High bone mass phenotype in a cohort of patients with Osteogenesis Imperfecta caused due to BMP1 and C-propeptide cleavage variants in COL1A1

E.H. Campanini, D. Baker, P. Arundel, N.J. Bishop, A.C. Offiah, S. Keigwin, S. Cadden, E. Dall’Ara, N. Nicolaou, S. Giles, J.A. Fernandes, M. Balasubramanian

Abstract

Objectives: Osteogenesis Imperfecta (OI) is a heterogeneous condition mainly characterised by bone fragility; extra-skeletal features in OI include blue sclerae, dentinogenesis imperfecta, skin laxity and joint hyper-extensibility. Most patients with OI are thought to have a low bone mass but contrary to expectations there are certain forms of OI with high bone mass which this study explores in further detail.

Method: A cohort of n=6 individuals with pathogenic variants in BMP1 and the C-propeptide cleavage variants in COL1A1 were included in this study. Detailed clinical and radiological phenotyping was done and correlated with genotype to identify patterns of clinical presentation and fracture history in this cohort of patients. This data was compared to previously reported literature in this group.

Results: 2 patients with BMP1 and 4 patients with pathogenic variants in C-propeptide region in COL1A1 were deep-phenotyped as part of this study and 1 patient with C-propeptide variant in COL1A1, showed low bone mineral density. In those with an elevated bone mineral density, this became even more apparent on bisphosphonate therapy. Patients in this cohort had variable clinical presentation ranging from antenatal presentation to more of an insidious course resulting in later confirmation of genetic diagnosis up to 19 years of age.

Conclusions: Patients with pathogenic variants in the C-propeptide region of COL1A1/A2 and BMP1 appear to have a high bone mass phenotype with increased sensitivity to bisphosphonate therapy. It is important to closely monitor patients with these genotypes to assess their response to therapy and tailor their treatment regime accordingly.

Introduction

Osteogenesis Imperfecta (OI), also known as brittle bone disease, is an inherited disorder of the connective tissues causing bone fragility and leading to a high fracture risk (Marini et al., 2016). It is a rare disorder, present in 1 in 15–20,000 births (Forlino and Marini, 2016), yet it is the most common form of inherited bone fragility (Marshall et al., 2016).

An initial diagnosis of OI is typically made from a combination of clinical and radiological findings. Key features on presentation include: fractures, growth deficiency, and skeletal deformities, in particular bowing of the long bones (other deformities include: flat midface and triangular facies, chest wall deformities and scoliosis) (Forlino and Marini, 2016). The fractures characteristically present following minimal trauma, in atypical locations and are recurrent (Marom et al., 2020). The fracture risk also involves the increased likelihood of vertebral compressions (Marini et al., 2017). The severity of the disease can range greatly, from mild forms with very few or no fractures, to very severe forms presenting with in utero fractures and perinatal death (Sangsin et al., 2017).

There are also common secondary features, which include: blue
sclerae, relative macrocephaly, dentinogenesis imperfecta (DI), and hearing loss. It has been noted that these secondary features are commonly absent in the recessive forms of OI (Forlino and Marini, 2016). Other potential features include short stature, joint laxity (Sangsin et al., 2017), basal invagination, and cardiac/pulmonary problems (Marini et al., 2017).

The revised Sillence Classification recognises OI types I to V (Van Dijk and Sillence, 2014). This describes the different presentations caused by variants in the COL1A1 and COL1A2 genes, the IFTM5 gene, and some of the recessive forms of the condition (Barnes et al., 2019). This classification has since been adapted to include OI types V-XX of OI which are associated with variants in other genes. Variants in the BMP1, as seen in this cohort, cause Type XII OI (Marini and Dang Do, 2020).

Type I OI is caused by a null allele in COL1A1 (Barnes et al., 2019) and presents with mild features, with infants typically fractures at the time they begin mobilising, but these decrease in frequency after puberty. They may have some bowing of the long bones, and commonly have blue sclerae. Type II OI is lethal in the perinatal period, commonly presenting with many skeletal deformities in utero. Type III OI (progressive deforming type) is the most severe survivable form, patients may sustain hundreds of fractures throughout their life mainly following minimal trauma. Other common features in type III OI include dentinogenesis imperfecta (DI), chest wall abnormalities and severe scoliosis. Type IV OI presents with mild to moderate bone deformity and patients typically have white sclerae although they may have been lightly blue at birth. Variants in BMP1 cause a moderate to severe presentation and patients often have early recurrent fractures and possible long bone bowing, despite having a high bone mass (Marini and Dang Do, 2020).

The most common genes associated with OI are those which encode Type 1 collagen (COL1A1 and COL1A2) (S克斯 et al., 2015); however, variants in other genes account for up to 15% of cases (Marom et al., 2020). This paper discusses 18 genes (from Online Mendelian Inheritance in Man (OMIM)) with variants associated with OI, including: BMP1, COL1A1, COL1A2, CREB3L1, CRTAP, FKB10, IFTM5, MBTPS2, MESP, P3H1, PIP, SERPINF1, SERPINH1, SP7, SPARC, TENT5A, TMEM38B, WNT1. These genes are associated with different types of OI and have a variety of proposed pathways and mechanisms. Variants in COL1A1, COL1A2 and IFTM5 have an autosomal dominant pattern of inheritance. Variants in other genes, including BMP1, are responsible for autosomal recessive forms of OI, except for MBTPS2 which has an X-linked recessive inheritance (Marom et al., 2020).

Type 1 collagen is made up of two alpha1 chains (encoded for by COL1A1) and one alpha2 chain (encoded for by COL1A2). As these genes are involved in the synthesis, structure and assembly of type 1 collagen, variants can decrease the production (e.g., type I OI) or alter the structure (e.g., types II to IV OI) (Marini and Dang Do, 2020). Research has shown that ~6.5% of patients with OI have variants in the C-propeptide (carboxyl terminal propeptide) of type 1 procollagen (Barnes et al., 2019). The C-propeptide is important during the synthesis of procollagen and is cleaved extracellularly before the fibril assembly of mature collagen (Barnes et al., 2019). A high bone mass has been associated with variants involved with the cleavage of C-propeptide (Rolvien et al., 2018). BMP1 is involved in the processing of type I collagen, cleaving the C-propeptide moiety; variants in this gene have been associated high bone mineral density (Marini and Dang Do, 2020).

A high bone mass form of OI could require a different management approach due to the paradigm of clinical presentation. Bisphosphonate treatment is the standard treatment for patients with OI and is known to increase bone mineral density. However, in the context of high bone mass, it is difficult to be precise regarding dosing of bisphosphonate treatment. In addition, surgical interventions may be complicated by the quality of bone and risk of further fractures thereafter. It is also possible that background genetic variants can have an influence on certain OI phenotypes, especially with the dominant C-propeptide cleavage variants.

This paper describes the clinical data of patients with variants in the BMP1 and C-propeptide of COL1A1/A2 genes, with a focus on those with OI associated with high bone mass.

2. Materials and methods

2.1. Clinical information

A cohort of six patients with OI and variants in BMP1 or COL1A1/A2 was examined in this study. With their informed consent (and their parent’s where necessary), relevant clinical data were extracted from the patients’ medical records to put together a history and phenotype.

2.2. DNA extraction

Standard extraction methods were used to collect 2 to 5 ml of peripheral blood from the patients and their parents. In one instance, a blood sample was also extracted from the umbilical cord. The QiAmp DNA Blood Midi kit (Qiagen, Venlo, The Netherlands) was used to extract the DNA from these samples.

2.3. DNA sequencing

The samples where then prepared for analysis using Next Generation Sequencing. Osteogenesis Imperfecta autosomal dominant and autosomal recessive panels were used, containing; NM_006129.5 (BMP1); NM_000942.4 (COL1A1); NM_000918.2 (COL1A2); NM_025854.3 (CREB3L1); NM_006371.4 (CRTAP); NM_001025295.1 (IFITM5); NM_022356.3 (P3H1); NM_000942.4 (PPIB); NM_021939.3 (FKBP10); NM_152860.1 (SP7); NM_002615.4 (SERPINF1); NM_001235.2 (SERPINH1); NM_182943.2 (PLOD2); NM_018112.1 (TMEM38B); NM_005430.3 (WNT1), NM_000918.2 (P4HB); NM_005032.6 (PLS3); NM_014822.2 (SEC24D); NM_003118.3 (SPARC); NM_153365.2 (TAP1); NM_022167.3 (XYLT2).

2.4. Library preparation

The Covaris E220 sonicator was used for the shearing of genomic DNA. The SureSelectCT library system (Agilent Technologies) was used to perform the end repair, A tailing and ligation of adaptors. SureSelect target enrichment (Agilent Technologies) performed target enrichment using custom in house designed probes. Sequencing was performed on the Illumina HiSeq using the HiSeq Rapid SBS Kit v2 performing 2 × 108 base pair paired end reads.

2.5. Data analysis

Data analysis was carried out using the Best Practice Guidelines from Broad Institute.

BWA alignment was used in order to map the reads to the human reference sequence (GRCh37/hg19) (for additional information see http://www.broadinstitute.org/gatk/guide/best-practices).

A read depth of 30-fold was set as a minimum threshold for exonic sequences and intronic sequences up to and including 5 bp from the ends of each exon. A read depth of 18-fold was set as the minimum threshold got intronic sequences from 6 to 25 bp from the ends of each exon. Haplotype Caller (Broad Institute) was used to identify variants and the variants were filtered against this in-house polymorphism list.

2.6. Variant reporting

Once identified, the variants were compared to cDNA reference sequences NM_003118.3 and then assessed with Alamut Visual version 2.11 QT v5.5.1 (Interactive Biosoftware, Rouen, France). The sequence variants were all classified using guidelines for variant interpretation: ACMG/AMP (Richards et al., 2015) and ACGS Best Practice Guidelines for variant classification 2020 (https://www.acgs.uk.com/quality/
3.0 and total body less head at 1.4. She was switched to oral risedronate at 12 years old beginning with 35 mg weekly, reducing to alternate weeks a year later.

At 13 years and 9 months, her lumbar spine BMD z score was +4.3 (previously +4.7) and TBLH was unchanged at +3.1. By this age, she had scoliosis concave to right, believed to be a result of a leg length discrepancy (at age 15, her left leg was 3 cm shorter than the right). She also had increased lumbar lordosis thought to be due to a low sacral kyphosis with a bony shelf. Due to the persisting high BMD despite reducing the bisphosphonate to low levels, it was decided to consider testing for rarer gene mutations causing OI. Fig. 1 shows her BMD trend with age and dense bones on radiographs.

At 14 years, genetic testing identified that she was homozygous for a pathogenic variant in BMP1 and familial testing showed that both her mother and father were carriers of a variant each.

Patient 1 continued to have occasional fractures and at age 14 ½, she fractured her distal left humerus after her wheelchair tipped over. By 17 years old, she had both femurs treated with locked, intramedullary nails and both tibiae had telescopic rods which were somewhat bent. There was some metalwork in the left elbow remaining from a previous fixation. There were hemihipphyesodesis plates in situ in medial distal femora bilaterally and proximal medial tibia on the left.

She was discharged from the bone clinic age 17 and 8 months, taking vitamins but no bisphosphonate treatment since age 14 ½. She was mainly mobilising with a wheelchair but was able to walk short distances if needed. She was seeing a dentist every 6 months, but had no DI.

2.7.2. Patient 2

Patient 2 is a 12-year-old female, the only child of non-consanguineous parents. She was born at 39 weeks following an uncomplicated IVF pregnancy and had a birth weight of 6 lb. 9 oz. She was well at birth and breast fed without any problems. She had one episode of pneumonia at 8 months but was otherwise well. Other than both parents having some degree of hypermobility, there is no other family history of note.

Through her first year of life, she met all developmental milestones. The first concerns did not arise until around 16 months when she was still not walking; however, this followed her mother's development who did not start walking until 18 months, so the parents were not overly concerned. The patient was bottom shuffling at 17 months and, at this time, it was noted that she was very hypermobile. At 21 months, she was still not walking and so was referred to the paediatric clinic. She was found clinically to have bilateral dislocated hips.

She had three operations for her hips, the first of which was at 25 months, and each operation led to at least 6 weeks of cast immobilisation. On her third birthday, she had her first fracture to her fibula and was casted. Within a week, she presented to A&E with a fracture to the left tibia having slipped while leaning against the sofa. Over the next 8 months, she suffered a further 3 fractures to her lower limbs following minimal trauma. Her speech and fine motor development were both normal for her age.

It was initially thought that the fractures were most likely due to osteopenia secondary to long periods of immobilisation; however, she had extensive investigations and was referred to genetics for a possible diagnosis of Ehlers Danlos syndrome due to her joint hypermobility and skin laxity. Treatment with pamidronate was offered, but her parents declined.

Investigations included an MRI head and spine, urine amino-organic acid, acylcarnitine and Prader Willi Screen; all of which were normal. Blood results from FBC, U&E, bone profile, PTH, vitamin D showed that PTH was 0.8 pmol/l and vitamin D was 58 nmol/l but the rest were normal.

As necessary, clinically significant sequence variants were confirmed using Sanger sequencing.

2.7. Clinical reports

2.7.1. Patient 1

Patient 1 is a 19-year-old female, born to non-consanguineous parents. She was born at term following an uncomplicated pregnancy and a normal vaginal delivery. Her birth weight was 2400 g and her length was 62 cm. She is the youngest of three children and both siblings are well. Despite frequent fractures, her general health has been fine. She was investigated several times for possible cardiac problems, including heart murmur and episodes of tachycardia and palpitations, but all tests were negative. She has normal intelligence and attended mainstream school; she reached menarche at age 13.

She presented with a history of multiple fractures, not consistent with injury which had increased in frequency once she became independently mobile. It was also reported that she had hypermobile joints and a blue tinge to her sclerae. Her first fracture was to her right lower leg at 14 months old, and by the time she was 5 years old, she had sustained more than 10 fractures including multiple fractures to both lower limbs. And at age 6, she had bilateral tibial Sheffield rodding.

She arrived in the United Kingdom with her family at the age of 4 and was referred to the paediatric bone disease specialists due to her diagnosis. On examination, she had white sclerae and no evidence of dentinogenesis imperfecta. She did have bilateral tibial bowing and genu valgum with an intermalleolar distance of 10 cm. There was normal bone densitometry, and the spinal films had no major findings, although three vertebrae in the thoracic spine showed a slight decrease in anterior height.

She was started on pamidronate at age 7 (1 mg/kg/day on 3 successive days every 3 months) following a finding of an increasingly deformed crush fracture to the thoracic vertebrae. She was also found to have Grade I spondylolisthesis at the L5/S1 level. Around this time, she had extended investigations and was referred to genetics for a possible diagnosis of Ehlers Danlos syndrome due to her joint hypermobility and skin laxity. Treatment with pamidronate was offered, but her parents declined.

She was diagnosed with type I OI by the Genetics service in Poland. She was started on pamidronate at age 7 (1 mg/kg/day on 3 successive days every 3 months) following a finding of an increasingly deformed crush fracture to the thoracic vertebrae. She was also found to have Grade I spondylolisthesis at the L5/S1 level. Around this time, she had extensive investigations and was referred to genetics for a possible diagnosis of Ehlers Danlos syndrome due to her joint hypermobility and skin laxity. Treatment with pamidronate was offered, but her parents declined.

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2.7.2. Patient 2

Patient 2 is a 12-year-old female, the only child of non-consanguineous parents. She was born at 39 weeks following an uncomplicated IVF pregnancy and had a birth weight of 6 lb. 9 oz. She was well at birth and breast fed without any problems. She had one episode of pneumonia at 8 months but was otherwise well. Other than both parents having some degree of hypermobility, there is no other family history of note.

Through her first year of life, she met all developmental milestones. The first concerns did not arise until around 16 months when she was still not walking; however, this followed her mother's development who did not start walking until 18 months, so the parents were not overly concerned. The patient was bottom shuffling at 17 months and, at this time, it was noted that she was very hypermobile. At 21 months, she was still not walking and so was referred to the paediatric clinic. She was found clinically to have bilateral dislocated hips.

She had three operations for her hips, the first of which was at 25 months, and each operation led to at least 6 weeks of cast immobilisation. On her third birthday, she had her first fracture to her fibula and was casted. Within a week, she presented to A&E with a fracture to the left tibia having slipped while leaning against the sofa. Over the next 8 months, she suffered a further 3 fractures to her lower limbs following minimal trauma. Her speech and fine motor development were both normal for her age.

It was initially thought that the fractures were most likely due to osteopenia secondary to long periods of immobilisation; however, she had extensive investigations and was referred to genetics for a possible diagnosis of Ehlers Danlos syndrome due to her joint hypermobility and skin laxity. Treatment with pamidronate was offered, but her parents declined.

Investigations included an MRI head and spine, urine amino-organic acid, acylcarnitine and Prader Willi Screen; all of which were normal. Blood results from FBC, U&E, bone profile, PTH, vitamin D showed that PTH was 0.8 pmol/l and vitamin D was 58 nmol/l but the rest were normal.
normal. She also had a DXA scan at this time. It showed L2–4 BMD: 0.728 g/cm$^2$ and TBLH BMD: 0.577 g/cm$^2$, the latter scan included numerous metal pins. There was no paediatric reference data but at the time it was thought that the lumbar result was likely high for her age.

At age 4, she was started on pamidronate infusions for about 12 months. After 3 cycles, her BMD increased considerably, and she had chalk stick fractures of the tibiae and fibulae. A clinical diagnosis of osteopetrosis was considered; however, bone biopsy findings did not support this and showed active osteoclasts and osteobone resorption. On examination, she was found to have a high forehead and light blue sclerae. She also had generalised joint hypermobility and her skin was lax over her abdomen.

By age 6 and ½, she had had a total of 16 fractures. She was just above the 50th centile for weight and between the 50th and 75th centiles for height. She had also had dislocations of her left shoulder while swimming, on three separate occasions, all with spontaneous reduction. She was able to walk with a stander and was attending mainstream school regularly, also mobilising with a wheelchair. She was having weekly hydrotherapy sessions.

She was referred to Genetics Clinic at 6 years and 9 months in order to rule out a possible diagnosis of OI. She had previously tested negative for variants in type I collagen genes (COL1A1 and COL1A2), but went onto to have further testing which identified her to be compound heterozygous for variants in BMP1. Subsequently, familial testing showed her parents were carriers of one variant each.

1 year later, she had had only one further fracture. She was, however, still having regular dislocations of her shoulder and hypermobility. She was lax at both large and small joints with skin hyper laxity. She also had increased range of movement in both hips and a leg length discrepancy which led to scoliosis on standing.

Quantitative CT imaging showed very high values in the peripheral skeleton with a radius z score of +8.7 and a trabecular z score at distal radius +9.2. Her lumbur spine BMAD Z score was +4.02 and her lumbar spine QCT Z score was +3.6. Fig. 2 shows her BMD trend with age and dense bones on imaging.

Throughout her education, she attended mainstream school, with one-to-one help for physical education (PE) and later use of writing aids including an angled board and Y shaped pencils as she was tired easily when writing. She was able to walk short distances independently but preferred to wear splints and used her wheelchair for longer distances. By 9 years and 8 months, she had 25$^\circ$ curvature of the spine and had also had recent correction of genu valgum by hemi-epiphysiodesis. She was wearing new AFOs and using a recumbent bike for exercise.

By age 10 ½, she had bilateral intramedullary rods in situ and had not had any fractures or dislocations for 2 years, since before her tibial rods and her most recent DXA scan (6 months previously), showed the lumbar spine Z score to be +1.8. She was taking multivitamins and cholecalciferol, but no other medications. On examination, she had scoliosis concave to the right with the left scapula higher than the right, which was confirmed radiologically with no suggestion of segmentation anomaly, no syrinx, and no focal marrow oedema. Recent hip films showed coxa valga with the right femoral head significantly more
uncovered than the left.

By age 12, she was walking independently again, however her scoliosis was worsening. She was also receiving psychology input due to increasing frustration with her condition. 3 months later, she underwent correction of her scoliosis. She had slow tooth eruption and dental crowding but no DI.

2.7.3. Patient 3

Patient 3 is a 12-year-old male who was diagnosed antenatally with a severe form of OI and it was believed he would not survive for long postnatally. He had 2 older siblings including a half-sister with a history of fractures, and he was born following a dichorionic, diamniotic twin pregnancy with the female twin unaffected. The labour was induced at 27 weeks and both babies were born well by normal vaginal delivery and no resuscitation was required. Patient 3 had a birth weight of 2440 g and was born with multiple fractures of almost every bone in his body.

He was initially given analgesia to help with discomfort and on the day of delivery, he had cast immobilisation applied to four limbs to stabilise fractures for pain relief. Initially he was demand fed but had been vomiting after feeds and so was given an NG tube.

He did not die shortly after birth as expected and instead was transferred from the neonatal unit to a hospice. He continued to be managed with the expectation that he would die; however, a week later, he was still alive. It was decided that a palliative approach was not appropriate for his phenotype and he would benefit from bisphosphonate treatment. He was then transferred to the HDU where he was given 2 infusions of pamidronate on successive days. At 17 days, he had a skeletal survey and clinical phenotyping. He had multiple Wormian bones, but a normal head shape and no mid-face hypoplasia. His skeletal phenotype was reported to be unusual; he had a mixture of wider, slightly cramped femora and thinner distal segments. Despite multiple rib fractures, he did not have any vertebral crush fractures. The high number of fractures was thought to be a possible consequence of sharing the uterine space with an unaffected twin.

He had several health complications over the first few months of life, including ventilation problems meaning he was still requiring oxygen at 9 months of age. He was also found to have a patent foramen ovale although this had spontaneously closed by the age of 5. His feeding was also complicated by gastroesophageal reflux. He was discharged home at 7 months, although this late discharge was largely due to the family finding suitable housing. At 4.5 years old, genetic testing showed he had a de novo variant in the C-propeptide region in \( \text{COL1A1} \).

As a consequence of his diagnosis, his gross motor skills were delayed; he was able to roll from side to side in lying position by age 1 and was crawling on his hands and knees at age 4. While his gross motor skills development was delayed, in all other areas, he was meeting milestones. There were no concerns regarding his intelligence, and he was able to attend mainstream school with 1:1 support.

By age 5.5, he had undergone multiple surgeries including osteotomies and telescopic rodding bilaterally to the femora and tibiae, to attempt to straighten the long bones. By this age he was confident in using a wheelchair for mobilising. He was also wearing glasses due to astigmatism and was under the dentists for a diagnosis of dentinogenesis imperfecta. A year later he was investigated for possible sleep apnoea following reported lengthy pauses in breathing with associated pallor; however, this was found to not be an issue.

His BMD was constantly low, with a DXA scan at age 5.5 showing the lumbar spine BMD to be 0.483 g/cm\(^2\) (Z score of −2.0) and the total body less head to be 0.443 g/cm\(^2\) (Z score of −1.1). This further decreased to −2.2 and −1.8 respectively a year later and at 7 years old, the L2–4 BMD Z score was −2.7 (BMAD Z score −3.3) and the TBLH Z score was −1.6.

At 7 years and 3 months, while in plaster following a recent revision surgery for the tibia rodding, he was able to stand with the support of the
therapist for the first time and was able to take some steps with a walker. 3 months later, he was switched from pamidronate to zoledronic acid (0.04 mg/kg for one day every 6 months) as there were difficulties around cannulation and it was thought he would benefit from fewer infusions. It had also been noted that he had been developing blisters on the tips of his fingers following the treatments. However, the zoledronic acid was found to not be as effective and so he was switched back again shortly afterwards. The dose was later changed to 1.5 mg/kg on one day to minimise the cannulation required.

He began to encounter problems where he had outgrown the previous corrective surgeries. At 8 years of age, he was found to have outgrown the intramedullary wire in his right forearm which had begun

Fig. 3. A. DXA performed at 9 years and 4 months shows a BMD below 2SD of normal for age and sex (this is the only patient in our cohort with low BMD). B. Anteroposterior radiograph of the chest aged 1 day. The bones are slender. There are multiple acute and healing rib fractures, an acute fracture of the left humerus and healing fractures of the right humerus and both clavicles. There were also fractures of the lower limbs (not shown). The acute fractures may have been sustained during the birth process; however, the healing fractures will have been sustained in utero. C. Vertebral fracture assessment (lateral spine DXA) shows multilevel mild and moderate vertebral body collapse of thoracic and lumbar spine.
to irritate the soft tissue, causing him significant pain. At 9 years and 3 months, pain in his left buttock showed a palpable rod which required revision rodding. Just before he turned 10, he was having slight pain over his right proximal humerus and a radiograph confirmed that the telescopic rod had cut out. He subsequently underwent removal of the rod and diaphyseal osteotomy with revision rodding.

Patient 3 also had bone local mechanical properties assessed on a bone specimen mainly of cortical bone, obtained from the upper limb (hardness and elastic modulus) assessed utilizing nano-indentation tests (Pepe et al., 2020) tested in different locations (192 tests within different regions of the specimen). The results showed high heterogeneity in the local material properties of the bone in different locations, with coefficient of variations in the range of 43–48% (Elastic modulus equal to 11.20 ± 5.39 GPa; Hardness equal to 0.49 ± 0.21 GPa). The elastic modulus measured in the bone was 30–40% lower than the average values found in bones extracted from the lower limb or iliac crest of OI type I or type III young patients (3–18 years old) (Albert et al., 2013; Fan et al., 2007). This difference could be due to the different disease, the tested anatomical site and/or the age of the patients. Nevertheless, given absence of normative paediatric literature and analysis of only one sample, it was difficult to draw any firm conclusions from this data.

At 9 ½ years, his DXA scan showed a TBLH BDM of –2.8 (previously –1.5) (Fig. 3). Densitometric vertebral fracture assessment (VFA) showed little change, but possible worsening of shape of L2 and L3 vertebrae and there were still multiple vertebralloss with loss of height. At this age, he was mobilising using a walking frame and his spine was straight with no obvious tenderness.

At 10 years and 3 months, he was noted to have bilateral thyroid nodules however they appeared benign and microcytic and it was decided that there was no need for further investigations unless indicated by a change in biochemistry results. At this age, his BMD was still at the lower range of normal. His L2–4 BMD z score was –2.1 (BMAD –2.0) and his total body less head BMD z score was –0.6 (BMAD –2.6). VFA showed a continued mild-moderate loss of height of several thoracic vertebral bodies, moderate loss of height of L2, and marked loss of height of L3 and L4. His weight was 17.95 kg and his length was 97.5 cm and 102 cm (due to leg length discrepancy). His spine was straight.

At age 11, he had further interventions including trimming and reinsertion of the K wire which was eroding at the elbow following revision rodding of the right ulna. This rod was later removed. He had pain in the left hip due to left migration of the femoral telescopic rod which required readjustment. Recent scans had shown no spondylolisthesis, but an almost parallel sacrum and a further significant bend in the lower sacrum coccyx.

At 11 ½ years old, he had radiological investigations following a possible finding of hyperextension of C2 and C3 on a skull radiograph a year previously. He was also occasionally having pain at the back of the neck, associated with some headaches. It was reported that the tip of the dens was at C2 and he had a wide anterior gap. It was decided that there was no need for neurosurgical intervention at that time.

A month later, he suffered an undisplaced horizontal fracture through his right radius while reaching out for a video game controller and he was placed in a below the elbow soft cast. 12 weeks later, radiograph showed fractures to the distal 1/3 shaft of radius and ulna which were not uniting and causing deformity and required a splint. By 11 years and 10 months, the deformity had worsened although the non-union was painless. His arm was neurovascularly intact and so surgical correction was planned for.

2.7.4. Patient 4

Patient 4 is a 9 year old male born to non-consanguineous parents. He was born at 40 + 4 weeks following an uncomplicated pregnancy and normal vaginal delivery. His birth weight was 3855 g and he had no clinical evidence of fractures. His father had recently been diagnosed with type 1 OI following familial genetic testing due to a diagnosis in his nephew. Therefore, Patient 4 had pre-symptomatic testing of the cord blood at birth which showed him to be heterozygous for a variant in COL1A1, like others in his family including his father and grandfather. Affected members of the family appeared to follow a similar pattern of fractures at birth and then none until age 15.

At 3 days old, the patient was referred with swelling to the right thigh and was found to have a fracture of the right femur, which was confirmed on radiographs and managed conservatively. At 4 weeks old, he attended hospital again with a fracture of the left distal humerus after his left arm had got caught in his vest sleeve. At this time, he also had a skeletal survey which showed an old fracture to the right femur with no callus, a fracture to the right radius with some callus and a fracture to the 6th rib with some callus, as well as two new fractures, including the one to the left humerus and one to the right femur (Fig. 4). He was admitted and placed in gallows traction for 11 days and then a pelvic harness. At 6 weeks, he was started on pamidronate at the standard dose of 0.5 mg/kg.

At 6 months, he had sustained no further fractures. He was now sitting unsupported for short periods, was comfortable pushing up from prone, but was not yet pulling to stand. He was fully weaned and growing well with a weight of 8.8 kg (50th centile) and a length of 69 cm (75th centile). On examination, he had blue sclerae and some flattening of the left side of the occiput. His teeth, that had erupted appeared normal, he had normal shaped facies, normal chest shape and a straight spine with no tenderness. Lateral spine films showed minor loss of height in two of the thoracic vertebrae and he had a mild degree of bowing of both femora, but no other long bone deformities.

Other than a single skull fracture at 11 months, Patient 4 had no further fractures until he was 2 years old; he fractured his right femur while at soft play and was treated with a fixed rod. He was standing and cruising around furniture at 11 months and by 16 months, his development was age appropriate. A DXA scan at 13 months showed his lumbar spine BMD to be 0.405 g/cm². There was little change in the appearance of the vertebrae and persisting loss of height in two thoracic vertebrae. At 22 months, his spine was straight, he had good muscle bulk and there was no evidence of long bone deformity. A lateral radiograph showed that the appearance of the spine had improved although there was still mild reduction in the vertebral body height in the thoracic vertebrae and L3. There was also spondylolisthesis at L5/S1 but no spondylolisthesis.

At age 2 ½ he was speaking words, but no sentences. He lacked approximately 5° knee extension on the right following the rodding. He also lacked end of range supination at the elbows bilaterally. He was switched to zoledronic acid with a dose of 0.025 mg/kg as a single infusion once every 6 months. This was due to the lower frequency of infusions as there were difficulties with cannulation; at 4 years, this was switched again to oral risedronate 35 mg once fortnightly.

At 4 years old, he had another DXA scan which showed L2–4 BMD to be 0.743 g/cm² (increased from 0.663 g/cm² 10 months previously) and TBLH BMD to be 0.592 g/cm² including a metal rod in the right femur (increased from 0.561 g/cm²). He had undergone removal of fixed rods and telescopic rodding of the right femur 7 months previously. Lateral spine radiography showed no definite loss of height of any vertebral bodies and lateral skull radiography showed no basilar invagination. It was soon noted that his bone density was relatively high for his age and the surgeons had commented that his bones were particularly dense. He also had delayed healing of the osteotomy and it was also noted that the patient’s father had had problems with delayed healing and it had taken 14 surgeries to get his right femur to heal properly. It was confirmed that patient 4 had a paternally inherited COL1A1 C-propeptide variant.

After a repeat DXA scan showed further increase in BMD, the decision was made to discontinue the risedronate. He had had no further fractures having been off risedronate for 3 months following surgery for femoral de-rotation on the right side with plate fixation, and the surgeons again commented that his bone was difficult to drill through due to hardness. It was decided to consider reintroduction of bisphosphonates approaching growth spurt given the family history of femoral
fractures in teenage years. At 5 years, his DXA scan showed another increase in BMD with L2–4 at 0.792 g/cm² and TBLH at 0.674 g/cm². By 5 ½ years old, Patient 4 had had a stress fracture in the right subtrochanteric area with malunion and de-rotation osteotomy subsequently. Medial cortical healing was almost complete after removal of the de-rotation plate, but there was still a lateral cortical defect. He was not in any pain but was walking with a short leg gait with segment deficient being on the contralateral side. On examination, he was short in the left tibia by an estimated 15 mm which accounted for his limp when walking and his right knee had a small, fixed flexion contracture of about 10°. His hip exam was normal and he was able to play football with his father and participate in school PE. His DXA scan showed a slight decrease in BMD; L2–4 was 0.777 g/cm² (Z score +1.8) and TBLH was 0.662 g/cm² (Z score +3.1). VFA showed preservation of vertebral height.

Patient 4 continued to have repeat DXA scans (Fig. 4): At 6 years old the L2–4 and TBLH Z scores were +1.9 and +1.5 and at 7 years old they were +1.9 and +2. He had no further fractures and continued to show L4 and L5 Grade I spondylolysis. There was good disc height, slip angle was acceptable, sacral promontory was reasonable and it was decided to observe for symptoms and risk of slip progression. 1 month later, he underwent revision rodding of the right proximal femur with the plan being to start graded weight bearing 2 months later.

At 8 years and 3 months, he had a recent re-fracture of his right femur with rod in situ. He tripped and fell at school onto his right knee and bent the intramedullary rod. He underwent revision rodding of the femur with corrective osteotomy and plating a few months later and was mobilising with crutches 6 weeks later. By 8 years and 11 months, he was healing well, walking inside the house without support; using crutches for short distances outside the house and his wheelchair for longer distances. His most recent DXA scan showed L2–4 +0.6 (previously +1.9), TBLH +2.0 (previously −0.1). VFA showed persistent loss of height in at least two thoracic and two lumbar vertebrae but there was probable minor improvement in one thoracic vertebra. He had occasional discomfort in his back after prolonged sitting where leaning forward and he had a recent fracture to his toe, but on clinical examination his spine was straight. The decision was made to restart the risedronate at a dose of 35 mg once weekly due to the persistent loss of vertebral height and the fact that the BMD was now decreasing.

2.7.5. Patient 5

Patient 5 is a 10 year old male born following an uncomplicated pregnancy to a 19 year old mother with known OI with an identified COL1A1 variant. She contacted the genetics service at 6–7 weeks pregnant, aware that there would be a 50% chance of passing on the condition and unsure of whether or not to continue with the pregnancy. She was offered a test to find out whether the baby had OI but declined, and decided to continue with the pregnancy.

Patient 5 was born via c-section at 38 weeks. He was healthy at birth and weighed 3400 g. He was active and alert with no external stigmata of OI. Examination at 1 month showed that he had blue sclerae and there was laxity of the small joints, but rest of his examination was normal. 2 months later, he had normal range of active movements although he had mild head lag in the sitting position. His weight was tracking around the 25th centile. By 10 months, he was meeting most developmental milestones. He had some reluctance to crawl but this was put down to spending so much time in a walker. He had had no fractures or dislocations. There was no deformity of long bones, his spine was straight, and a previous lateral spine film showed no crush fractures. He had some flattening at the back of his head and hyperlaxity of joints, especially the...
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Fig. 5.

OI variant.
sclerae and it was thought that he most likely had inherited the familial
wrist, ankle and hips, but had reasonable trunk control. He still had blue
perextension of the knees (20 months later at a dose of 1 mg/kg on each of 3 successive days.
vertebrae having lost some height. He was started on pamidronate a few
other fractures until he was 5 years and 8 months, when he fractured his
left tibia.
At 2½ years old, he was seen to have an increase in BMD of lumbar
spine and a deterioration in vertebral morphology, with at least 3 vertebrae having lost some height. He was started on pamidronate a few
months later at a dose of 1 mg/kg on each of 3 successive days.
By 3½ years old, he was noted to have very pronated feet and hyperextension of the knees (20°), with poor lower limb alignment. His most recent DXA scan showed LS BMD to be 0.531 g/cm² (increased from 0.461 g/cm² 6 months previously) and TBLH BMD to be 0.432 g/cm² (increased from 0.392 g/cm²). Lateral skull radiography showed no basilar invagination. Possible fusion of the posterior elements of C2 and C3 was noted as an incidental finding. Lateral spine radiography showed anterior loss of height in several thoracic vertebrae. He was also complaining of back pain in the weeks leading up to his pamidronate infusion.
At 4 years old, he was able to cycle his bike independently with stabilisers; however, he had some degree of limitation of activity compared to his peers and his general wellbeing had been impacted by frequent tonsillitis and ear infections. He also had bilateral foot calcaneovalgus deformity and was wearing orthotics regularly.
He started mainstream primary school at age 5. VFA showed a continuing decrease in anterior height of 3 or 4 thoracic vertebrae. His lumbar spine z score was +0.9 and his total body less head z score was +2.1. 6 months later, following his 12th cycle of pamidronate, a repeat DXA scan showed his lumbar spine z score had increased to +1.3, but his TBLH had decreased to +1.6. VFA showed a decrease in anterior height

Fig. 5. A. TBLH BMD trend (ages 5 to 10) shows consistently raised BMD.
B. AP pelvis at 2 years and 1 month shows an acute fracture of the left proximal femoral shaft.
2.7.6. Patient 6

Patient 6 is the 30-year-old mother of Patient 5. At age 13, she was found to have a COL1A1 variant along with a clinical diagnosis of OI. There is also a more significant family history of OI. It was understood that there was a history of fractures on her father’s side involving her father, aunt and cousin, but there was limited information on this.

Unfortunately, there was not much information available on her early history and presentation of OI. She has had many fractures in her life and while her eldest son (Patient 5) was growing up, she frequently encountered problems regarding her ability to carry him and push his buggy due her own diagnosis.

At 10 months, it was noted that she had congenital musculoskeletal anomalies, but these were not otherwise specified. At 4 years and 5 months, she fractured her tibia and fibula, and 6 months later she fractured her radius and ulna. Since then, she has suffered multiple fractures to bones in her foot including recurrent fractures to her right and left 5th metatarsals.

She had a routine echocardiogram just before she turned 29 due to a perceived risk of valve and aortic root diseases associated with OI but this was normal, and she was scheduled for a repeat in 5 years-time. She was being administered pamidronate from age 10 to 18 and stopped when she found out that she was pregnant. When she was 19 ½ years old her son (Patient 5) was born following an elective caesarean section. Her most recent fracture was 2 years previously to the 5th metatarsal. The most recent imaging of her lumbar spine (at 18 years and 10 months) showed no crush fracturing deformity, spondylosis or spondylolisthesis below T10. She had suffered with severe back pain for a long time and found that this was much worse following the lumbar puncture she had for the caesarean section.

Over the following years, she continued to have severe back pain requiring analgesia. At 20 years old, she was noted to have no spinal tenderness, but had exaggerated lordosis and the movement of her lumbar spine was limited. She had no reduction in the height of vertebrae and her BMD was stable (Fig. 6). At 21 years, her lumbar spine and total hip scores were 0.8 and 1.4, which had showed no significant change from the results 2 years previously. She had a long-standing problem of not consuming enough dairy in her diet and so was regularly taking calcium and vitamin D supplements since the birth of her first son.

She suffered another fracture to her left 5th metatarsal at age 22 after stumbling; she decided that she wanted to restart bisphosphonates and was started on zoledronate at 5 mg. This was not continued as it only

![Fig. 6. A. TBLH BMD trend (ages 9 to 18) shows consistently normal BMD, trending towards the upper end of the normal range with increasing age. B. Lateral spine radiograph aged 18 years and 9 months demonstrates mild loss of height of mid thoracic vertebral bodies and dense vertebral end plates due to bisphosphonate therapy.](image-url)
improved her symptoms for a few weeks. She later decided not to start on other bisphosphonates as was having problems finding a method of contraception that worked for her and did not want to risk getting pregnant while on bisphosphonates.

At 27 years of age, she gave birth to her second son following a caesarean section. He was well at birth with no signs of OI and was found to be negative for the familial variant for OI. In both cases, she found that her back pain increased following the pregnancy. 10 weeks after delivery, her lumbar spine and total hip z scores were 1.0 and 1.8 respectively, which showed no significant change from the results three years previously.

By age 30, she continued to have frequent generalised aches and pains (which was improving following regular physiotherapy exercises), as well has problems with her left knee when kneeling, and her left foot around the site of previous stress fractures. Her most recent lumbar spine and total hip scores were 0.8 and 1.7 respectively, which again showed no significant change from 18 months previously.

3. Results

3.1. Patient 1

Genetic testing identified Patient 1 to be homozygous for the c.355C>T missense variant, predicted to result in a p.Arg119Trp amino acid change in exon 3 of BMP1. Further testing showed that both her mother and father were heterozygous for the c.355C>T variant.

3.2. Patient 2

Genetic testing showed Patient 2 to be compound heterozygous for variants in BMP1. She had a c.1148G>A missense variant in exon 9 predicted to result in the p.Arg383Gln protein change and a c.1293C>G nonsense variant in exon 10 predicted to result in the replacement of the tyrosine at position 431 with a premature termination codon. These mutations had not previously been reported but other mutations in this gene were known to be associated with OI.

Subsequently, her parents were also tested for the variants. Her father was found to be heterozygous for the c.1148G>A variant and her mother was found to be a heterozygous for c.1293C>G variant.

3.3. Patient 3

Patient 3 was found to be heterozygous for a c.4160C>T variant in the C-propeptide region of COL1A1 predicted to result in a p.Ala1387Val missense amino acid change. This variant had been previously reported in one family with a recurrence of type IIC OI although the functional effect was undetermined and in silico analysis did not support the pathogenicity. Further familial testing showed that this variant to be de novo.

3.4. Patient 4

Pre-symptomatic testing of the cord blood, showed Patient 4 to be heterozygous for the familial c.4343G>A, p.Gly1448Asp pathogenic missense variant in exon 51 of COL1A1.

3.5. Patient 5

Patient 5 was found to be heterozygous for a c.3584delG deletion variant, predicted to result in a p.Gly1195fs frameshift in COL1A1. This is the same variant found in his mother (Patient 6).

3.6. Patient 6

Patient 6 was found to be heterozygous for a c.3584delG deletion variant located in the C terminal telopeptide domain of COL1A1, not previously reported in literature. It was predicted to generate a shift in the reading frame (resulting in a p.Gly1195fs) and produce premature termination codon 44 amino acids downstream. Similar variants have been associated with OI of variable phenotype.

4. Discussion

COL1A1 and COL1A2 genes (located on chromosomes 17q21.3 and 7q21.3 respectively (OMIM) encode for three peptide chains: two proα1 chain and one proα2 chain. Together, these make up the type I procollagen molecule, which is the precursor for type I collagen. Each of the chains has a central triple helix and they combine into a trimeric structure (Cundy et al., 2018). The amino (N) and carboxyl (C) terminal propeptides are then cleaved and the collagen molecules are arranged into fibrils (Sangsin et al., 2017). The BMP1 gene (found on chromosome 8p21.3) encodes BMP1 and mTLD which are two of the four proteins involved in the cleavage of C-propeptide from procollagen (the others are mTLL1 and mTLL2) of which, BMP1 was found to have the greatest cleavage activity (Sangsin et al., 2017). In addition, it also affects regulation of TGFβ and activins which impact osteostegenesis. Inactivating variants in BMP1 have been shown to decrease the levels of both BMP1 and mTLD and therefore impairing the cleavage of the C-propeptide (Cundy et al., 2018).

It has been explained that the type I procollagen C-propeptides are involved in both the intracellular procollagen assembly and the extracellular assembly of collagen fibrils (Symoens et al., 2014). It is needed for the regulation of bone mineralisation (Syx et al., 2015) and is involved in regulating the expression of collagen genes via negative feedback (Barnes et al., 2019). One study found that where there was C-propeptide retention, the structure of type 1 collagen fibrils was less organized (Syx et al., 2015). The C-propeptide domains are also important in the selection and association of proc chains (Symoens et al., 2014).

Variants in the collagen I genes (COL1A1 and COL1A2) are the most common in OI (Marshall et al., 2016). With the most common variant being glycine substitutions in the helical domain (Marshall et al., 2016). This variant is known to cause problems with the structure of collagen and has been associated with the more severe forms of OI. The other main variants are nonsense mutation which causes there to be less normal collagen (Hoyer-Kuhn et al., 2015). C-propeptide variants account for only ~5% (as of 2014) of the variants that have been identified in type I collagen (Symoens et al., 2014).

Variants in the COL1A1/A2 genes have been shown to lead to varying phenotypes. These have been classified into OI types I–IV and these vary from lethal in the neonatal period (type II) to more mild forms (types I and IV) (Marini and Dang Do, 2020). The phenotypes of patients with a C-propeptide variant in these genes can also be very varied, which is shown in the patient descriptions of this cohort. This is also supported in the literature, with examples of mild to lethal types of OI in patients with heterozygous variants in the C-propeptide regions (Lindahl et al., 2011). There are also examples of patients with variants in the type I procollagen C-propeptide domain or in the C-propeptide cleavage sites of the proα1 and α2 chains who have high bone mass (Syx et al., 2015).

The cohort reported here includes descriptions of 4 patients from 3 different families with C-propeptide variants in the COL1A1 and 2 patients with BMP1 variants. Table 1 describes the clinical phenotype of BMP1 patients in this cohort in comparison to published literature. Table 2 describes the cohort of patients here with COL1A1 variants in comparison to previously published literature on C-propeptide variants in association with high bone mass while Table 3 gives an overview of their genotype and variant classification based on ACMG criteria (Richards et al., 2015; ACGS Best Practice Guidelines for variant classification 2020 https://www.acgs.uk.com QUALITY/best-practice-guidelines/).

The severity of their phenotypes was very different from case to case. Patient 3, who was de novo heterozygous for variant in COL1A1, had a
| Table 1  | Clinical phenotype in patients with BMP1 variants and comparison to published literature. |
|----------|-------------------------------------------------------------------------------------------------|
| Patient 1 | Family 1 |
| Source   | This cohort (Sangsin et al., 2017)                                                             |
| Age (at time of publication) | 19 | 12 | 6 | 14 | 5.2 | 43 | 31 | 47 | 45 |
| Sex      | F | F | M | M | F | M | M | M | M |
| Ethnicity | Polish | Northern European | Thai | Asian | Asian | Caucasian | Portuguese | Scottish | Scottish |
| Consanguinity | No | No | No | Yes | Yes | No | No | No | No |
| Family history | None | Parents have hypermobility | Sister (patient C) | Brother (patient B) | no | Younger brother | Brother (patient G) | Brother (patient F) | |
| Pregnancy | Uncomplicated IVF | Uncomplicated IVF | Data unavailable | Data unavailable | Data unavailable | Data unavailable | Data unavailable | Data unavailable | Data unavailable |
| Gestation | Term 39 weeks | Term 37 weeks | 36 weeks | Full term | Data unavailable | Data unavailable | Data unavailable | Data unavailable | Data unavailable |
| Birth weight | 2400 g | 2900 g | 3600 g | 2500 g | 2420 g | 3650 g | Data unavailable | Data unavailable | Data unavailable |
| 1st fracture | 14 months Right lower leg | 5 months Right arm | 12 months Right wrist | 7 months Bilateral ulnar and radius | 8 months Femur | Birth | Long bone fractures shortly after walking | Long bone fractures shortly after walking | |
| Bisphosphonates started (age) | 7 years | 4 years | 8 months | –6.5 years | 3 years | None | Not reported | Not reported | Not reported |
| Genotype | c.355C>T | c.1484G>A | c.1484G>A | c.2188dupC | c.925deG | c.34G>C | c.34G>C | c.34G>C | c.34G>C |
| Protein Change | p.Arg191Trp | p.Arg383Gln & p.Tyr431* | p.Phe666Argfs+25 | p.Gln730Profs*294 | p.Assp09Thrfs*54 | p.Gly12Arg | p.Asp614Thrfs*188 | p.Gln730Profs*294 | |
| Homozygous/compound heterozygous | Homozygous | Compound heterozygous | Compound heterozygous | Homozygous | Homozygous | Composite heterozygous | Composite heterozygous | |
| Cognitive development | Normal | Normal | Normal | Not reported | Normal | Normal | Not reported | Not reported | Not reported |
| Motor development | Normal | Delayed – not walking at 21 months | Early development was normal | Not reported | Normal | Delayed | Not reported | Not reported | Not reported |
| Motor skills | Mainly using wheelchair, but can walk for short distances | Able to walk independently | Able to walk until 6 years | Not reported | Never able to walk. | Wheelchair dependent until around 14 years | Able to walk very short distances supported | |
| Age matched DXA Z score (pre-bisphosphonates) | None measured | 0.728 g/cm² (aged 4 years, no Z score available) | –0.9 (aged 6 years) | Not reported | Not reported | Not reported | Not reported | Not reported | Not reported |
| Age matched DXA Z score (post-bisphosphonates) | +4.3 (aged 13 years 9 months) | +4.02 (aged 7 years 9 months) | 0.3 (aged 14 years) | 0.95 (BMAD aged 2 years) | No bisphosphonates | Not reported | Not reported | Not reported | Not reported |
| Sclerae | Blue then white | Light blue then white | White | Grey | Blue then white | White | White | White | White |
| Teeth | No DI | Light blue then white Dental crowding and slow tooth eruption. | White | White | Blue then white | Opalescent teeth | White | White | Not reported |
| Hearing | Normal | Normal | Normal | Normal | Normal | Normal | Normal | Normal | Normal |
| Vision | No problems | No problems | No problems | No problems | No problems | No problems | No problems | No problems | No problems |
| Hypermobility | ++ | ++ | ++ | ++ | ++ | ++ | ++ | ++ | ++ |
| Scoliosis | Concave to right due to leg length discrepancy – age 13 | Lumbar lordosis | No reported | Compression fracture | No vertebral compression fractures | Multiple vertebral fractures | Multiple vertebral fractures | Not reported | Not reported |
| Patient | 14 | 15 | 16 | 17 | 18 | 19 | 20 | 21 | 22 | 23 | 24 | 25 |
|---------|----|----|----|----|----|----|----|----|----|----|----|----|
| Family  | 11 | 12 | 13 | 14 | 15 | 16 | 17 | 18 | 19 | 20 | 21 | 22 |
| Source  | (Xu et al., 2019) | (Cho et al., 2015) | (Choksi et al., 2021) | (Asharani et al., 2012) | (Martínez-Glez et al., 2012) | (Valencia et al., 2014) | (Fahiminiya et al., 2015) |
| Age (at time of publication) | 15 | 7 | 11 | 7 | 15 | 5 | 2.8 | 17.4 | 28.8 | 22.6 | 8.1 |
| Sex | M | F | M | M | F | F | M | M | F | F | M | F |
| Ethnicity | Chinese | Korean | Hispanic | Turkish | F | Turkish | Egyptian | M | M | Pakistani | F | French Canadian | M | F | French Canadian |
| Consanguinity | No | No | No | Yes | Yes | Yes | Yes | Yes | No | No | No | No |
| Family history | No | No | No | Sister (Patient 18) | Yes | Brother (Patient 17) | Yes | Brother (Patient 20) | Yes | No | No | No |
| Pregnancy | Not reported | Not reported | Uncomplicated | Not reported | Not reported | Femoral bowing detected intrauterine | Unremarkable | Not reported | Not reported | Not reported | Not reported |
| Gestation Birth weight | Full term | 3300 g | 22 months left femur | NVD at 37 weeks | 3400 g | Immediately after birth left humerus | Full term | Not reported | 2.5 years right tibia | Not reported | Not reported | Full term |
| 1st fracture | Not reported | 23 months | Not reported | Not reported | 14 months | NVD full term | Not reported | Immediately after birth left tibia | Not reported | Not reported | Not reported |
| Bisphosphonates started (age) | Treated for 12 months. Not reported when. | None | 5.4 years | 2.8 years | Not reported | Not reported | 2 years | None | None | 9.6 years | 4 years |
| Genotype | c.1324G>T | c.808A>G | c.34G>C | c.747C>G | c.148+1G>A | c.808A>G | c.34G>C | c.747C>G | c.1297G>T | c.C505T | c.34G>C |
| Protein change | p.Asp442Tyr | p.Arg169Cys | p.Gly12Arg | p.Phe249Leu | p.Phe249Leu | p.Phe249Leu | p.Gly12Arg | p.Phe249Leu |
| Homo/heterozygous | Compound heterozygous | Homozygous | Homozygous | Homozygous | Homozygous | Homozygous | Homozygous | Homozygous |
| Cognitive development | Not reported | Normal | Mild speech delay | Moderately delayed | Normal intelligence | Not reported | Delayed | Delayed | Delayed | Not reported | Not reported |
| Motor development | Sit at 7 months | Not reported | Not reported | Not reported | Not reported | Not able to sit unsupported | Not reported | Not able to stand unsupported | Not able to stand unsupported | Not reported |
| Motor skills | Not reported | Not reported | Not reported | Not reported | Not reported | Not able to stand unsupported | Not reported | Not reported | Not reported | Not reported | Not reported |
| Age matched DXA Z score (pre-bisphosphonates) | +2.1 (aged 15.7 years) | +4.6 (aged 16.7 years) | +1.4 (aged 18 years) | Total hip 1.76 (aged 7 years) | High above normal | High above normal | –2.13 (aged 15 years) | Normal | –0.4 (aged 11.5 years) | 1.7 (aged 5.7 years) | 0.0 (aged 9.4 years) |
| Age matched DXA Z score (post-bisphosphonates) | +2.1 (aged 15.7 years) | +4.6 (aged 16.7 years) | +1.4 (aged 18 years) | Increased by 27% following bisphosphonate treatment | Increased further | Increased further | Increased further | No bisphosphonates | No bisphosphonates | 1.1 (aged 22.6 years) | 0.1 (aged 8.1 years) |
| Sclerae | White | No DI | No DI | White | Blue-grey | White | Blue | Blue | White | White | White |
| Teeth | White | No DI | No DI | White | No DI | No DI | No DI | No DI | No DI | No DI | No DI |
| Hearing | Normal | Not reported | Not reported | Normal | Normal | Normal | Normal | Normal | Not reported | Not reported | Not reported |
| Vision | Not reported | Hyperlaxity | Elbow, wrist, interphalangeal joints | Not reported | No | Not reported | Not reported | Not reported | Not reported | Not reported | Not reported |
| Hypermobility | Slightly hypermobile | Hyperlaxity | Slightly hypermobile | No | No | No | No | No | No | No | No |

(continued on next page)
| Patient | Family | Scoliosis | Vertebral compression fractures | Thoracic spine |
|---------|--------|-----------|---------------------------------|----------------|
| 14      | 11     | MILD      | Not reported                    |                |
| 15      | 12     | MILD      | Not reported                    |                |
| 16      | 13     | MILD      | Not reported                    |                |
| 17      | 14     | MILD      | Not reported                    |                |
| 18      | 15     | MILD      | Not reported                    |                |
| 19      | 16     | MILD      | Not reported                    |                |
| 20      | 17     | MILD      | Not reported                    |                |
| 21      | 18     | MILD      | Not reported                    |                |
| 22      | 19     | MILD      | Not reported                    |                |
| 23      | 20     | MILD      | Not reported                    |                |
| 24      | 21     | MILD      | Not reported                    |                |
| 25      | 22     | MILD      | Not reported                    |                |

**Table 1 (continued)**

| Patient | Family | Scoliosis | Vertebral compression fractures | Thoracic spine |
|---------|--------|-----------|---------------------------------|----------------|
| 14      | 11     | MILD      | Not reported                    |                |
| 15      | 12     | MILD      | Not reported                    |                |
| 16      | 13     | MILD      | Not reported                    |                |
| 17      | 14     | MILD      | Not reported                    |                |
| 18      | 15     | MILD      | Not reported                    |                |
| 19      | 16     | MILD      | Not reported                    |                |
| 20      | 17     | MILD      | Not reported                    |                |
| 21      | 18     | MILD      | Not reported                    |                |
| 22      | 19     | MILD      | Not reported                    |                |
| 23      | 20     | MILD      | Not reported                    |                |
| 24      | 21     | MILD      | Not reported                    |                |
| 25      | 22     | MILD      | Not reported                    |                |

very severe form of the disease, initially being diagnosed with the lethal type antenatally. He has continued to have fractures and problems with deformities throughout his life. He was also the only patient from this cohort to have a significantly low BMD. In contrast to this, patients 5 and 6 (son and mother respectively) had much milder forms of the condition. Bone biopsies were taken from patient 3 (Pepe et al., 2018) and nanoindentation was performed. The study was limited as the exact location that the specimen was extracted from was not comparable to published normative paediatric literature. The results showed that the elastic modulus and hardness of the bone specimen were lower than those in healthy adult bone and the elastic modulus was lower than results from other paediatric patients with type III and type I OI. The results also showed that the mechanical properties were very heterogeneous across different regions of the specimen, and one side of the bone specimen showed a larger proportion of denser cortical bone compared to the other side. Further bone biopsy samples from this group were difficult to obtain, limiting the possibility of generalising the findings. Nevertheless, future analysis on a larger sample size should shed more light into bone material properties in this group of patients with OI. In addition, evidence of enhanced osteoclastic activity through analyses of urinary collagen derived crosslinks and studying osteoclastic activity in bone fragments from surgical bone tissue may provide further data on those patients that might benefit from bisphosphonate therapy and those that may become hypermineralised.

Patients 4 and 5 were both found to have significantly raised BMD. In patient 4 this was significant enough to be remarked upon by the operating surgeons who were finding that the bones were particularly difficult to drill in to due to their high density. While patients 5 and 6 shared the same familial variant, Patient 6 did not have a high BMD, instead it was consistently normal.

The phenotypes of patients with mutations in the BMP1 gene have been described as high mineral bone density associated bone fragility (Pollitt et al., 2016), but the phenotypes do differ in the literature. It has been noted that some patients with milder phenotypes had a homozygous variant (c.*241T>C) which only affected the transcription of the BMP1 protein, leaving the mTLD protein functional. There were also milder phenotypes demonstrated in patients with the p.Gly12Arg signal peptide variants and 2, out of the 3 patients with this variant, had high bone mass. This was thought to be due to a higher mineral incorporation in the bone matrix following a delay in bone mineralisation (Syx et al., 2015). Biopsies that were carried out from patients with BMP1 variants and high BMD demonstrated that there was a simultaneous excess of osteoid (hyperostoideosis) and increased mineralisation (hypermineralisation). This was the same as was seen in those with variants in the C-propeptide cleavage site (Cundy et al., 2018).

This cohort of patients included 2 patients with variants in the BMP1 gene. Despite differing initial presentations, where patient 2 had a fairly complicated path to diagnosis, they both did have many similarities. They both began getting fractures once they were independently mobilising, although in patient 2 this did not occur until later largely due to problems with hip dislocation and hypermobility. They also both began with light blue sclerae which became white with age and had notable scoliosis when older. Both had increased BMD; however, this was a more significant finding in Patient 2 who was investigated for osteopetrosis early on in her presentation.

Bisphosphonates are the main pharmacological treatment used in children with OI. They work by inhibiting the activity of osteoclasts and inducing apoptosis. They have been shown to increase bone mass and improve the internal structure of bone tissue, thereby helping to reduce the incidence of fractures (Rossi et al., 2019). There is some suggestion of delayed healing following surgery in patients having bisphosphonate therapy (Marom et al., 2020). Studies have shown that along with reducing the rate of fractures, it increases the bone mineral density (Bradbury et al., 2012); but does not alter bone mineralisation density distribution i.e. tissue bone density (Boyd et al., 1999).

All the patients in this cohort were started on intravenous
| Patient | 3 | 4 | 5 | 6 | 26 | 27 | 28 | 29 | 30 | 31 | 32 |
|---------|---|---|---|---|----|----|----|----|----|----|----|
| Family  | 3 | 4 | 5 | 5 | 21 | 22 | 23 | 24 | 24 | 25 | 25 |
| Source  | This cohort | (Lindahl et al., 2011) | (Mcinerney-Leo et al., 2015) | (Rolvien et al., 2018) | (Le Quesne Stabej et al., 2017) |
| Age (at time of publication) | 12 | 9 | 10 | 30 | 13.5 | 14 | Not reported | 80 | 56 | Not reported | Not reported |
| Sex | M | M | M | F | F | M | F | F | F | F | M |
| Ethnicity | Caucasian | Caucasian | Caucasian | Caucasian | Swedish | Caucasian | American | Not reported | Not reported | Not reported | Pakistani |
| Consanguinity | No | No | No | No | Not reported | No | Not reported | Son (6) | No | Not reported | Yes |
| Family history | History of #s in half sister | Father (6) | Son (5) | Paternal side | Possibly on paternal side | Data unavailable | Full term NVD | Data unavailable | 3.5 years right tibia and fibula | Not reported | Yes |
| Pregnancy | Diagnosed antenatally, twin pregnancy | Uncomplicated | Uncomplicated | Data unavailable | Full term NVD | Data unavailable | Full term NVD | Data unavailable | 3 years right tibia and fibula | Not reported | Data unavailable |
| Gestation | 36 weeks | 2440 g | 3400 g | Data unavailable | 2950 g | 3690 g | Not reported | Data unavailable | 3.5 years right tibia and fibula | Not reported | Data unavailable |
| 1st fracture | In utero | 38 weeks | 2 years 2 months | Left femur | Data unavailable | Full term NVD | Data unavailable | Data unavailable | Full term NVD | Early childhood | Data unavailable |
| Bisphosphonates started (age) | 6 weeks–5 years | Restated at 8 years | 11 months | 2.5–3 years | None | 6 years | 10–18 years | 2 years | 10–18 years | None | Data unavailable |
| Gene | COL1A1 | COL1A1 | COL1A1 | COL1A1 | COL1A1 | COL1A1 | COL1A2 | COL1A1 | COL1A2 | COL1A1 | COL1A1 |
| Genotype | c.4160C>T | c.4343G>A | c.4343G>A | c.3584delG | c.3584delG | c.3584delG | c.3655G>A | c.3655G>A | c.3655G>A | c.3655G>A | c.3655G>A |
| Protein change | No | Heterozygous | Heterozygous | Heterozygous | Heterozygous | Heterozygous | Heterozygous | Heterozygous | Heterozygous | Heterozygous | Heterozygous |
| Cognitive development | Normal | Normal | Normal | Normal | Normal | Normal | Normal | Normal | Normal | Normal | Normal |
| Motor development | Very delayed | Age appropriate by 16 m | Data unavailable | Normal | Walking independently | Not reported | Not reported | Not reported | Not reported | Not reported | Not reported |
| Motor skills | Using a walking frame at 9 ½ years | Delayed Walking at 21 m | Data unavailable | Normal | Not reported | Not reported | Not reported | Not reported | Not reported | Not reported | Not reported |
| Age match DXA Z-score (pre-bisphosphonates) | None measured | None measured | Data unavailable | +3.9 (aged 12.5 years) | Not reported | 7.7 (aged 28) | Not reported | Not reported | Not reported | BMAD 0.45 (aged 5 years) | Data not available |
| Age matched DXA Z-score (post-bisphosphonates) | −2.1 (aged 10 years 3 months) | +0.6 (aged 8 years 11 months) | +2.0 (aged 10 years) | LS 0.8 (aged 30 years) | No bisphosphonates | 0.00 (aged 12 years) | Not reported | Not reported | Hip +3.9 | Spine normal | Data not available |
| Sclerae | Grey | Blue | Blue | Blue | White | White | White | White | White | White | Data not available |
| Teeth | Dentinogenesis imperfecta | No DI | Normal | Data unavailable | Light grey | No DI | Delayed eruption | Normal | Normal | Normal | Data not available |
| Hearing | Normal | Normal | Normal | Normal | Not reported | Normal | Not reported | Normal | Non-syndromic sensorineural hearing loss | Normal | Data not available |
| Vision | Normal | Normal | Normal | Normal | Not reported | Not reported | Not reported | Not reported | Not reported | Not reported | Data not available |

(continued on next page)
Table 2 (continued)

| Family | Glasses for hypermobility | Hypermobile | Scoliosis | Vertebral fractures | Genotype of reported patients in this cohort. |
|--------|--------------------------|------------|-----------|--------------------|-----------------------------------------------|
| 3      |                          |            |           |                    |                                               |
| 4      |                          | Hypermobile |           |                    |                                               |
| 5      |                          | Hypermobile |           |                    |                                               |
| 6      |                          | Hypermobile |           |                    |                                               |

ACMG criterion applied: PS2: De novo (both maternity and paternity confirmed) in a patient with the disease and no family history, used at strong level; PVS1: null variant (nonsense, frameshift, canonical ±1 or 2 splice sites, initiation codon, single or multiexon deletion) in a gene where LOF is a known mechanism of disease; PM1: Located in a critical functional domain without benign variation, used at moderate level; PM2: Absent from controls in gnomAD database; PM3: Detected in trans with a pathogenic variant; PM3: Multiple lines of computational evidence support a deleterious effect on the gene or gene product, used at supporting level; PP4: Highly specific for a single genetic aetiology; PM3: Detected in trans with a pathogenic variant; PP4: Pathogenic PM2, PVS1 – Class 5.

5. Conclusions

This project aimed to establish whether there was a common phenotype in OI patients with BMP1 or C-propeptide variants, particularly in regards to a high bone mass presentation in order to aid decision making when considering treatment in this cohort of patients.

This cohort shows that 2/2 BMP1 patients and 2/4 C-propeptide patients have a high bone mass phenotype; however, it remains unclear why some patients do not have high bone mass and why the phenotypes can range so greatly.

We show that some patients with these variants have a significant increase in BMD following bisphosphonate treatment that often exceed the upper limit of the normal range. Further work is required in a larger cohort of patients to establish whether this increase would have occurred due to the clinical presentation with high bone mass.
phenotype, regardless of bisphosphonate treatment or whether these genotypes give the patients an increased susceptibility and elevated sensitisation to treatment with bisphosphonates and if this may be agent-specific. A more systematic phenotyping of this group of patients from early clinical presentation with close monitoring of response to treatment is required to draw further conclusions regarding timing of intervention and whether alternate modalities of emerging treatment options for OI needs to be considered in this cohort.

CRediT authorship contribution statement

EC collected patient data and collated all genetic and radiology information; MB supervised data collection and write-up; all authors contributed to patient data collection and approved manuscript.

Declaration of competing interest

The authors declare that they have no known competing financial-interest personal relationships that could have appeared to influence the work reported in this paper.

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Statement

This study has been approved by the South Yorkshire Research Ethics Committee (REC reference: 249448) and appropriate institutional boards and the research has been performed in accordance with the 1964 Helsinki Declaration. All authors contributed to this manuscript.

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