Case Report

Hypophosphatasia: Novel Mutation Associated With An Atypical Newborn Presentation

Short title: Hypophosphatasia: Novel Mutation & Presentation

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What is already known on this topic?
Hypophosphatasia is a rare Mendelian disease affecting bone metabolism by loss-of-function mutations in the gene encoding the tissue non-specific isoenzyme of alkaline phosphatase. To date, 381 different mutations have been described. The clinical expression of the disorder varies widely and can even differ between patients with the same mutation.

What this study adds?
We describe a novel mutation that has not been described in the literature to date. We also present an atypical neonatal presentation of the syndrome, with mild phenotype, different from the classic neonatal form. However, it could also be an early diagnosis of the childhood form, with better prognosis.

Abstract
Hypophosphatasia, a rare genetic disease affecting bone metabolism, is characterized by decreased activity of tissue non-specific alkaline phosphatase (TNAP). The gene encoding TNSP (ALPL) has considerable allelic heterogeneity, which could explain different degrees of enzyme activity determining a wide clinical variability. We report the case of a preterm newborn in whom a corneal opacity was detected at birth. Blood tests performed to investigate this finding showed low alkaline phosphatase concentrations. The corneal opacity disappeared within a week but alkaline phosphatase remained persistently low. With persistently decreased levels of alkaline phosphatase upon suspicion of hypophosphatasia, plain radiography detected changes suggestive of rickets. Sequencing of the ALPL gene revealed a heterozygous variant that has not been described in the literature to date.

Our patient’s condition could be an atypical neonatal form of the syndrome, with a mild phenotype, very different from the classic neonatal form which can lead to severe skeletal disease and respiratory failure. However, it could also be an early diagnosis of the childhood form, with better prognosis.

Keywords: Hypophosphatasia, mutation, newborn, alkaline phosphatase

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Introduction
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Case Report
A late preterm male infant (34+3 weeks) was admitted to the minimal care neonatal unit for prematurity and infection risk, due to delivery 5 days after premature rupture of membranes. Cleft palate had been shown at the second trimester ultrasound study. The patient’s family background included 3 paternal family members with cleft palate. A corneal opacity was detected at birth. On blood testing to investigate this finding at 5 days of life, phosphate value was 8.2 mg/dL (normal value in preterm infants,<7.9 mg/dL) with normal calcium (9.6 mg/dL), vitamin D (25.4 ng/mL) and parathormone hormone (19 pg/mL) but alkaline phosphate was 52 IU/L (normal value, >75 IU/L). At 1 week of life, the corneal opacity had disappeared and the infant was asymptomatic. In subsequent analyses alkaline phosphate remained low, up to a maximum of 67 IU/L with phosphate value under 8 mg/dL. The main potential causes of low alkaline phosphate values—hypothyroidism, anemia, low magnesium or zinc values, vitamin D intoxication, low vitamin C, Wilson’s disease, or intake of certain drugs were ruled out. This led to a possible diagnosis of hypophosphatasia.

Radiographic studies on day 8 of life revealed low radiological density. Subsequent radiography at 2.5 months of life (Figure 1) showed irregularities and widening of the radial and ultrar metaphyses, suggestive of rickets. A periostal reaction in the femur with discrete metaphyseal widening was also observed.

Molecular genetic studies were performed using polymerase chain reaction amplification with specific primers. The sequencing reaction was carried out with the BigDye Terminator v3.1 Sequencing Kit (Applied Biosystems), capillary electrophoresis was performed on an ABI Prism 3730XL DNA Analyzer Sequencer (Applied Biosystems), and chromatograms were generated with the SeqPilot software (JSI Medical Systems). A heterozygous variant, c.1292T>A, was found in exon 11 of the ALPL gene, producing an amino acid change: p.(Val431Asp). This specific mutation, in the gene affected in hypophosphatasia, has not yet been described [5]. Bone densitometry of the lumbar spine was performed. It showed a bone mineral density of 0.359 g/cm², consistent with normal reference values from Spain.

Clinical evaluation of first-degree relatives to investigate possible affected individuals within the family was negative, but the patient’s mother showed low plasma levels of alkaline phosphate (28 IU/L). She did not present stigmas of rickets or early tooth loss and its height was within the limits of normality. TNAP gene analysis was carried out to determine whether the mutation was de novo or inherited, and the same mutation was found in the mother.

Currently, the patient is being monitored with periodic follow-up visits and is not receiving replacement therapy. Two years after the diagnosis the alkaline phosphate values remain low, always inferior to 70 IU/L. Radiographic studies (Figure 2) show signs of diffuse osteopenia with affected distal metaphysis of both ulnas with marked deflection, loss of trabecular density and slight significant widening on the left ulna. The rest of metaphyses are enlarged with only a slight sclerotic line on the physis. There are no other skeletal findings.

Written informed consent was obtained from the patient’s parents for publication of this case report and the accompanying images.

Discussion
Hypophosphatasia is a rare inherited disease characterized by decreased alkaline phosphatase levels due to reductions in the tissue non-specific isoenzyme. The presumptive diagnosis of this disease is based on physical examination and consistent radiographic findings, in association with low concentrations of alkaline phosphatase in blood tests. The molecular diagnosis is established by sequencing the gene encoding TNAP [1].

The casual finding of corneal opacities in this patient was the starting point of the diagnosis, although the opacities disappeared by 1 week of life. An association between hypophosphatasia and ocular signs of hypercalcemia, morphologically identical to band keratopathy, has been reported [6]. However, the fast, spontaneous resolution of this finding suggests that it was not an abnormality related with the genetic disease. This case exemplifies an atypical presentation of the disorder, as the patient was asymptomatic at the time of the diagnosis, but had low alkaline phosphate levels. Physical examination at birth provided no evidence of hypophosphatasia and radiographs only detected non-specific osteopenia. However, in subsequent examinations, the patient had the characteristic decrease in alkaline phosphatase activity and radiographic findings suggestive of hypophosphatasia. Imaging typically shows a generalized decrease in bone density and metaphyseal abnormalities in long bones, similar to those found in severe forms of rickets [1,2]. However, unlike what occurs in rickets, serum alkaline phosphate levels are decreased in hypophosphatasia [1,3].

Reported evidence has indicated that the lower the alkaline phosphate levels, the more severe are the manifestations of the disorder [1,7]. The mutations in severe hypophosphatasia produce a protein that fails to reach the cell membrane, but instead, accumulates in the cis-Golgi apparatus and is then degraded in the proteasome without producing adequate enzymatic activity. However, mutations found in the mild forms produce enzymes that are, in part, properly located at the cell membrane and exhibit significant residual activity [8].

Our patient did not have very low levels of the enzyme, which suggests that the newly identified genetic variant may cause a mild phenotype. It may be an atypical, mild neonatal form, considering the age of presentation and absence of clinical features, despite the presence of radiographic abnormalities. Nonetheless, we cannot exclude that it may represent an early diagnosis of childhood hypophosphatasia, with a better prognosis than the neonatal forms. It cannot be considered as benign prenatal hypophosphatasia, as the diagnosis was made after birth and the patient had no visible abnormalities on prenatal ultrasound studies.

The definitive diagnosis of hypophosphatasia is established by sequencing the TNAP gene, located on chromosome 1p36.1-p34 [1,5], and subject to very strong allelic heterogeneity. To date, 381 different mutations have been described in ALPL, 70.6% of which are missense mutations [5]. Allelic heterogeneity explains the different degrees of enzyme activity and the great variability in the clinical expression of the disease [1,3]. In our case, a previously unidentified, a heterozygous variant in exon 11 of ALPL (c.1292T>A) was detected in both the patient and his mother. This mutation produces an amino acid change: p.(Val431Asp). Bioinformatic studies using SIFT [9] and Polyphen2 algorithms [10] predicted a harmful effect.
of the mutation on the structure or function of the protein. According to the American College of Medical Genetics and Genomics guidelines for the interpretation of sequence variants [11], the mutation would be considered probably pathogenic. As to inheritance, severe forms of hypophosphatasia (perinatal and infantile) have an autosomal recessive transmission, whereas in mild forms, transmission can be recessive or dominant [2]. Sporadic cases are rare. Our patient had a heterozygous mutation, allegedly dominant. The negative effects of some heterozygous mutations, ranging the TNAP activity from 20 to 40% of wild-type, explain the dominant transmission of hypophosphatasia. A mild phenotype is expected at the heterozygous state even if the mutation has a severe effect [1,2].

The dominant negative effect is corroborated being a missense mutation in the localization of Val431. A mutation in the same allele—p.(Val431Ala)—leading to a change from valine to alanine has been described in relation with odontohipphosphatasia [5]. The clinical form of the condition in our patient remains to be defined, but as bone was affected on radiography, it is unlikely to be odontohipphosphatasia.

The severity of the condition can differ between patients showing the same mutation [1], and there is significant variability in the expected clinical features [3]. In our patient, the mutation supposedly had a dominant inheritance; hence, relatives with the mutation might or might not develop mild hypophosphatasia (his mother is still asymptomatic). It could be that this patient and his mother will have low alkaline phosphatase levels and harbor a recessive variant in a TNAP allele but never experience symptoms, in which case they could be considered carriers [1]. The variability in the inheritance models and the variable penetrance complicates genetic counseling, although it could be of value in couples with an affected child [2,3]. Considering that the accumulation of the substrates will be a key point in the future follow-up [1,4], to be able to predict any clinical manifestations of progression of the disease the next step to confirm the phenotype would be the determination of the substrates of the enzyme.

Asfotase alfa, a recombinant human tissue non-specific alkaline phosphatase coupled to a deca-aspartate motif for bone targeting, has demonstrated efficacy for healing the skeletal manifestations of hypophosphatasia. In addition, it has been found to improve respiratory and motor function in perinatal and infantile forms, with a good safety profile [12]. Given the age of the patient, the disease course should be monitored so that management can be adapted to the abnormalities that may occur.

Conclusions

A previously unidentified heterozygous variant located in exon 11 of the ALPL gene—c.1292T>A— is described as a cause of hypophosphatasia. The case reported could be considered an atypical presentation of the neonatal form, with a mild phenotype and different from the classical neonatal form, or the childhood form with a prompt diagnosis, which is associated with a better prognosis. The child’s condition should be closely monitored, and if deemed appropriate, replacement therapy with asfotase alfa can be provided. The persistently low level of alkaline phosphatase should alert the clinicians for HPP. Definite diagnosis can be done by genetic analyse.

Declarations

The authors declare that there are no conflicts of interest regarding the publication of this paper. Written informed consent was obtained from the patient’s parents for publication of this case report and the accompanying images.

References

Figure 1. Radiographs at the time of the hypophosphatasia diagnosis
Figure 2. Radiographs at the age of 14 months