the large clinical spectrum of phenotypes, with disease presentation that can differ from very mild to perinatal lethal, even with the same causative mutation. The underlying molecular basis of this variability remains largely not understood. The zebrafish mutants described here similarly show significant variation in their expressivity. Intragenotype variability is illustrated in \textit{coll1a1a}^{+/−};\textit{coll1a1b}^{+/−} mutant siblings, as some show fractured ribs or vertebral compressions, while others do not. In addition, there is a large variability in vertebral morphology and mineralization in mutant siblings, in contrast to control siblings. A likely explanation for
The intragenotype phenotypic variability in zebrafish mutants is the highly polymorphic nature of its genome, which closely resembles human genetic variability. This makes them particularly useful to unveil genetic determinants that can be used to map potential regulation of phenotypic variability in humans. This is less obvious in mouse models, where mostly inbred strains are used for disease modeling (37, 38).

In zebrafish, type I collagen has been shown to have a different composition than in mammals, with two orthologs of the human COL1A1 gene, namely col1a1 and col1ab, encoding the α1- and α3-chain, respectively. Some insights in these differences have already been addressed previously (19), and by studying different knockout mutants for the type I collagen genes we further extend this knowledge. Heterozygous loss of either α1(1) or α3(1) did not induce any skeletal abnormalities, while reduced levels of both α1(1) and α3(1) (col1a1α+/−; col1abβ+/− mutants) causes a mild skeletal phenotype with increased bone fragility, arguing for interchangeability and functional redundancy between α1(I) and the homologous α3(I). Complete loss of the α1(I) chain causes lethality after 7 dpf, while loss of α3(I) only shows increased early lethality if the amount of α1(I) is also diminished. This suggests a gene/protein dosage effect, where α1(I) is more abundant than α3(I), as hypothesized in our previous work (19). The functional similarity between α1(I) and α3(I) is further illustrated by the fact that glycine substitutions in both genes can cause severe skeletal phenotypes, arguing for a dominant-negative effect on type I bone collagen and thus incorporation of a substantial amount of both α1(I) and α3(I) into the type I collagen triple helix. However, to obtain a full understanding of zebrafish type I collagen, the exact trimer stoichiometry in different tissues, and the abundance of each possible trimer composition should be examined, which is technically challenging due to the small specimen size of zebrafish compared with the large amount of input material needed for standard biochemical analysis techniques.

In humans, haploinsufficiency for the α2(I)-chain does not cause a clinically significant phenotype, while complete loss of α2(I) leads to the very rare cardiac valvular subtype of the EDS, associated with soft connective tissue defects, including skin and joint manifestations and propensity to cardiac valvular defects, and relatively mild skeletal manifestations (6). These trends were mostly recapitulated in zebrafish, as col2aα−/− mutants are asymptomatic, whereas col2a2α−/− mutants display an EDS-like phenotype with different connective tissue defects; µCT-scanning showed local vertebral abnormalities, including kyphosis, and a mild reduction of bone thickness and mineralization. Histological analysis demonstrated dislocation and distortion of the fibrous connections between some of the vertebral bodies, caused by degeneration of the layers comprising the connecting intervertebral ligament. Most likely the latter defect is the cause of the kyphosis seen in these mutants, with the location of the kyphosis being the area bearing the highest mechanical force in the fish vertebral column (34). Similar to these observations, patients with the same molecular defect display joint dislocations but no severe skeletal involvement (6). Interestingly, col2a2α−/− mutant fish showed fragile skin upon handling and histological analysis demonstrated a much thinner dermis in the skin of mutant fish compared with wild-type siblings. Consistent with these findings, bio-mechanical testing argued for a strongly reduced strength of soft connective tissues in col2a2α−/− mutant fish. This relates to other clinical hallmark features of the skin in human cardiac valvular EDS patients (6). Preliminary experiments in col2a2α−/− mutant fish could not provide any proof of defects in cardiac morphology or functioning. However, more extensive analyses are needed to further explore a possible impaired cardiac function in a follow-up study, focusing on an in-depth characterization of this mutant, which was beyond the scope of the study presented here.

Taken together, we have analyzed a large set of zebrafish models with mutations in type I collagen that represent different types I collagenopathies, including the classic types of OI and a specific subtype of EDS, and illustrate the high phenotypic and genetic similarity of these zebrafish mutants with human type I collagenopathies. We further report a zebrafish model resembling the human EDS, which harbors a number of characteristic defects in the soft connective tissues. Our study surpasses the analysis at the single-mutant model level but illustrates the occurrence of relevant phenotypic patterns and characteristics across a large set of genetically distinct zebrafish models. These results, taken together with the feasibility of zebrafish for easily accommodating large genetic and phenotypic screens, argue for zebrafish as a promising tool in future research aiming to dissect the underlying basis of phenotypical variability in human type I collagenopathies, such as OI. Furthermore, we provide more insight into the biology of zebrafish type I collagen and its consequences, which are relevant in the study of any human type I collagen related disease by using zebrafish as a model.

Material and Methods

Animals. The col1a1α+1748+; col1a1β−12931, col1a2α−17981, bmp1a+2415, and plod2α−1748 mutant zebrafish were generated by the zebrafish mutation project and together with wild-type AB fish obtained from the zebrafish international resource center (EZRC, zebrafish.org/home/guide.php) or the European Zebrafish Resource center [EZRC, www.ezrc.kit.edu]) (39). The col1a1α−/− were previously described (21) and the col1a1αmed mutant fish were
purchased from the EZRC. The colla1a1Δm13, colla1a1Δm14, colla1bΔm23, and colla2Δm59 mutant fish were generated in a forward genetics screen (40). Fish husbandry was performed as previously described (22). Studies were done in agreement with EU Directive 2010/63/EU for animals, permit Number: ECD 17/58. All efforts were made to minimize pain and discomfort.

**μCT Scanning and Analysis.** μCT-based phenotyping and quantification of adult zebrafish was performed as previously described (25). For each genotype five mutants and five control fish were analyzed.

**Collagen Analysis.** Collagen analysis of adult zebrafish bone was performed as described previously (22). To determine a relative amount of α1(1)- and α3(1)-chains, a unique tryptic peptide for each α1(1) and for α3(1) was selected and the relative amount of the corresponding peak manually measured on the full scan by determining peak height. For all mutant genotypes an expected ratio of α3(1)/α1(1) was calculated based on the mean relative α3(1)/α1(1) ratio observed in control samples and based on the assumption of absence of α3(1) and α1(1) in the case of knockout of colla1a and colla1b, respectively.

**Histological Analysis.** Histological analysis of zebrafish tissues was performed as described in Gistelinck et al. (22).

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