Bone health in patients with inborn errors of metabolism

M. Langeveld¹ • C. E. M. Hollak¹

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Abstract
Inborn errors of metabolism encompass a wide spectrum of disorders, frequently affecting bone. The most important metabolic disorders that primarily influence calcium or phosphate balance, resulting in skeletal pathology, are hypophosphatemic rickets and hypophosphatasia. Conditions involving bone marrow or affecting skeletal growth and development are mainly the lysosomal storage disorders, in particular the mucopolysaccharidoses. In these disorders skeletal abnormalities are often the presenting symptom and early recognition and intervention improves outcome in many of these diseases. Many disorders of intermediary metabolism may impact bone health as well, resulting in higher frequencies of osteopenia and osteoporosis. In these conditions factors contributing to the reduced bone mineralization can be the disorder itself, the strict dietary treatment, reduced physical activity or sunlight exposure and/or early ovarian failure. Awareness of these primary or secondary bone problems amongst physicians treating patients with inborn errors of metabolism is of importance for optimization bone health and recognition of skeletal complications.

Keywords Osteopenia • Osteoporosis • Inborn error of metabolism • Lysosomal storage disorder • Skeletal dysplasia • Bone metabolism

1 Introduction
Bone is frequently affected by inborn errors of metabolism. Some disorders primarily affect bone and present with prominent skeletal features such as the mucopolysaccharidoses, while in other disorders the alterations, such as decreased bone mass, may be secondary to nutritional deficiencies as a consequence of strict diets, inflammatory processes, hypogonadism and/or drugs, especially anti-epileptic treatments [1]. Mirroring the range of pathophysiological mechanisms involved, therapeutic approaches will be different for the wide range of disorders. However, some generalizations can be made: in all conditions, physicians should be aware of general measures to optimize bone health such as adequate intake of calcium/phosphate and vitamin D and optimal physical activity. In patients with cognitive dysfunction, the latter can be very challenging. Early identification and management of potential risk factors is essential for skeletal health in adulthood.

In this article, we discuss primary and secondary involvement of the bone in inborn errors of metabolism. The first group includes those primarily affecting the bone marrow, mineral component or cartilage, in particular the lysosomal storage disorders, hypophosphatasia and hereditary hypophosphatemic rickets (Table 1, Fig. 1). The second group encompasses mainly disorders of aminoacid metabolism such as phenylketonuria, lysinuric protein intolerance and homocysteinuria (Table 2).

Finally, diagnostic strategies and general recommendations are provided.

2 Primary involvement of the skeleton
2.1 Disorders affecting calcium and/or phosphate homeostasis
Calcium and phosphate are essential components of bone and have important roles in several physiological functions as well. For this reason homeostasis is tightly regulated involving not only the bone but also intestine and kidney. Mutations in several key regulatory genes may affect parathyroid hormone function, calcium sensing or phosphate transporters resulting in a variety of clinical symptoms [2]. The following section will focus be on inherited metabolic bone disorders
with an overt skeletal phenotype, i.e. hypophosphatasia and hereditary hypophosphatemic rickets.

### 2.2 Hypophosphatasia

Hypophosphatasia is caused by low activity of the tissue non-specific isoenzyme of alkaline phosphatase (TNSALP) caused by a mutation in the ALPL gene. The enzyme normally degrades extracellular inorganic pyrophosphate (PPi) into Pi (inorganic phosphate). Defects in this metabolic pathway leads to accumulation of PPi in the extracellular bone matrix and low alkaline phosphatase levels. Other substrates of TNSALP are pyridoxal-5-phosphate (PLP), and phosphoethanolamine (PEA). PLP is the biologically active form of vitamin B6 and high levels, as seen in hypophosphatasia, are believed to be involved in neurotoxicity. The pathophysiology of the disorder is complex: increased PPi levels inhibit normal mineralization resulting in rickets and osteomalacia, but other tissues and organs can be affected as well. For example, muscle hypotonia is a well-known feature of the disease [4]. The disorder is extremely variable, ranging from severe infantile phenotypes to almost asymptomatic adult cases, with only dental problems. Most common skeletal symptoms are those of rickets, with bone pain, fractures and bowing of legs. The muscle hypotonia may add to walking problems. In the most severe perinatal cases, hypotonia and respiratory distress can lead to early death. Children can develop craniosynostosis and frequently have retarded growth. In adults, normal height can be achieved, but patients may suffer from skeletal complications such as fragility fractures and chronic pain [5].

Treatment of hypophosphatasia consists of supportive care, including regular periodontal and dental care to avoid inflammation, sufficient physical activity and orthopedic interventions. In adult patients, several other modalities have been tried, including the use of teriparatide (parathyroid hormone amino acid 1–34) which has been shown to improve fracture healing and resolve stress fractures [6].

Recently, subcutaneous asfotase alfa (Strensiq®), a first-in-class bone-targeted human recombinant TNSALP replacement therapy, is approved in the EU for long-term therapy in patients with pediatric-onset hypophosphatasia. It was shown that asfotase alfa in this patient group can improve rickets as evidenced by an improvement in radiographically-assessed severity scores at 24 weeks [7]. Furthermore, patients experienced improvements in respiratory function, gross motor function, fine motor function, growth and quality of life [8]. In life-threatening perinatal and infantile hypophosphatasia, asfotase alfa also improved overall survival [9]. However, not all infantile cases have a favorable outcome [10]. Knowledge on long term effectiveness is still scarce and prognostic factors to determine eligibility for treatment insufficiently known. In view of the high costs of this therapy,
collaborative efforts are needed to support the decision making whom to treat. This will also become a challenge for the adult patient group, for whom this modality is still under study.

2.3 Hereditary hypophosphatemic rickets

The most common of the hereditary hypophosphatemic rickets is the X-linked form caused by a mutation in the \textit{PHEX} gene, encoding a phosphate-regulating endopeptidase homolog. Disruption of this enzyme results in a rise in FGF23 levels, suppressing transcription of sodium–phosphate co-transporters in the kidney eventually leading to renal phosphate wasting and hypophosphatemia [11]. An autosomal dominant form resulting from gain of function mutations in FGF23 gives rise to a similar phenotype. Elevated FGF23 levels decrease synthesis and increase catabolism of active vitamin D, resulting in low levels. Patients typically have short stature and lower extremity deformities secondary to rickets at an early age, but milder forms exist. Female carriers of the X-

Table 2 Inborn errors of metabolism with secondary bone disease

| Skeletal symptom/sign | Disorder                                      | Additional symptoms                                                                 |
|-----------------------|-----------------------------------------------|------------------------------------------------------------------------------------|
| Osteoporosis          | All inborn errors of metabolism that require  | Miscellaneous; frequently neurological symptoms                                     |
|                       | strict dietary treatment                      |                                                                                     |
|                       | Galactosemia                                  | Cognitive impairment, primary ovarian failure, cataract*                              |
|                       | Phenylketonuria                               | Cognitive impairment in untreated patients#                                           |
|                       | Homocystinuria                                | Marfanoid habitus, kyphosis, lens luxation, cognitive impairment                     |
| Pathological fractures| Lysinuric Protein Intolerance                 | Protein avoidance, gastrointestinal symptoms hyperammonaemia, lung disease           |
|                       | Wilson disease                                | Liver disease and/or neurological and psychiatric manifestations                     |
|                       | GSD type II (Pompe disease)                   | Muscle weakness, secondary respiratory impairment                                    |

*a*Cataract in untreated patients only  
#In patients born before the introduction of newborn screening or in countries were NBS for this disorder is not performed

**Fig. 1** Examples of typical skeletal abnormalities in patients with inborn errors of metabolism. 

- **a.** Erlenmeyer flask deformity in the distal femur of a patient with type 1 Gaucher disease (plain radiograph).
- **b.** Fragility fractures (arrows) in the femoral neck and diaphysis of a patient with X-linked hypophosphatemic rachitis (T2-MRI).
- **c.** Myelopathy (arrow) due to narrowing of the spinal canal in MPS-I. Note the platyspondylyia (flattened vertebral bodies) as part of the dysostosis multiplex (T2-MRI).
- **d.** Avascular necrosis on the right side, normal femoral head on the left in a patient with Gaucher disease (plain radiograph).
linked variant in particular may present late, sometimes with nonspecific bone pain, fatigue, and weakness. At a later age they may have more localized complaints relating to enthesopathy and early arthropathy. Hypophosphatemia is an important diagnostic finding.

An early diagnosis is important to improve growth and prevent complications [12]. Children should be treated with the active form of vitamin D (calcitriol or alfalcacidol) and phosphate [12, 13]. Not all patients tolerate liquid phosphate very well and thus compliance may be a challenge. For older patients, phosphate tablets may be an alternative. Additional treatment with growth hormone has been tried in children, with inconsistent results and potential side effects. A recent trial suggests that some improvement in pre-pubertal short children can be achieved [14]. Treatment of adults is debated: once adult height has been achieved, the indications for therapy are to reduce osteomalacia and related pain symptoms. It may be difficult in clinical practice to distinguish pain symptoms from related irreversible skeletal complications such as arthropathy. When there is biochemical evidence of osteomalacia or insufficiency fractures, treatment may be of use. In all cases, treatment should be monitored carefully to prevent complications and avoid nephrocalcinosis. For further review of treatment options, please refer to the excellent review article: A clinician’s guide to X-linked hypophosphatemia [13]. A new treatment for both XLH and tumor-induced osteomalacia with high FGF23 levels is the administration of an Anti-FGF23 Antibody (Burosumab, Crysvita®, Ultragenyx). Recent results from a phase 2 and phase 3 studies have shown that burosumab can reduce the loss of phosphate from the kidney, improve abnormally low serum phosphate concentrations and reduces the severity of rickets as shown in x-rays [15, 16]. Treatment has recently received marketing approval in the EU for children with XLH and is being reviewed in the US for both pediatric an adult indications.

2.4 Lysosomal storage disorders

Primary involvement of the bone marrow in inborn errors of metabolism is mainly found in the sphingolipidoses, particularly in Niemann Pick diseases and Gaucher disease. In Gaucher disease, severe bone marrow infiltration with lipid-laden macrophages can occur. Gaucher disease is caused by deficient activity of the lysosomal enzyme β-glucocerebrosidase (GBA; EC 3.2.1.45). GBA hydrolyzes the natural glycosphingolipid glucocerebroside (or glucosylceramide; GC) into glucose and ceramide. Storage of GC in macrophages gives rise to hepatosplenomegaly and involvement of the bone marrow. Severe skeletal pathology can occur in this disorder when left untreated [17]. The pathophysiology involves mass effect of storage with cortical thinning, necrosis, fibrosis and probably low-grade inflammation. Osteopenia or osteoporosis is frequent. Timely intervention with intravenous enzyme replacement therapy or oral substrate inhibitors can prevent skeletal complications. Frequently, bone marrow examination is the clue to a diagnosis of Gaucher disease, since most patients present with splenomegaly and cytopenia, mimicking lymphoma or leukemia. Also in Niemann Pick disease, caused by acid sphingomyelinase deficiency (types A and B), skeletal features can be present as a consequence of storage cells in the marrow. This disorder mainly presents with hepatosplenomegaly and interstitial lung disease, and, in severe cases with neurological symptoms [18]. Bone marrow involvement is less prominent than observed in Gaucher disease. Other skeletal manifestations of the disease include joint and bone pain, decreased bone mineral density and osteoporosis with a risk of fragility fractures [19]. In young adults, short stature and delayed onset of puberty may occur, but most patients eventually attain a normal adult height [20]. Of note, so called “pseudo-Gaucher cells” can be found in a number of disorders and should not be mistaken for the disorders mentioned above: proper diagnostic tests should be employed including genetic and enzymatic tests to confirm any disorder presenting with marrow storage cells.

2.5 The mucopolysaccharidoses and mucolipidoses

Skeletal abnormalities are the hallmark of lysosomal storage disorders caused by accumulation of glycosaminoglycans. There are eleven types of mucopolysaccharidosis (MPS; type I, II, III (with subtypes A,B,C and D), IV (subtypes A en B), VI, VII and IX), each caused by deficiency of one of 11 different GAG degrading enzymes [21, 22]. These are multisystemic disorders, with a wide range in presence and severity of the different clinical features. Skeletal involvement however, is universal in these disorders (with the exception of MPS III) and is of very early onset. Radiological studies in young children with MPS I, II and VI show abnormal bone development [23] and a histological study of a MPS IV foetus show prenatal accumulation of GAGs in chondrocytes [24].

Virtually all bones can be affected in MPS, but alterations in the shape of the skull, the bones of hips, hands, feet and spinal column are most prominent [23]. The term most commonly used for the combination of developmental skeletal alterations in MPS is dysostosis multiplex. Length growth in MPS is reduced and the spinal abnormalities and hip dysplasia results in altered posture. Range of motion of the spine, hips, knees, shoulders, ankle and often also the elbows is reduced to variable degree in all MPS types [25, 26] apart from type IVA, where there is joint laxity, often associated with (sub) luxation of joints [27]. During life, the use of these abnormally shaped joint bones leads to progressive joint damage and secondary arthritic changes, limiting mobility [28, 29]. Hip surgery and later on hip replacement are often performed in order to limit pain and to try to maintain walking ability [28, 29]. Skeletal Pain is a frequent complaint in MPS patients [30]. Other complications of skeletal disease in the MPSes are restrictive
pulmonary disease due to altered thorax shape and thoracic wall stiffness and meninx compression resulting from spinal abnormalities.

How GAG accumulation leads to abnormal skeletal development and the occurrence of (secondary) osteoarthritis is only partially understood. In the growing skeleton, the accumulation of GAGs affects growth plate functioning, as can be seen from the disturbed growth plate architecture in MPS mouse models and a limited number of human bone specimens [31]. Joint inflammation is apparent in MPS hips and knees and is most likely due to secondary to damage from movement of abnormally developed and malformed joints and a direct effect of increasing GAG accumulation, potentially mediated through activation of the Toll Like Receptor 4 [32, 33].

Mucolipidosis type II and III α/β or γ are disorders caused by deficiency of the enzyme UDP-N-acetyl glucosamine-1-phosphotransferase (GLcNAc-P-Tase), which is involved in phosphorylation of lysosomal enzymes, ensuring correct targeting to the lysosome. Aberrant targeting of these degradation enzymes results in lysosomal accumulation of their substrates, including glycosaminoglycans and (glyco)sphingolipids. Mucolipidosis type II is the most severe form, in which rapidly progressive airway, cardiac, skeletal and nervous system disease results in death in early childhood. Mucolipidosis type III has a broader phenotypic range, with milder affected patients surviving into adulthood and displaying primarily debilitating skeletal symptoms. Skeletal symptoms in mucolipidosis bear great similarities to those of mucopolysaccharidosis, with dysostosis multiplex, secondary osteoarthritis of the joints and complications such as meninx compression and severe arthrosis of (amongst others) the hip joints, requiring surgical interventions at an early age [28].

Treatment in the form of enzyme replacement therapy (ERT) is available for MPS type I, II, IV, VI. There may be some positive effect of ERT on joint mobility (specifically range of motion of the shoulder and hips) but by and large this effect is limited and even early treated patients go on to develop severe skeletal complications [34, 35]. Hematological stem cell transplantation, performed in MPS I to halt neurodegeneration, does not have a significant effect on the occurrence of skeletal symptoms either [36]. A small, short duration trial with an anti-inflammatory drug, pentosan polysulphate in MPS I suggests a positive effect on pain [37]. There are no disease specific therapies for mucolipidosis type II and III. In all these disorders supportive care is of great importance, focusing on pain management, physiotherapy and well timed surgical interventions executed by a team experienced in treating these complex multisystem disorders.

### 2.6 Mannosidoses

Alpha mannosidosis is a lysosomal storage disorder in which deficiency of the enzyme α-mannosidase leads to accumulation of mannose-rich oligosaccharides. Similar to the mucopolysaccharidoses and mucolipidoses, this is a clinically widely variable multisystemic disorder with neurocognitive and psychiatric symptoms, corneal clouding or cataract, hearing loss, immune deficiencies, myopathy and skeletal abnormalities. Dysostosis multiplex with skull abnormalities, kyphoscoliosis, pectus carinatum, hip dysplasia and hand and feet deformities are present in variable degree in the majority of patients [38, 39]. Secondary osteoarthritis of the large joints can occur [40] and an increasing incidence of arthrosis is found as patients get older [41]. Alpha mannosidosis has been treated with hematopoeitic stem cell transplantation [42, 43], but there are not enough data to evaluate the effect on the musculoskeletal symptoms of the disorder. Enzyme replacement therapy has been recently granted marketing authorization under exceptional circumstances in the EU. Long term the data on the effectiveness of this treatment, including the effect on musculoskeletal symptoms, will be evaluated.

Beta mannosidosis is an extremely rare disorder caused by accumulation of complex disaccharides due to deficiency of the lysosomal beta-mannosidase enzyme, with few published cases. It has a wide clinical spectrum of which skeletal abnormalities can be a feature [44].

### 2.7 Pycnodysostosis

Pycnodysostosis is an autosomal recessive metabolic bone disorder caused by mutations in the cathepsin K (CTSK) gene [45]. Cathepsin K is a lysosomal protease which is secreted into the sealed extracellular compartment, where it is involved in degradation of bone matrix proteins (e.g. collagen type I, osteonectin and osteopontin). Deficient cathepsin K activity results fragile bones with increased mineral content. Clinically, the disorder is characterized by short stature, pathological fractures and bone dysplasia [46, 47]. Especially craniofacial and dental abnormalities and stubby hands and feet with acroosteolysis are typical of pycnodysostosis, though the latter is not found in all patients [47–49]. No specific treatment for pycnodysostosis is available. Multidisciplinary supportive care, including attention for the risk of upper airway obstruction, is indicated [50].

### 2.8 Alkaptonuria

Alkaptonuria is an autosomal recessive metabolic disorder in the catabolic pathway of the amino acids phenylalanine and tyrosine. Decreased activity of the enzyme homogentisate 1,2-dioxygenase leads to accumulation of homogentisic acid (HGA) and its oxydated derivative benzoquinone acetic acid (BQA) in connective tissue, cartilage and body fluids (e.g. urine, sweat). The deposition of BQA polymer in connective tissue triggers the progressive tissue damage observed in this
disorder. The disorder is sometimes recognized early in life because of discoloration of urine when exposed to oxygen or alkali, but symptomatic disease generally does not develop until the third decade of life. Back and joint pain are the first symptoms to occur and over time, progressive disease results in progressive deformation of the spine and arthropathy of the large joints, necessitating joint replacement at an early age [51, 52]. Other complications of the disease are the formation of stones in the genitourinary tract and cardiac valve disease.

Limiting HGA production can be attempted by a protein restricted diet, though this is hard to comply with, especially in a period of life when individuals are still asymptomatic [53]. Nitisinone (2-(2-nitro-4-trifluoromethylbenzoyl)-1,3-cyclohexanedione, NTBC), inhibiting the enzyme 4-hydroxyphenylpyruvate dioxygenase, can be used to reduce HGA synthesis. In a first clinical treatment trial, urinary HGA excretion was reduced by >95% in response to the highest administered daily dose (8 mg) [54]. Safety of this treatment (NTBC increases tyrosine, with unknown long term consequences) and its effectiveness in preventing clinical complications remains to be proven and a phase 3 trial (Suitability of Nitisinone in Alkaptonuria 2 (SONIA 2)) is currently ongoing.

3 Secondary involvement of the skeleton

Several inherited metabolic disorders have been reported to be associated with reduced bone density (summarized in Table 2). Factors contributing to the reduced mineralization in these disorders can be the disease itself (e.g. lysinuric protein intolerance), dietary treatment (all inborn errors of protein metabolism), hypogonadism (e.g. galactosemia) and/or low vitamin D status due to reduced sunlight exposure (e.g. erythropoietic protoporphyria).

3.1 Galactosemia

In classic galactosemia, deficiency of the galactose-1-phosphate uridyltransferase enzyme results in accumulation of galactose and its derivatives, from both exogenous ingestion and endogenous production. Clinically, this results in life threatening liver disease, often complicated by E Coli sepsis, very early in life. Treatment with a lactose free, galactose restricted diet reverses (or in case of newborn screening prevents) the severe early illness, but long term complications such as ovarian failure and neurocognitive deficits still occur [55]. Bone density has been reported to be below the average values for the general population in several studies in children and adults with classic galactosemia, included in the 2017 meta-analysis by van Erven et al. [56]. This meta-analysis showed that bone mineral density was within two standard deviation of the mean in the majority of patients. An estimated 10–25% had a clinically significant reduced bone mass (Z score ≤−2 SD). Whether this results in an increased fracture risk is unknown [56].

Factors contributing to low bone density in classic galactosemia are reduced dietary calcium and vitamin D intake, ovarian insufficiency, reduced physical activity in some patients and potentially the metabolic disorder itself [57, 58]. Treatment with calcium and vitamin D supplementation, and estrogen in patients with early ovarian insufficiency, improves bone mass in classic galactosemia patients [59, 60]. The additional effect of aging on bone mineral density and fractures remains to be established as early treated patients have not yet reached late adulthood.

3.2 Homocystinuria

Classic homocystinuria is caused by a deficiency of the enzyme cystathionine beta-synthetase (CBS), leading to grossly elevated levels of homocysteine and methionine. Symptoms include lens dislocation, thromboembolisms and cognitive impairment. In bone, high levels of homocysteine impairs collagen type I crosslinking, resulting in reduced bone strength [61]. Patients have no overt skeletal abnormalities at birth, but untreated they develop skeletal abnormalities such as genu valgum, pes cavus and excess growth of long bones as well as osteopenia or osteoporosis [62, 63]. Vertebral fractures and spine deformities can occur. Treatment may consist of a methionine restricted diet and administration of pyridoxine and betaine, depending on whether or not a patient is pyridoxine responsive [64]. Treatment from an early age, with resultant lower homocysteine levels, prevents the occurrence of bone complications [65, 66].

3.3 Glycogen storage disorders

This group of inborn errors are characterized by reduced glycogen degradation and or gluconeogenesis. There over 12 different types of glycogen storage disorders (GSDs) with enzyme deficiencies in these pathways. Dependent on where the enzymes are expressed, deficiency leads to a liver defect (hepatomegaly, hypoglycemia, lactate acidosis), muscle defect (exercise intolerance, myopathy) or a combination of both. In GSD I type a and b additional symptoms such as renal tubulopathy (1a and b) neutropenia and neutrophil and autoimmune phenomena (1b) occur. Low bone density has been described in GSD I type a and b, GSD II, GSD III, GSD V and GSD IX [67–70] 1995 [71–73].

In GSD type 1, factors contributing to the reduction in bone mineral density are poor metabolic control [71, 74, 75] and possibly the treatment with gCSF in GSD1b [74], though the latter relationship was not found in all studies [76]. In a recent study in 4 male GSD I (a and b) patients, all had either osteoporosis or osteopenia, suffered from hypogonadotropic hypogonadism and one patient had a spinal compression fracture at a very young age [75].
In GSD II (Pompe disease) a high incidence of osteoporosis, as well as (asymptomatic) vertebral fractures has been observed [68, 69]. Contributing factors to altered bone composition in Pompe disease are reduced muscle strength [68], lack of physical activity, but there may be a direct effect of the storage disease on bone metabolism as well [69].

3.4 Phenylketonuria

Phenylketonuria (PKU), one of the most prevalent inborn errors of metabolism, is caused by the inability to adequately degrade the amino acid phenylalanine, due to a deficiency of the enzyme phenylalanine hydroxylase. Phenylalanine is neurotoxic and its accumulation causes progressive neurological damage. Since the 1960s newborn screening for PKU is implemented and treatment with a protein restricted diet, combined with phenylalanine free amino acid supplementation, results in preservation of neurocognitive development [77]. A meta-analysis of all publications up to 2015 on bone density in PKU indicated that bone density is on average lower in PKU patients compared to the control population, but not in the range of osteoporosis or osteopenia [78]. Contributing factors to the reduced bone mineral density may be the low protein diet (despite amino acid supplementation) and/or the elevated phenylalanine levels. Low calcium, vitamin D or other vitamins and micronutrient intake is unlikely to influence bone density in the current PKU population, since these are sufficiently present in the presently used amino acid supplements. There is currently no conclusive evidence indicating a higher fracture risk in adults with early treated PKU [79, 80]. Since worldwide newborn screening for PKU has been introduced from the late 1960ties onwards, the effect of aging on fracture rate in early PKU patients cannot yet be established.

3.5 Urea cycle defects, organic acidurias and lysinuric protein intolerance

Urea cycle disorders (UCDs) are a group of inborn errors in which conversion of nitrogen, from either dietary protein or body protein catabolism, to urea is disturbed, causing an increase in neurotoxic ammonia. Clinically, these disorders are characterized by episodes with progressive symptomology of lethargy, anorexia, altered behavior, drowsiness, epilepsy and ultimately coma. Organic acidurias are caused by specific enzyme deficiencies in the intermediary pathways of carbohydrate, protein and fatty acid metabolism, leading to the production of large amount of organic acids. The phenotypical spectrum of these disorder is broader compared to urea cycle disorder and apart may include for example renal, hematological and cardiac complications in addition to the neurocognitive manifestations. As in urea cycle disorders, acute metabolic decompensation, characterized by a high anion gap metabolic acidosis and/or hyperammonemia, may occur as a result of a catabolic state or high protein intake. Treatment of UCDs and organic acidurias, dependent on the type and severity of the disorder, exists of a protein restricted diet, avoidance of catabolic state and ammonia scavengers.

For the urea cycle disorders and organic acidurias the composition of the therapeutic low protein diet and/or the disorders itself may cause low bone mineral density, but limited data on bone density and factors influencing bone density are available for these conditions. In a case series of 81 methylmalonic and propionic aciduria patients, the bone density Z score was low in the majority of patients [81]. In another eight adult organic aciduria patients studied, osteoporosis was present in 2 and osteopenia in 4 patients, with one propionic aciduria patient suffering spinal fractures [82]. In a very small study in male urea cycle defect patients, bone mineral density was lower compared to aged matched control subjects [83].

Lysinuric protein intolerance (LPI), a condition in which reduced activity of the cationic amino acid transporter SLC7A7 leads to reduced intestinal uptake and increased urinary loss of amino acids, is associated with frank osteoporosis and an increased fracture rate [84]. Fractures may even be the presenting symptom in LPI [85, 86]. Classic symptoms of this disorder are intolerance for protein rich food, symptomatic hyperammonemia, failure to thrive. Long-term complications include pulmonary alveolar and progressive renal disease. Reduced amino acid availability, resulting from the low protein content of diet and the deficient cationic amino acid transporter activity, may lead to reduced bone density through a decreased collagen synthesis [87]. A study in healthy discordant monozygotic twins suggest that specifically lower alanine and glycine intake are associated with lower bone mineral density [88]. Treatment of LPI consist of protein restriction, citrulline and/or lysine supplementation and nitrogen scavenger drugs [89]. Whether the pharmacological treatment, either through a direct effect of citrulline and/or lysine availability or through improvement in protein tolerance, has a positive effect on bone density and reduces fracture rate is currently unclear. Individual patients have also been treated with bisphosphonate therapy [90].

3.6 Erythropoietic protoporphyria

In this inborn error of the haem biosynthesis, resulting from either ferochelatase deficiency or increased ALAS2 activity, protoporphyrin IX accumulates in skin and liver. Protoporphyrin IX reacts with light, damaging skin endothelium by reactive oxygen species formation. Clinically this manifests as first severe pain, and second, swelling and erythema of the skin if light exposure continues. As a result, patients avoid exposure to high intensity light, which in practice means they lead their life mainly indoors. This in turn leads to vitamin D deficiency and reduced physical activity,
causing osteopenia and osteoporosis in a significant proportion of patients [91, 92].

3.7 Wilson disease

In this autosomal recessive disorder, copper accumulation occurs due to mutations in the ATP7B gene, encoding for a copper transporting ATPase. The disease is characterized by hepatic, neurological and psychiatric manifestations. Osteoporosis as well as osteopenia occurs more frequently in adult patients compared to control subjects [93] and may be associated with a higher fracture rate [94]. Early onset osteoarthritis of the large joints (predominantly the knees) as well as lower back pain with spinal radiological abnormalities have been described in Wilson patients [95, 96].

4 Diagnosis: When to suspect an inborn error of metabolism

In this section, diagnostic clues to uncover an underlying inborn error of metabolism are discussed for patients presenting with skeletal symptoms. Hence this refers to the first group discussed in this article: those with primary involvement of the skeleton. In clinical practice, patients will be referred to a specialist when they have skeletal deformities, retarded growth or both. This is particularly the case for children. In adults, more frequently subtle deformities, pain or fractures can be initial symptoms of an underlying metabolic defect. Hence, in some attenuated phenotypes, the diagnostic delay can be very long. In the current era of rapidly expanding genetic screening, whole exome sequencing may occur nowadays at an earlier stage. However, these techniques are not readily available in every center. In addition, variants of unknown significance may be discovered, which implicates that a combined approach, using important clinical, biochemical, radiological and other tools are needed for a correct diagnosis [97]. To improve awareness for metabolic disorders amongst the high number of acquired and inherited skeletal dysplasias and other factors that may affect bone, a brief summary of clues to a diagnosis of inborn errors of metabolism will be discussed (summarized in Tables 1 and 2).

4.1 Additional clinical signs or symptoms

Within the group of lysosomal storage disorders, Gaucher disease is an example of a disorder that may present with pain as a first symptom. Although in most cases splenomegaly or cytopenia are the first symptom, in rare instances patients may come to medical attention because of a skeletal problem. Severe bone crises may occur, as well as avascular necrosis of the femoral head [17]. Additional signs are typical radiographic images such as Erlenmeyer flask deformities of the femurs. Both in Gaucher disease and pycnodysostosis, pathological fractures are reported. In acid sphingomyelinase deficiency, skeletal disease is much less prominent, but arthropathy may occur. The mucopolysaccharidoses have as most prominent features the dysplasia of the skeleton, but may also also present as a first sign in attenuated phenotypes with femoral head abnormalities resembling avascular necrosis [98]). True avascular necrosis of the femoral head is more frequent in MPS III, Sanfilippo disease [99]. Early arthropathy is common, and is also the main symptom of skeletal disease in the mucolipidoses. Joint stiffness and carpal tunnel syndrome are frequent symptoms of ML III, which resembles features of MPS I and VI. In mild MPS cases, such as MPSI-Scheie, arthropathy of the hands may mimic rheumatological disease [100]. Arthropathy is also common in alkaptonuria and hypophosphatasia. In the latter, chronic pain may also be the result of myopathy [4]. Abnormalities of the spine can be found in Gaucher disease (pathological fractures, crises, kyphosis) and in MPS disorders (dysplasia, kyphosis, scoliosis). Osteoporosis is rarely an isolated finding and unlikely to be a presenting symptom.

Thus, in patients with joint stiffness, early arthropathy, rheumatoid factor negative arthritis, femoral head necrosis, unexplained bone pain, radiographical evidence of marrow expansion or skeletal dysplasia, especially in the presence of other features such as abnormal growth, underlying metabolic disorders should be part of the differential diagnosis.

4.2 Biochemistry

The presence of very low levels of bone-specific alkaline phosphatase and calcific deposits are a clue to the diagnosis of hypophosphatasia [11]. In hereditary hypophosphatemic rickets hypophosphatemia and low-normal circulating 1,25-dihydroxyvitamin D [1,25(OH)2D] levels are typical biochemical findings. Serum alkaline phosphatase (ALP) activity can be elevated in children and also sometimes in adult patients. Especially in patients with mild phenotypes, a low 25-vitamin D level with elevated ALP will be easily mistaken for vitamin D deficiency. The low phosphate level is the clue to the diagnosis in this case. In all other metabolic disorders presenting with skeletal features, routine biochemistry is usually normal.

4.3 Genetics

In addition to the diagnostic clues described above, appropriate genetic testing should always follow for confirmation of the diagnosis, although in some cases, enzymatic testing is the gold standard for diagnosis (e.g. in the lysosomal storage disorders). Genetic counseling is essential to find other affected family members, assist in decision making for family planning and in some cases to be able to predict disease outcome.
5 Therapy and management

Skeletal symptoms, in the form of dysplasia, arthropathy and/or reduced mineral density are part of the symptomatology of a large number of inborn errors of metabolism. Due to better targeted therapies, as well as supportive care, more patients survive into adulthood and in many cases these skeletal symptoms become a more prominent feature of these disorders. This makes awareness of these skeletal symptoms as well as the options for management of the accompanying complaints all the more important.

Factors positively influencing healthy bone formation in the general population are adequate intake of calcium and vitamin D, preventing underweight, avoiding tobacco and excessive alcohol use and performance of weight bearing exercise. Their importance needs to be emphasized in the care of patients suffering from IEMs, in whom some of these aspects may provide a challenge. In case of primary ovarian failure (galactosemia) or hypogonadism (e.g. in glycogen storage disorder type I) appropriate hormone replacement therapy should be started to prevent osteopenia or osteoporosis. The role of bisphosphonate therapy in the treatment of patients with IEMs is not well established and currently the best advice is to treat osteopenia and/or osteoporosis according to general guidelines. In hereditary hypophosphatemic rickets treatment with active vitamin D and phosphate in childhood is important.

For several disorders, disease specific therapy with an effect on bone metabolism is or will become available, e.g. Asfotase alfa for hypophosphatasia, anti-IGF23 for hypophosphatemic rickets and Nitisinone for alkaptonuria. The long term effect of these drugs on the skeletal manifestation of these disorders remains to be established. Pain management, physiotherapy and well timed surgical interventions, executed by a team experienced in treating these complex multisystem disorders, remain the cornerstone of disease management and their importance in maintaining mobility and improving quality of life should not be underestimated.

Compliance with ethical standards

Conflict of interest  Mirjam Langeveld and Carla Hollak have no conflict of interests concerning any aspect of the submitted work. Outside of the submitted work, they are involved in pre-marketing studies with Genzyme, Protalix and Idorsia. Financial arrangements are made through AMC Research BV. No fees, travel support or grants are obtained from the Pharmaceutical Industry.

Ethical approval This article does not contain any studies with human participants performed by any of the authors.

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