Research Article

Mutational Screening of Skeletal Genes in 14 Chinese Children with Osteogenesis Imperfecta Using Targeted Sequencing

Wei Tan,1 Yuelun Ji,1 Yuepeng Qian,1 Yongchang Lin,1 Ruolian Ye,1 Weiping Wu,1 Yibin Li,1 Yongjian Sun,1 and Jianyin Pan2

1Department of Pediatric Orthopedic, Center for Orthopedic Surgery, The Third Affiliated Hospital of Southern Medical University, Guangzhou 510515, China
2Department of Joint Surgery, Center for Orthopedic Surgery, The Third Affiliated Hospital of Southern Medical University, Guangzhou 510515, China

Correspondence should be addressed to Yongjian Sun; nfyysyj@163.com and Jianyin Pan; storm0132002@163.com

Received 15 March 2022; Revised 28 April 2022; Accepted 3 May 2022; Published 19 May 2022

Academic Editor: Fu Wang

Copyright © 2022 Wei Tan et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Background. As a heterogeneous hereditary connective tissue disorder, osteogenesis imperfecta (OI) is clinically characterized by increased fracture susceptibility. Analysis of genetic pathogenic variants in patients with OI provides a basis for genetic counseling and prenatal diagnosis. Methods. In this study, 14 diagnosed OI patients from sporadic Chinese families were enrolled to be screened for potential mutations from these patients by next-generation sequencing technology. Results. 34 different variants were identified. 18 variants were from 4 OI-related genes including COL1A1, COL1A2, P3H1, and WNT1, and 10 variants are novel. Most OI patients (11 out of 14, 78%) harbor variants in type I collagen genes. Conclusions. Our results support previously established estimates of the distribution and prevalence of OI mutations and highlight both phenotype and genetic heterogeneity among and within families. We report several novel variants of OI, which expands the clinical spectrum of OI. In summary, our data provides disease-causing genes information for genetic counseling towards OI patients and families and also provides a reference for clinicians in the diagnosis of OI, also in prenatal diagnosis of this disease.

1. Introduction

Osteogenesis imperfecta (OI), also known as brittle bone disease, comprises a class of connective tissue disorders characterized by increased bone fragility and vulnerability to fracture due to minor trauma. Other features include brittle teeth, low bone mass, hearing loss, and blue sclera. The incidence of OI was estimated to be 6 to 7 per 100,000 people worldwide [1]. OI is clinically and genetically heterogeneous and historically had various other names. The name OI dates back to at least 1895, and the current four-type system with Sillence began in 1979 [2]. Using clinical, radiographic, and genetic criteria, OI individuals were categorized into four types: type I (mild OI with blue sclera, Mendelian Inheritance in Man (MIM 166200)), type II (pre- or perinatal lethal, MIM 166210), type III (severe deforming, MIM 259420), and type IV (intermediated severity, MIM 166220) in 1979 [3]. However, the correlation between specific genetic defects and clinical manifestations is confused due to the apparent overlap between clinical manifestations and disease severity. Therefore, a new classification system was proposed based on the specific genetic defects that cause the disease [4]. The new revision of classification types 1-4 is closely related to Sillence’s previous I-IV classification system. It is mainly based on specific phenotypic characteristics and genetic findings describing other OI types (V and
higher). There are currently 23 distinct OI types according to the affected genes according to the Online Mendelian Inheritance in Man (OMIM) resource (https://omim.org). It is established that 90% of OI was caused by variants in the two collagen coding genes (COLIA2 and COLIA1), which cause a lack of type I collagen [5]. These genetic defects are often inherited from a person’s parent in an autosomal dominant manner or occur via a de novo mutation. Currently, more and more pathogenic genes have been discovered, which further facilitate OI classification. Variants of CRTAP (MIM 605497) and SERPINF1 (MIM 172860) may be the cause of type VII and VI OI, respectively [6, 7]. The remaining types are usually associated with mutations in some autosomal recessive genes, such as P3H1 (MIM 610339) [8], WNT1 (MIM 164820) [9], and TENT5A (MIM 611357) [10]. The genetic classification incorporated a new type for each defective mutation. In total, OI is considered to be a spectrum with considerable phenotype overlap between the subtypes. The application of next-generation sequencing technologies (NGS), such as targeted sequencing, whole exome (WES), or whole genome sequencing (WGS), has greatly improved the diagnosis and classification of OI. In particular, targeted sequencing of candidate genes is considered as a cost-effective strategy for the clinical diagnosis of heterogeneous disorders [11]. A recent study reported that targeted sequencing has been applied to prenatal diagnosis of skeletal dysplasia [12].

Considering the increasing popularity of molecular diagnosis during OI classification, we assembled a cohort of 14 OI patients of Chinese ethnicity and investigated their mutation spectrum using a targeted skeletal gene panel. Other than known variants, we identified novel variants associated with OI and revealed the genotype-phenotype relationships.

2. Methods

2.1. Patients. There were 14 unrelated probands with OI recruited at The Third Affiliated Hospital of Southern Medical University, Guangdong, China, in this project. After obtaining appropriate informed consent from participants and/or legal guardians, the whole blood samples were collected from the affected individuals. All affected individuals including the probands and their family members were of Chinese Han ethnicity. The complete pedigree information for the individuals is summarized in Figure 1. The details of the medical history, phenotype features, and physical examination were obtained and summarized in Table 1. This study was approved by the Institutional Review Board of the Third Affiliated Hospital of Southern Medical University.

2.2. Targeted Sequencing and Variant Identification. Due to the clinical and genetic heterogeneity of OI, targeted sequencing was performed on all 14 patients to identify potential variants. Briefly speaking, genomic DNA was extracted from the peripheral blood of patients, using a customized array (NimbleGen, Roche Madison, USA) to capture the whole exons and flanking intronic sequences (±100bp) from 225 genes known to be associated with inherited skeletal disorders and then followed by high-throughput sequencing on Illumina HiSeq 2000 platform with 150 bp paired-end reads (supplementary Table S1). Bioinformatics analysis was performed using public software and independently developed pipelines. In bioinformatics analysis, all cleaned data after trimming were aligned against human reference genome build hg19 using Burrows-Wheeler Alignment (BWA) tool. The mean depths was 150x across all patients. With the integration of customized databases, SNP and Indels were then discovered by HaplotypeCaller module of the Genome Analysis Toolkit (GATK) and annotated by ANNOVAR. Analysis was restricted to coding and flanking intronic regions which generally ensure good data coverage. Variants with minor allele frequency (MAF) greater than 1% in any of the Exome Aggregation Consortium (ExAC), the Genome Aggregation Database (gnomAD), NHLBI GO Exome Sequencing Project (ESP6500) databases, and 1000 genome project were excluded from subsequent analysis. Then, filtering and prioritizing were performed to look into potentially deleterious variants, including missense, nonsense, indel, frameshift, and variants affecting splicing, for variant interpretation and classification according to American College of Medical Genetics (ACMG) guidelines.

Fragments covering the mutation sites of the relevant genes were amplified by PCR followed by Sanger sequencing to confirm identified mutations of interest. According to the manufacturer’s instructions of Premix Taq® Hot Start Version/TaKaRa LA Taq® with GC Buffer (Takara, Osaka, Japan), PCR amplification was performed under the following conditions: 1 cycle of initial denaturation at 95°C for 2 min and 35 cycles of 95°C for 30 s, 60°C for 30 s, 72°C for 30 s, and 5 min at 72°C. Using an ABI 3730 DNA Analyzer with BigDye™ Terminator Cycle Sequencing Kit (Applied Biosystems, Foster, CA, USA), polymerase chain reaction (PCR) fragments were purified and further sequenced.

3. Results

Fourteen clinically and radiologically confirmed patients with OI were selected for this study. Out of them, 9 were males and the remaining 5 were females. Most of them were children, and the average age was 7 years old. All the patients were type III except patient 11. Blue sclera is an important clinical characteristic of OI patients and was observed in 13 patients in our cohort. 10 patients suffered tooth agenesis, and almost all patients (13 in 14) have fracture history. The number of fractures in these patients ranged from 0 to 8. Out of 34 heterozygous variations in total identified, 18 polymorphisms in 4 OI-related genes including COLIA1, COLIA2, P3H1, and WNT1 and the remaining 16 were in other genes (Figure 2).

The 18 polymorphisms in the 4 OI-related genes were shown in Table 2. 10 variants are novel. 3 mutations are localized within the triple helix repeat of the COLIA1 gene, 2 in COLIA2 gene, and 1 in P3H1 gene. All mutations of WNT1 gene are novel. Four probands carried splicing variants, including three from COLIA1 (c.2451+1G>A, c.1821+1delG, and c.693+5G>A) and one from WNT1 (c.104
These four splicing variants were closely related with mild to moderate phenotype features predominantly characterized by short stature and recurrent fractures. Three probands had frameshift mutations, including two from COL1A1 (p.G1046Vfs*62 and p.A606Cfs*27) and one from WNT1 (p.R73Tfs*82). One proband carried null mutation in P3H1 (p.R210X), and the rest were nonsynonymous mutations. Consistent with the previous reports, COL1A2 and COL1A1 were the two most frequently mutated genes (11 out of 14 patients). Locations of variants in COL1A2 and COL1A1 in individuals were shown in Figure 3. 7 probands

M: male; F: female; y: year; T: times.

| Patient ID | Gender | Age (y) | Blue sclera | Tooth agenesis | Fracture (T) | Contractures | Hearing loss |
|------------|--------|---------|-------------|----------------|--------------|--------------|--------------|
| 1          | Male   | 13.8    | Yes         | No             | 7            | No           | No           |
| 2          | Male   | 11.1    | Yes         | No             | 3            | No           | No           |
| 3          | Male   | 9.3     | Yes         | Yes            | 7            | No           | No           |
| 4          | Female | 3.6     | Yes         | Yes            | 2            | No           | No           |
| 5          | Male   | 2.5     | Yes         | Yes            | 1            | No           | No           |
| 6          | Female | 0.75    | Yes         | No             | 0            | No           | No           |
| 7          | Male   | 15      | No          | Yes            | 5            | No           | No           |
| 8          | Female | 14.8    | Yes         | Yes            | 8            | No           | No           |
| 9          | Male   | 7       | Yes         | Yes            | 3            | No           | No           |
| 10         | Male   | 0.6     | Yes         | Yes            | 3            | No           | No           |
| 11         | Male   | 10.9    | Yes         | Yes            | 3            | No           | No           |
| 12         | Female | 8.7     | Yes         | Yes            | 3            | No           | No           |
| 13         | Male   | 2.75    | Yes         | No             | 7            | No           | No           |
| 14         | Female | 5       | Yes         | Yes            | 2            | No           | No           |
had a COL1A1 variant. Three variants including c.1721G>A, c.1816_1817del, and c.2069C>T were novel, and the rest four had been reported previously, which include c.2451+1G>A, c.2443G>A, c.1821+1delG, and c.3135delT [13–16]. 5 probands had a COL1A2 variant, out of which three variants have been reported in previous studies, including c.1801G>A in patient 7, c.693+5G>A in patient 9 [17], and c.964G>A in patient 10 [18]. The splicing variant in patient 9 occurred at a highly conserved site. With respect to P3H1, there was a previously reported variation (c.628C>T) [19] and a novel compound heterozygote variant (c.1703G>A). Four novel compound heterozygote variants were detected in the WNT1 gene, including two nonsynonymous mutations, one splicing variant, and one frameshift variant.

In this report, we present 4 OI probands with substitutions of glycine residue including p.G815S from COL1A1 and p.G601S, p.G391V, and p.G322S from COL1A2. The phenotypes of these probands are consistent with the previous reports that these variants of substitutions are associated with severe OI. For example, patient 5 suffered fracture of left tibia 20 days after birth and fractured right femoral shaft 2 months later. His mother carried this mutation and presented excess fractures. However, the proband with OI may be significantly affected clinically, whereas family
members with the same mutation may have no obvious symptoms (patient 6 and her father), demonstrating that the clinical manifestations vary substantially within families with different types of genetic variants.

Additionally, we also found 16 variations in 14 other genes, as shown in Table 3. There were two mutation sites in \textit{RPGRIP1L} (c.1960G > A and c.3272G > A) in patient 14. The protein encoded by \textit{RPGRIP1L} localizes to the basal body-centrosome complex in ciliated cells and takes part in Hedgehog and organelle biogenesis and maintenance pathways. Defects in \textit{RPGRIP1L} are a cause of Meckel syndrome type 5 (MKS5 and MIM 611561) and Joubert syndrome type 1464.

**Table 3: Variations in other genes.**

| Patient ID | GENE   | Inheritance | Nucleotide variant | Predicted consequence | Zygosity | ACMG pathogenic | Paternal/maternal |
|------------|--------|-------------|--------------------|-----------------------|----------|----------------|-----------------|
| 1          | SLC34A1| AR/AD       | c.1322A > G        | p.Y441C               | Het      | VUS            | Paternal        |
| 1          | LRSAM1 | AR/AD       | c.1367C > T        | p.A456V               | Het      | VUS            | Paternal        |
| 3          | ACVR1  | AD          | c.454C > T         | p.R152C               | Het      | VUS            | Maternal        |
| 4          | ACAN   | AR/AD       | c.488G > A         | p.R163H               | Het      | VUS            | Maternal        |
| 4          | FLNB   | AR/AD       | c.1669C > T        | p.P557S               | Het      | VUS            | Paternal        |
| 4          | NF1    | AD          | c.2991-11C > G     | Splicing              | Het      | VUS            | Maternal        |
| 5          | SMARCAL1| AR         | c.662G > T         | p.R221I               | Het      | VUS            | Maternal        |
| 6          | FLNA   | XL          | c.2027A > G        | p.K676R               | Het      | VUS            | Maternal        |
| 8          | TRIP11 | AR          | c.4434A > T        | p.E1478D              | Het      | VUS            | Maternal        |
| 8          | TRIP11 | AR          | c.701A > G         | p.H234R               | Het      | VUS            | Maternal        |
| 8          | COL9A1 | AD          | c.432A > G         | p.I144M               | Het      | VUS            | Maternal        |
| 12         | COL2A1 | AD          | c.4393C > T        | p.R1465C              | Het      | VUS            | Paternal        |
| 12         | HBA2   | AD          | c.427T > C         | p.X143Q               | Het      | VUS            | Paternal        |
| 13         | PTH1R  | AD/AR       | c.629C > T         | p.A210V               | Het      | VUS            | Maternal        |
| 14         | RPGRIP1L| AR         | c.3272G > A        | p.R1091Q              | Het      | VUS            | Paternal        |
| 14         | RPGRIP1L| AR         | c.1960G > A        | p.E654K               | Het      | VUS            | Paternal        |
7 (JBTS7, MIM 611560) [20]. Our results suggested that RPGRIP1L mutation may be related to OI, which is worthy of further investigation.

4. Discussion

OI is an inherited connective tissue disorder with a wide genotypic and phenotypic spectrum and up to 23 subtypes. Mutations in type I collagen or proteins involved in post-translational modification of type I collagen are the most common cause of OI. Type I collagen containing two α1 and one α2 chains is a heterotrimeric fibrillar collagen, and α1 and α2 chains are encoded by COL1A1 and COL1A2, respectively. Genetic defects include reducing the amount of type 1 collagen (quantitative defects) or affecting its structure (qualitative defects), resulting in a milder phenotype to more severe phenotype [21]. The remaining cases were caused by genetic defects in genes involved in the posttranslational modification and intracellular trafficking of type 1 collagen or genes related to osteoblast differentiation and function. In our study, 11 out of 14 patients exhibited mutation in COL1A1 or COL1A2, accounting for 78% of our cohort. There is no hotspot mutation identified in COL1A1 and COL1A2 genes.

Other than COL1A1 and COL1A2 genes, mutations in P3H1 and WNT1 were found in our OI patients. It is known that cartilage-associated protein (encoded by CRTAP), cyclophilin B (encoded by PPIB), and prolyl 3-hydroxylase 1 (encoded by P3H1) form a collagen posttranslational modification complex in a 1:1:1 ratio on the endoplasmic reticulum and regulate collagen function [8]. The null mutation in P3H1 would result in a protein decrease or absence, leading to the collapse of the complex [8, 22]. The defective structural collagen exhibits a delay in the collagen helix folding and exposes the constituent chains to lysyl hydroxylase and prolyl 4-hydroxylase for a longer period of time, leading to overmodification of the chain [8]. The truncation of P3H1 gene has been previously reported in OI individuals. In addition, we reported a novel missense variant c.1703G>A, which is the first one ever reported to the best of our knowledge. This missense variant was predicted to be a disease causing by Sift [23] and Polyphen-2 [24].

The WNT1 encodes a ligand for the WNT signalling pathway, which acts as a primary regulator in bone formation. Wnt1 promotes bone formation by binding to the LRP5-FZD receptor complex and activating the canonical WNT signalling pathway for promoting bone formation. Loss-of-function mutations in WNT1 result severe OI, severe skeletal abnormalities, white sclerae, or possible neurological defects [25]. Herein, we report four autosomal recessive (AR) forms of WNT1 variants and two are likely pathogenic. More than 3200 variants have been observed across COL1A1, COL1A2, P3H1, and WNT1 genes (http://www.le.ac.uk/ge/collagen/).

The type I collagen genes are also associated with other genetic bone diseases including Caffey disease [26], arthrochlasia Ehlers-Danlos syndrome (aEDS) [27–29], cardiac valvular EDS (cvEDS), and OI/EDS disease [30, 31]. The core phenotype is closely related to the location of deleterious defects. Mutations near the N-terminal peptide, which links the N-propeptide to the triple helical domain of collagen, are often associated with aEDS or OI/EDS [32]. Clinical symptoms of OI and OI/EDS overlap. Given the clinical and genetic heterozygosity of these disease, their diagnosis and treatment are always challenging. Hence, detailed clinical characterization of rare diseases is extremely important. In our study, mutations in type I collagen gene cluster in collagen a chain are far from the N-terminal peptide. The patient’s clinical symptoms did not show increased laxity of the ligaments and skin.

In addition to variations in 4 OI-related genes, our study also identified novel mutations in 14 other genes, such as 2 heterozygous mutations in RPGRIP1L. Whether these variants are OI causing or not is worthy of further investigation. Our data support that the autosomal dominant (AD) forms are most prevalent in our Chinese OI population. These results highlight the importance of genetic analysis in autosomal dominant disease including OI, which can significantly reduce the risk of this serious disease. 6 probands were AR forms of OI. Previous studies have reported that approximately 10-20% of germline pathogenic mutations may alter mRNA splicing. There were four splicing mutations found in our analysis. In one of these cases, no pathogenic variants were identified in known OI genes, making them suitable for WGS or WES to identify novel underlying genetic defects of OI.

In summary, our study provides an estimate of the prevalence and the distribution of OI mutations and further highlight both phenotypic and genetic heterogeneity among and within families. The technological improvements and decreasing cost will make NGS approaches an effective first-tier screening method, and the targeted sequencing of interested gene panels has become part of routine clinical genetic practice for the diagnosis of heterogeneous disorders [33]. However, our study was limited to a single center and was retrospective. A larger number of phenotype-genotype correlative studies are required in order to provide better insights into OI.

Data Availability

The data used to support the findings of this study are available from the corresponding author upon request.

Ethical Approval

This investigation was approved by the Institutional Review Board of the Third Affiliated Hospital of Southern Medical University.

Consent

Patients or legal guardians signed informed consent regarding publishing their data and photographs. Informed consent was obtained from all participants or legal guardians included in the study.
Conflicts of Interest

All authors declare no conflict of interest.

Authors’ Contributions

Conception and design was carried out by W Tan, Y Sun, and J Pan; Y Sun and J Pan were responsible for the administrative support; W Tan, Y Ji, and Y Qian were responsible for the provision of the study materials or patients; Y Ji, Y Lin, Y Li, and R Ye were responsible for the collection and assembly of data; W Tan, Y Ji, and Y Sun were responsible for the data analysis and interpretation; all authors wrote the manuscript; all authors approved the final manuscript. Wei Tan and Yuelun Ji contributed equally to this work.

Acknowledgments

We thank all patients and their parents who participated in this study. We thank Dr. Yongchu Liu from the Aegicare (Shenzhen) Technology Co., Ltd., for the advice and discussions.

Supplementary Materials

Supplementary material of the description of NGS. Supplementary Table S1: 225 genes included in the skeletal gene panel. (Supplementary Materials)

References

[1] A. M. Barnes, W. Chang, R. Morello et al., “Deficiency of cartilage-associated protein in recessive lethal osteogenesis imperfecta,” The New England Journal of Medicine, vol. 355, no. 26, pp. 2757–2764, 2006.
[2] J. C. Marini, A. Forlino, H. P. Bächinger et al., “Osteogenesis imperfecta,” Nature Reviews Disease Primers., vol. 3, no. 1, p. 17052, 2017.
[3] D. O. Sillence, D. L. Romin, and D. M. Danks, “Clinical variability in osteogenesis imperfecta-­variable expressivity or genetic heterogeneity,” Birth Defects Original Article Series, vol. 15, no. 5B, pp. 113–129, 1979.
[4] L. Bonafe, V. Cormier-Daire, C. Hall et al., “Nosology and classification of genetic skeletal disorders: 2015 revision,” American Journal of Medical Genetics. Part A, vol. 167A, no. 12, pp. 2869–2892, 2015.
[5] B. Sykes, “Linkage analysis in dominantly inherited osteogenesis imperfecta,” American Journal of Medical Genetics, vol. 45, no. 2, pp. 212–216, 1993.
[6] J. Becker, O. Semler, C. Gilissen et al., “Exome sequencing identifies truncating mutations in human SERPINF1 in autosomal-recessive osteogenesis imperfecta,” American Journal of Human Genetics, vol. 88, no. 3, pp. 362–371, 2011.
[7] R. Morello, T. K. Bertin, Y. Chen et al., “CRTAP is required for prolyl 3-hydroxylation and mutations cause recessive osteogenesis imperfecta,” Cell, vol. 127, no. 2, pp. 291–304, 2006.
[8] W. A. Cabral, W. Chang, A. M. Barnes et al., “Prolyl 3-hydroxylase 1 deficiency causes a recessive metabolic bone disorder resembling lethal/severe osteogenesis imperfecta,” Nature Genetics, vol. 39, no. 3, pp. 359–365, 2007.
[9] R. Shanshan, B. Koen, and A.-A. Zaid, “Efficient acceleration of the pair-HMMs forward algorithm for GATK Haplotype-Caller on graphics processing Units,” Evolutionary Bioinformatics Online, vol. 14, 2018.
[10] M. Doyard, S. Bacrot, C. Huber et al., “FAM46A mutations are responsible for autosomal recessive osteogenesis imperfecta,” Journal of Medical Genetics, vol. 55, no. 4, pp. 278–284, 2018.
[11] V. Nigro and M. Savarese, “Next-generation sequencing approaches for the diagnosis of skeletal muscle disorders,” Current Opinion in Neurology, vol. 29, no. 5, pp. 621–627, 2016.
[12] X. Y. Zhou, N. Chandler, L. B. Deng, J. Zhou, M. Z. Yuan, and L. M. Sun, “Prenatal diagnosis of skeletal dysplasias using a targeted skeletal gene panel,” Prenatal Diagnosis, vol. 38, no. 9, pp. 692–699, 2018.
[13] J. Schleit, S. S. Bailey, T. Tran et al., “Molecular outcome, prediction, and clinical consequences of splice variants in COL1A1, which encodes the proα1(I) chains of type I procollagen,” Human Mutation, vol. 36, no. 7, pp. 728–739, 2015.
[14] I. M. Ben Amor, P. Roughley, F. H. Glorieux, and F. Rauch, “Skeletal clinical characteristics of osteogenesis imperfecta caused by haploinsufficiency mutations in COL1A1,” Journal of Bone and Mineral Research, vol. 28, no. 9, pp. 2001–2007, 2013.
[15] J. C. Marini, A. Forlino, W. A. Cabral et al., “Consortium for osteogenesis imperfecta mutations in the helical domain of type I collagen: regions rich in lethal mutations align with collagen binding sites for integrins and proteoglycans,” Human Mutation, vol. 28, no. 3, pp. 209–221, 2007.
[16] K. Kataoka, O. Ogura, K. Hasegawa et al., “Mutations in type I collagen genes in Japanese osteogenesis imperfecta patients,” Pediatrics International, vol. 49, no. 5, pp. 564–569, 2007.
[17] F. Malfait, S. Symoens, N. Goemans et al., “Helical mutations in type I collagen that affect the processing of the amino-­propeptide result in an osteogenesis imperfecta/Ehlers-Danlos syndrome overlap syndrome,” Orphanet Journal of Rare Diseases, vol. 8, no. 1, p. 78, 2013.
[18] H. Hartikka, K. Kuurila, J. Korkko et al., “Lack of correlation between the type of COL1A1 or COL1A2 mutation and hearing loss in osteogenesis imperfecta patients,” Human Mutation, vol. 24, no. 2, pp. 147–154, 2004.
[19] A. Willaert, F. Malfait, S. Symoens et al., “Recessive osteogenesis imperfecta caused by LEPRE1 mutations: clinical documentation and identification of the splice form responsible for prolyl 3-hydroxylation,” Journal of Medical Genetics, vol. 46, no. 4, pp. 233–241, 2009.
[20] M. Delous, L. Baala, R. Salomon et al., “The ciliary gene RPGRIP1L is mutated in cerebello-­oculo-­renal syndrome (Joubert syndrome type B) and Meckel syndrome,” Nature Genetics., vol. 39, no. 7, pp. 875–881, 2007.
[21] T. Symeon and A. D. Dede, “Osteogenesis imperfecta - A clinical update,” Metabolism: clinical and experimental, vol. 80, pp. 27–37, 2018.
[22] W. Chang, A. M. Barnes, W. A. Cabral, J. N. Bodurtha, and J. C. Marini, “Prolyl 3-hydroxylase 1 and CRTAP are mutually stabilizing in the endoplasmic reticulum collagen prolyl 3-hydroxylation complex,” Human Molecular Genetics, vol. 19, no. 2, pp. 223–234, 2010.
[23] R. Vasar, S. Adusumalli, S. N. Leng, M. Sikic, and P. C. Ng, “SIFT missense predictions for genomes,” Nature Protocols, vol. 11, no. 1, pp. 1–9, 2016.
[24] I. A. Adzhubei, S. Schmidt, L. Peshkin et al., “A method and server for predicting damaging missense mutations,” *Nature Methods*, vol. 7, no. 4, pp. 248-249, 2010.

[25] S. Alhamdi, Y. C. Lee, S. Chowdhury et al., “Heterozygous WNT1 variant causing a variable bone phenotype,” *American Journal of Medical Genetics. Part A*, vol. 176, no. 11, pp. 2419–2424, 2018.

[26] R. C. Gensure, O. Makitie, C. Barclay et al., “A novel COL1A1 mutation in infantile cortical hyperostosis (Caffey disease) expands the spectrum of collagen-related disorders,” *The Journal of Clinical Investigation*, vol. 115, no. 5, pp. 1250–1257, 2005.

[27] W. G. Cole, D. Chan, G. W. Chambers, I. D. Walker, and J. F. Bateman, “Deletion of 24 amino acids from the pro-alpha 1(I) chain of type I procollagen in a patient with the Ehlers-Danlos syndrome type VII,” *The Journal of Biological Chemistry*, vol. 261, no. 12, pp. 5496–5503, 1986.

[28] D. Weil, M. D’Alessio, F. Ramirez et al., “A base substitution in the exon of a collagen gene causes alternative splicing and generates a structurally abnormal polypeptide in a patient with Ehlers-Danlos syndrome type VII,” *The EMBO Journal*, vol. 8, no. 6, pp. 1705–1710, 1989.

[29] D. R. Eyre, F. D. Shapiro, and J. F. Aldridge, “A heterozygous collagen defect in a variant of the Ehlers-Danlos syndrome type VII. Evidence for a deleted amino-telopeptide domain in the pro-alpha 2(I) chain,” *The Journal of Biological Chemistry*, vol. 260, no. 20, pp. 11322–11329, 1985.

[30] U. Schwarze, R. Hata, V. A. McKusick et al., “Rare autosomal recessive cardiac valvular form of Ehlers-Danlos syndrome results from mutations in the COL1A2 gene that activate the nonsense-mediated RNA decay pathway,” *American Journal of Human Genetics*, vol. 74, no. 5, pp. 917–930, 2004.

[31] Y. Lu, Y. Wang, F. Rauch et al., “Osteogenesis imperfecta type III/Ehlers-Danlos overlap syndrome in a Chinese man,” *Intractable & Rare Diseases Research*, vol. 7, no. 1, pp. 37–41, 2018.

[32] N. Zhai, Y. Lu, Y. Wang et al., “Splice receptor-site mutation c. 697-2A > G of the COL1A1 gene in a Chinese family with osteogenesis imperfecta,” *Intractable & Rare Diseases Research*, vol. 8, no. 2, pp. 150–153, 2019.

[33] C. Gonzaga-Jauregui, J. R. Lupski, and R. A. Gibbs, “Human genome sequencing in health and disease,” *Annual Review of Medicine*, vol. 63, no. 1, pp. 35–61, 2012.