

## Review

# Adult Presentation of X-Linked Hypophosphatemia

Nobuaki Ito<sup>1,2</sup><sup>1</sup> Division of Nephrology and Endocrinology, The University of Tokyo Hospital, Tokyo, Japan<sup>2</sup> Osteoporosis Center, The University of Tokyo Hospital, Tokyo, 113-8655, Japan  
nobitotky@gmail.com

**Abstract:** Adult X-linked hypophosphatemia (XLH) patients present with specific symptoms, including enthesopathies (e.g., ossification of the longitudinal ligament (OPLL), osteophytes around the large joint, and enthesopathy in the Achilles tendon), the development of severe secondary and tertiary hyperparathyroidism (SHPT/THPT) and the subsequent progression of chronic kidney disease (CKD). In addition, these patients exhibit the typical phenotypes of osteomalacia, such as pseudofracture and fracture in weight-bearing bones, odontitis, and tooth abscess. The mechanism underlying enthesopathy development is unknown; however, a common underlying mechanism among XLH and autosomal recessive hypophosphatemic rickets, ARHR1.2, due to mutations in *PHEX*, *DMP1*, and *ENPP1*, is assumed. Clarification of the pathogenesis and drug discovery for this complication is an urgent issue to address, as many adult XLH patients suffer subsequent debilitating nervous symptoms or impingement syndrome, and existing treatments are ineffective. Severe SHPT and THPT are associated with conventional therapy, including active vitamin D and phosphate supplementation, and complicated and careful adjustment of the dosage by experienced clinicians is required to avoid SHPT/THPT. Burosumab is a very effective therapy without risk for the development of SHPT/THPT. However, the indication of this drug should be carefully considered along with the cost-effectiveness, guidelines or recommendations and health care system of each country.

**Keywords:** X-linked hypophosphatemia; fibroblast growth factor 23; osteomalacia; enthesopathy; secondary hyperparathyroidism; tertiary hyperparathyroidism; chronic kidney disease; oral disease; quality of life; burosumab

## 1. Introduction

X-linked hypophosphatemia (XLH) is a genetic disease caused by inactivating mutations of the phosphate-regulating endopeptidase gene (*PHEX*). Symptoms that develop during childhood are typical rachitic phenotypes, including leg deformity, short stature, odontitis, tooth abscess, and craniosynostosis. In contrast, adult XLH patients present atypical symptoms for osteomalacia, such as enthesopathy, secondary/tertiary hyperparathyroidism (SHPT/THPT), and chronic kidney disease (CKD), in addition to characteristic phenotypes for osteomalacia, such as pseudofracture and fracture in weight-bearing bones [1, 2]. Some clinicians believe that the treatment could be terminated after epiphyseal closure even among patients with severe XLH; however, adult XLH patients are currently recognized to be at risk of pseudofractures and fractures in the weight-bearing bones due to low turnover even with normal to high bone mineral density (BMD) [3, 4]. The mechanism for the development of enthesopathy in adults with XLH is unknown, and even among adult XLH patients treated with active vitamin D and phosphate supplementation or burosumab, remarkable enthesopathies sometimes develop and these patients experience significant difficulty in activities of daily life (ADL) and lower quality of life (QOL) with nervous symptoms due to ossification of the posterior longitudinal ligament (OPLL) or impingement syndromes in the hip [4].

Treatment for adult XLH patients with active vitamin D and inorganic phosphate is very difficult, as well as for child patients with XLH, because excess of this treatment is

associated with the development of SHPT/THPT and subsequent nephrotic diabetes insipidus, prerenal renal failure, postrenal renal failure and the progression of CKD [5]. Therefore, the transition of XLH patients to specialized clinicians is vital, especially when patients are treated with conventional therapy, although there are a limited number of specialists for bone and mineral disorders in adults.

In this review article about the adult presentation of XLH, the estimated underlying mechanisms for the development of adult XLH-specific problems, including enthesopathy, SHPT/THPT, and CKD, are illustrated, and the recommended indication and selection of the pharmaceuticals and detailed adjustment procedure of dosage according to previous publications and personal experience are introduced.

## 2. Diagnosis of adult XLH

Mild cases of undiagnosed XLH are infrequently suspected among adults with pseudofractures and fractures in weight-bearing bones or remarkable enthesopathies in the spinal ligament, around the hip joints, and in the Achilles tendons with concomitant chronic hypophosphatemia. To accelerate an accurate diagnosis, appropriate treatment and prevention of additional pseudofracture, fracture, and dental disorders, clinicians should suspect low-turnover disorders such as FGF23-related hypophosphatemia, including XLH and tumor-induced osteomalacia, Fanconi syndrome, vitamin D deficient osteomalacia, mild vitamin D dependent rickets (e.g., heterozygous mutation in *CYP3A4*), mild hypophosphatasia (e.g., heterozygous mutation in *ALPL*), mild osteogenesis imperfecta (e.g., Sillence type I), and mild osteopetrosis (e.g., dominant negative type of heterozygous mutation in *CLCN7*). These ailments should be suspected among the adult patients who develop pseudofractures and fractures in weight-bearing bones (costa, pelvis, femoral head subchondral fragility fracture, diaphyses of femur/tibia/fibula, calcaneus, and metatarsal) spontaneously or with low-power trauma or with relatively short term use of anti-resorptive reagents for osteoporosis (e.g.,  $\leq 5$  years) or among the adult patients who develop odontitis, tooth abscess, necrosis of the jaw spontaneously or with relatively short term use of anti-resorptive reagents for osteoporosis.

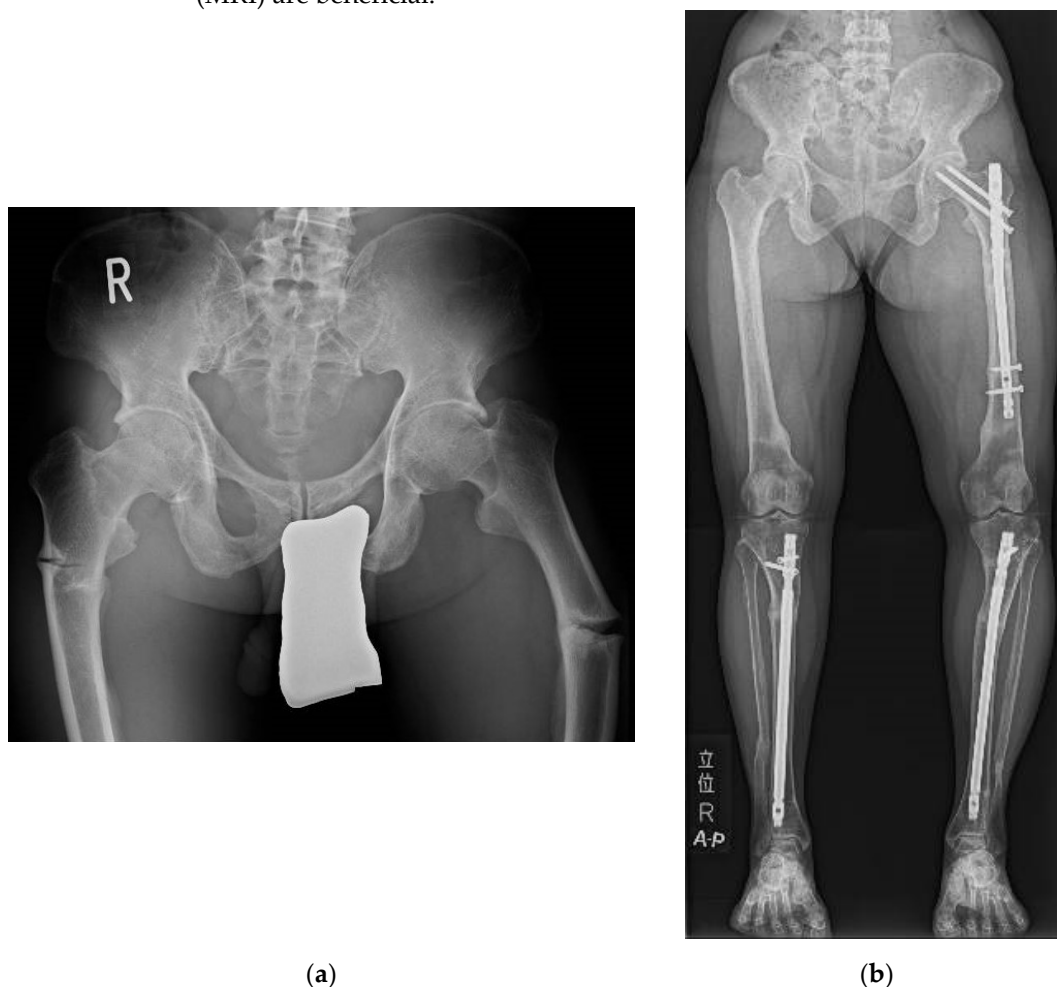
In Japan, once chronic hypophosphatemia with the relevant symptoms of rickets/osteomalacia is recognized, the measurement of serum intact fibroblast growth factor (FGF) 23 (Determinar CL FGF23; Minaris Medical, Tokyo, Japan) is encouraged to determine the etiology of hypophosphatemia with a cutoff value of 30 pg/mL to discriminate FGF23-related hypophosphatemic rickets/osteomalacia and others, which was revealed to possess high sensitivity and specificity [6, 7, 8]. Among adult patients with osteomalacia accompanied by FGF23 values of 30 pg/mL and more, diagnosis of XLH is strongly supported by the presence of one or more of the following symptoms: mild short stature, leg deformity (genu varum, genu valgus), enthesopathy or X-linked inheritance of rickets/osteomalacia. However, even among these cases, genetic diagnosis of *PHEX* mutation is recommended, if available, in an effort to make an accurate diagnosis [9]. In the cases of adult-onset FGF23-related hypophosphatemic osteomalacia without any of the symptoms above, the possibilities of tumor-induced osteomalacia, intravenous infusion of iron preparation-induced osteomalacia, and alcohol-induced FGF23-related hypophosphatemic osteomalacia should be explored [10, 11, 12].

## 3. Symptoms of adult XLH

### 3.1. Pseudofracture and fracture

As stated above, some clinicians used to believe that the treatment could be terminated after epiphyseal closure, probably due to the normal or relatively high bone mineral density detected in the majority of adults with mild XLH [3, 4]. Normal to high BMD observed in patients with mild XLH might stem from suppressed osteoclastic function and probably due to the excessive ossification caused by the same reason XLH patients tend to develop enthesopathy [4]. However, the risk of pseudofractures and fractures in weight-bearing bones (costa, pelvis, femoral head subchondral fragility fracture,

diaphyses of femur/tibia/fibula, calcaneus, and metatarsal) are usually not correlated with BMD and strongly associated with low turnover state of the bone (e.g., also shown in other hypophosphatemic rickets/osteomalacia, hypophosphatasia, osteogenesis imperfecta, osteopetrosis, and long-term use of anti-osteoclastic reagents), as accumulated micro bone crackles with delayed healing in weight-bearing bones eventually leading to pseudofracture and fracture. Therefore, the treatment for hypophosphatemia should be continued among adult XLH patients with typical rachitic phenotypes, including short stature and leg deformity, or a past history of surgical correction of leg deformity, with continuously elevated bone-specific alkaline phosphatase (BAP) in the absence of treatment [1, 2]. Figure 1 (a), (b) show typical femoral pseudofractures developed after years of treatment cessation, and surgically treated fractures in the diaphysis of the femur and bilateral tibiae developed under conventional therapy (Figure 1). To detect tiny pseudofractures, X-ray in multiple directions, bone scintigraphy ( $^{99m}\text{Tc}$ -methylene diphosphonate/hydroxymethylene diphosphonate), and T2-weighted fat-suppressed magnetic resonance imaging (MRI) are beneficial.



**Figure 1.** Pseudofracture and fracture in the femurs and tibiae in adult XLH patients. (a) Pseudofractures developed spontaneously in both femurs in an adult XLH patient after years of conventional therapy cessation. (b) Fractures developed spontaneously in the left femur and both tibiae in an adult XLH patient with conventional therapy.

To prevent the development of pseudofracture and fracture, long periods of strenuous exercise and labor (e.g., long period of walking, stomping exercise for osteoporosis, or weight bearing exercise/labor) should be avoided in adult XLH patients with uncontrolled chronic hypophosphatemia manifested by elevated BAP, and the patients should be encouraged to visit their attending clinician immediately when warning pain is recognized in the diaphysis of the femur/tibia/fibula. Adult XLH patients with new

pseudofracture or fracture should be treated with conventional therapy or burosumab. It is recommended that patients who developed pseudofracture or fracture under the treatment with conventional therapy consider changing the treatment to burosumab. When a surgical procedure is unavoidable for the developed pseudofracture/fracture, medical treatment should be provided for 1 to 3 months before surgery to prevent the loosening of prosthesis or the development of additional pseudofracture/fracture around prosthesis unless urgent surgery is required [2].

Please note that anti-resorptive reagents (bisphosphonates and denosumab) are contraindicated for patients with uncontrolled low bone turnover disorders, including adult XLH, as these reagents aggravate low bone turnover and increase the risk of developing pseudofracture and fracture in weight-bearing bones, tooth abscess and necrosis of the jaw. If coexisting osteoporosis is suspected in postmenopausal females with XLH or older patients with XLH, anti-osteoporotic treatment should not be initiated until osteomalacia improvement by normalization of BAP with conventional therapy or burosumab is confirmed.

### 3.2. Muscle weakness

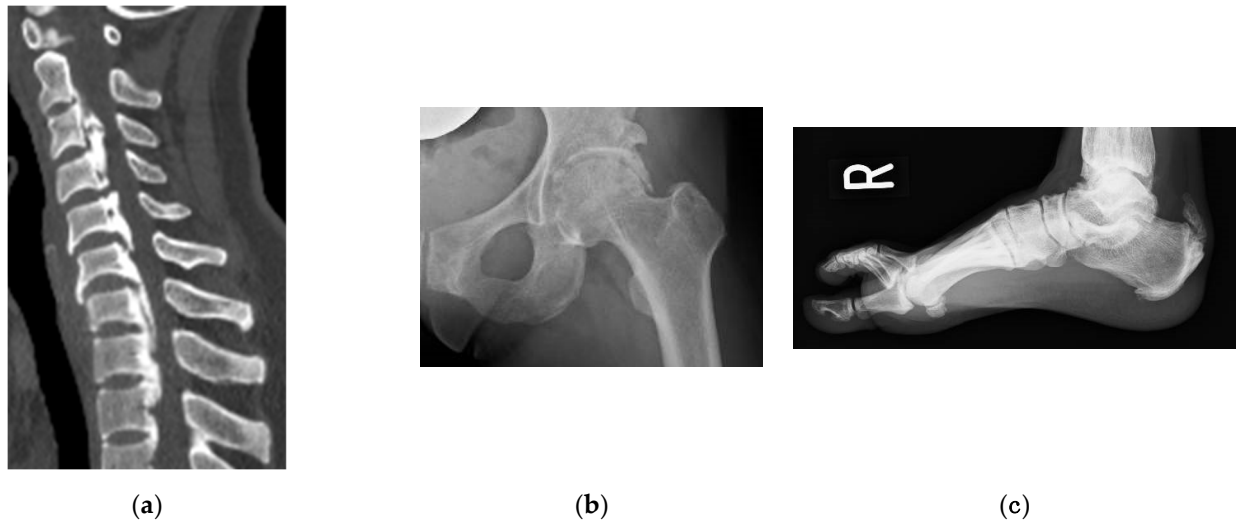
In 2013, Veilleux et al. reported that muscle force in the lower extremities in 13 XLH patients (6 to 60 years old) was significantly decreased despite normal muscle size in comparison with age- and sex-matched control participants [13]. Clinicians tend to ascribe this muscle weakness in XLH patients to the direct effect of hypophosphatemia on myocytes. However, the observed muscle weakness was also explained by bone pain due to osteomalacia, inefficient transduction of muscle contraction due to leg deformity or impingement syndrome stemming from enthesopathy. Of note, rapidly induced hypophosphatemia is not associated with muscle weakness [14]. Further clinical studies are warranted to clarify whether there is a direct effect of hypophosphatemia on myocytes.

### 3.3. Dental health

Uncontrolled rickets/osteomalacia is evidently associated with a high risk of periodontitis, odontitis, and tooth abscess in children and adults [15, 16]. The underlying mechanism for this might be the accumulation of microcracks with delayed healing in the enamel and dentin, which penetrates from the pit on the surface to the pulp, similar to how pseudofractures and fractures develop in weight-bearing bones. Orthodontic treatment sometimes results in the loss of permanent teeth in XLH patients with uncontrolled osteomalacia. Early intervention with conventional treatment was reported to have a prophylactic effect on the development of these dental diseases [17, 18].

### 3.4. Enthesopathy

We examined the prevalence of enthesopathies in 25 adult XLH patients (18 to 72 y) and revealed a high prevalence of OPLL (32%), osteophytes around the hip joints equivalent to a Kellgren-Lawrence grade of 2 and more (96%), and enthesopathies in the Achilles tendon (72%), which explained that XLH is an obvious genetic condition in which patients are prone to developing enthesopathies [4]. Normal to high BMD in adult XLH patients was also reported in the same article and was attributed to the same osteogenic nature of XLH [4]. In some adult XLH patients, neurological symptoms due to OPLL and limited range of motion (ROM) in the hip and intervertebral joints severely lowered ADL and QOL [19]. Figure 2 (a), (b), (c) are the typical images of OPLL, osteophytes around the hip joint, and enthesopathy in the Achilles tendon presented in adult XLH patients (Figure 2).



**Figure 2.** Enthesopathies in adult XLH patients (a) Severe ossification of the posterior longitudinal ligament and ossification of the anterior longitudinal ligament presented in an adult XLH patient. (b) A large osteophyte developed around the left hip joint in an adult XLH patient causing severe impingement syndrome. (c) Enthesopathy developed in the right Achilles tendon in an adult XLH patient.

Currently, the precise mechanism of the development of enthesopathy in adult XLH patients (typically over 30 years old) is unknown. It has been recognized that enthesopathy is also frequently present in patients with other inherited FGF23-related hypophosphatemic rickets, including autosomal recessive hypophosphatemic rickets 1 and 2 caused by homozygous mutations in *DMP1* and *ENPP1*, respectively, although other types of inherited or acquired FGF23-related hypophosphatemia are not associated with the development of enthesopathy [20, 21]. Recently, we reported that haploinsufficiency of ectonucleotide pyrophosphatase/phosphodiesterase (ENPP1) with heterozygous or compound heterozygous mutations of *ENPP1* is also associated with milder phenotypes of enthesopathy manifested by OPLL and diffuse idiopathic skeletal hyperostosis (DISH) [22]. This means that the enthesopathy present in patients with XLH and ARHR1/2 is not a consequence of chronic hypophosphatemia or high levels of serum FGF23, and there is a common mechanism to develop enthesopathy among XLH and ARHR1/2. Therefore, unfortunately, both conventional treatment and burosumab do not exert a prophylactic effect on the development of enthesopathy or an improving effect on enthesopathy. Severe neurologic symptoms, such as paraparesis due to OPLL, are an indication for surgical decompression of the spinal cord.

ENPP1 is a membranous enzyme that metabolizes adenosine triphosphate (ATP) into adenosine monophosphate (AMP) and inorganic pyrophosphate (PPi). PPi was identified to antagonize the formation of hydroxyapatite. Thus, lowering plasma PPi is the candidate mechanism for the development of enthesopathy in patients with homozygous, compound heterozygous or heterozygous ENPP1 mutations [23, 24]. Furthermore, Maulding et al. reported that low plasma PPi levels were identified in Hyp mice, the model mouse for XLH, implying the involvement of lowered PPi in the development of enthesopathy in XLH [25].

### 3.5. SHPT/THPT

XLH patients tend to develop SHPT (hyperparathyroidism with the value of calcium < middle of the reference range) due to low 1,25(OH)<sub>2</sub>D stemming from excessive action of FGF23 and exogenous supplementation of inorganic phosphate. SHPT is an undesired phenotype in XLH patients because hyperparathyroidism stimulates osteoclastic activity leading to bone loss in addition to rickets/osteomalacia and accelerates hypophosphatemia as parathyroid hormone (PTH) decreases the expression of sodium-phosphate



cotransporter independent of FGF23. Additionally, we recently reported that PTH directly stimulated the transcription of FGF23 in an osteocytic cell line [26]. Furthermore, a handful of XLH patients who already suffer from SHPT develop THPT (hyperparathyroidism with the value of calcium  $\geq$  the middle of the reference range), which is an equivalent clinical condition to primary hyperparathyroidism (PHPT) that causes hypercalciuria and nephrotic diabetes insipidus, leading to repetitive episodic prerenal and postrenal renal failure and progression of CKD.

In 2019, DeLacey et al. reported that among 84 patients with XLH (adult 40, child 44), 83.3% had SHPT or THPT, and THPT developed in 16.7% of patients [5]. Seventy-five percent (6/8) of the patients who underwent parathyroidectomy for THPT experienced persistent or recurrent THPT, which explains why other parathyroid cells are also ready to convert into autonomously PTH-producing cells in the XLH patients who once developed THPT [5]. We are now analyzing the relationship between the highest or cumulative dose of phosphate supplementation and the development of SHPT or THPT in the large cohort of XLH patients in Japan and Korea [27]. To prevent the development of severe SHPT and THPT, the dosage of conventional treatment with active vitamin D and phosphate supplementation should be adjusted with great care by experienced clinicians, and the change in treatment to burosumab should be considered for XLH patients who develop severe SHPT or THPT. Once THPT is developed, immediate cessation of conventional therapy and initiation of an allosteric modulator of the calcium sensing receptor should be considered until parathyroidectomy conduction occurs, and the treatment for hypophosphataemia should be resumed with burosumab afterward. Detailed guidance for conventional therapy recommended by the author is described in section “6.2. Conventional therapy (active vitamin D and phosphate supplementation).”

### 3.6. CKD

Some adult XLH patients with conventional treatment experience CKD progression uncommon for their age or for those with other medical conditions, and very rarely, a few patients need renal replacement therapy. The main reason for the progression of CKD in patients with adult XLH is described in the section above “3.5. SHPT/THPT”; that is, the main reason is hypercalciuria and nephrotic diabetes insipidus due to conventional therapy, leading to repetitive episodic prerenal and postrenal renal failure, which often follows after the development of THPT. In fact, in the article by DeLacey et al. introduced in the previous section including 84 XLH patients, nephrocalcinosis and CKD G3 and over ( $\text{eGFR} < 60 \text{ ml/min/1.73 m}^2$ ) were more prevalent in patients with THPT than in patients without THPT (60% vs. 18.6% and 35.7% vs. 1.5%, respectively) [5]. Therefore, to prevent the progression of CKD, adequate handling of severe SHPT and THPT by experienced clinicians is necessary, and appropriate water consumption and intake of salt should be recommended among XLH patients treated with conventional therapy, especially in those with febrile and gastrointestinal disorders or who labor and spend leisure time in the hot sun over a long period. Please note that the dosage of active vitamin D and phosphate supplementation needs to be decreased as CKD progresses; otherwise, a persistent excessive dose of conventional therapy might be associated with the development of THPT and further progression of CKD. The correlation between the highest or cumulative dose of phosphate supplementation and active vitamin D and the progression of CKD to stage G3 and over is also under examination in a large cohort of XLH patients in Japan and Korea [27].

### 3.7. Hypertension and left ventricular hypertrophy

A high prevalence of elevated blood pressure (27.3%, 6/22) unmatched to age has been reported in a small cohort of adult XLH patients, and this characteristic could be the consequence of the progression of CKD. A cohort study with a larger number of adult XLH patients will be necessary to address this issue [28].

#### 4. QOL of adult XLH

There are several reports about the QOL of adult XLH patients [29, 30, 31]. In 2019, Skrinar et al. reported that among 232 adult patients with XLH, 97% had bone or joint pain/stiffness, 44% had a history of fractures, 46% had osteophytes, 27% had enthesopathy in the Achilles tendon, and 19% had spinal stenosis. In addition, the mean score for age-specific patient-reported outcomes (PROs) evaluating pain, stiffness and physical function were worse than those of the control population [29]. In 2020, Seefried et al. conducted a systematic literature review including 91 articles and 44 congress abstracts, revealing that XLH had a substantial and wide-ranging negative impact on health-related quality of life (HRQOL), particularly relating to physical function and pain measured by the 36-item Short-Form Health Survey (SF-36), Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC), and Brief Pain Inventory (BPI) [30]. We evaluated the HRQOL in Japanese and Korean patients with XLH, including 32 adult patients; this investigation revealed that among adult XLH patients, 59.4% had bone pain, 65.6% had joint pain, 9.5% had CKD G3 or higher, 15.6% had nephrocalcinosis, 15.6% had SHPT or THPT, and 6.3% underwent parathyroidectomy [31]. In our study, the SF-36, WOMAC and BPI also revealed lower QOL in adult XLH patients [31]. Given the higher prevalence of joint pain (65.6%) than that of bone pain (59.4%) observed in our study, (1) early initiation of pharmaceutical intervention to prevent leg deformity and (2) exploration of the mechanism for the development of enthesopathy to develop new treatment options to conquer this debilitating complication are important to further improve the QOL among adult XLH patients.

#### 5. Transition of XLH patients

Clinical follow-up of XLH patients should be transferred from pediatricians to adult endocrinologists or rheumatologists when patients are approximately 18 to 20 years of age to encourage independence from their caregivers and facilitate the care of other adult-specific medical problems. At the time of transition, patients need to be educated or re-educated about XLH, including the genetic information. XLH patients should be followed by experienced clinicians, as stated above, in the institutions where multidisciplinary care, which involves physical and occupational therapy, dental care, pain clinics, etc., can be offered. For more information about the transition of XLH, please refer to the well-organized mini-review by Dahir et al. [32]. Most importantly, XLH patients should be referred to experienced clinicians, although there are fewer endocrinologists and rheumatologists worldwide specializing in bone metabolic disorder and skeletal dysplasia to meet the demand. Therefore, the need for an increased number of endocrinologists and rheumatologists who are educated about bone metabolic disorders and skeletal dysplasia is one of the urgent issues to resolve in this field.

#### 6. Treatment of adult XLH

##### 6.1. Indication and selection of treatment

The indication and selection of conventional treatment and burosumab among adult XLH patients might be influenced by the health care system of each nation. In Japan, I suggest the indication and selection of pharmaceutical treatment in adult XLH patients as described in Table 1; this information reflects my personal opinion based on my own experience with adult XLH patients (Table 1). The severity of the symptoms among adult XLH patients varies widely even within family members sharing the same mutation, and treatment for hypophosphatemia is not necessarily required among patients with mild symptoms defined by a normal value of BAP, stature  $\geq -1.0$  SD, lack of genu varum and genu valgus, and no history of pseudofracture or fracture in weight-bearing bones or odontitis/tooth abscess without any treatment. In contrast, adult XLH patients with severe phenotypes should be treated with burosumab, which is more effective than conventional therapy. In my opinion, severe phenotypes are defined by short stature ( $< -2.0$  SD), deviation of mechanical axis of the leg into zones 3 or more [33], history of corrective surgery

on the leg, history of pseudofracture in weight-bearing bones, odontitis, tooth abscess for more than two times without any treatment or more than once under conventional therapy, and history of fracture in weight-bearing bone. The indication for conventional treatment is between the indications for observation and burosumab. Additionally, I recommend changing the treatment approach to include burosumab for adult XLH patients who developed uncontrolled SHPT or THPT with conventional treatment and patients with moderate symptoms (between mild and severe symptoms) and eGFR< 45 ml/min/1.73 m<sup>2</sup> to prevent further progression of CKD.

Please note that enthesopathy could not be improved by therapeutically targeting the increase in serum phosphate. Supplementation of natural vitamin D preparation (ergocalciferol, cholecalciferol) or encouraged intake of natural vitamin D from diet targeting serum 25OHD of 30 ng/mL and more are recommended in XLH patients to avoid detriment in the bone due to vitamin D deficiency.

**Table 1.** Suggested indication and selection of the therapy for adult XLH patients.

Observation	Conventional therapy	Burosumab
BAP (≤ upper limit of the reference range)	BAP (> upper limit of reference range) with eGFR≥45 ml/min/1.73 m <sup>2</sup>	BAP (> upper limit of the reference range) with conventional treatment
Stature (≥ -1.0 SD)	Short stature (-2.0 ≤ -1.0 SD) with eGFR≥45 ml/min/1.73 m <sup>2</sup>	Short stature (< -2.0 SD)
Mechanical axis of the leg within zone 1 [33]	Deviation of the mechanical axis of the leg into zone 2 [33] with eGFR≥45 ml/min/1.73 m <sup>2</sup>	Deviation of the mechanical axis of the leg into zone 3 or greater [33] or history of corrective surgery for the leg deformity
No history of pseudofracture/fracture in the weight-bearing bone or odontitis/tooth abscess	Pseudofracture in the weight-bearing bone or odontitis/tooth abscess once with eGFR≥45 ml/min/1.73 m <sup>2</sup>  No response or trivial response to burosumab	Pseudofracture in the weight-bearing bone or odontitis/tooth abscess more than two times or more than once with conventional treatment  Fracture in the weight-bearing bone  Uncontrolled severe SHPT with conventional treatment (e.g., peak intact PTH ≥ double-folds of the upper limit of the reference range) THPT  Patients with symptoms described in the “conventional therapy” and eGFR<45 ml/min/1.73 m <sup>2</sup>

6.2. Conventional therapy (active vitamin D and phosphate supplementation)

Treatment of XLH patients with conventional therapy is the tricky part of the patient management, as inadequate and careless dosage adjustment results in the development of SHPT/THPT and consequent irreversible progression of CKD. Thus, clinicians should always prioritize the prevention of the development of severe SHPT and THPT as long as patients are treated with conventional therapy.

The range of daily dosages of conventional therapy described in the consensus statement introduced by specialists from European countries (calcitriol: 0.50 to 0.75 µg, alfalcidol: 0.75 to 1.5 µg, phosphate supplementation: 750 to 1,600 mg) appears very appropriate from the viewpoint of the prevention of severe SHPT and THPT [2]. Calcitriol should be taken twice daily due to its shorter half-life compared to alfalcidol, which is taken once daily. Initiation of active vitamin D should be preceded approximately one week before starting phosphate supplementation, and treatment with phosphate



supplementation should always be accompanied by active vitamin D to prevent the development of severe SHPT and THPT [1, 2]. Importantly, the serum phosphate level peaks at approximately 1.5 hours after the intake of phosphate supplementation and goes back to the trough value 2 to 3 hours after intake [34]. Thus, phosphate supplementation should be taken four or more times daily to maximize the antihypophosphatemic effect and minimize the risk of developing SHPT/THPT.

I propose laboratory tests, including serum phosphate, calcium, albumin, intact PTH, creatinine, and BAP, with blood samples drawn 1 to 2 hours after the intake of phosphate supplementation to detect the peak value of serum phosphate and intact PTH because these values are associated with the effect on the mineralization of the bone and risk for the development of severe SHPT/THPT. These effects and risks could not be inferred from the trough value of serum phosphate and intact PTH.

In our facility, conventional therapy usually starts with 1.0  $\mu\text{g}$  of alfacalcidol, and 1 to 2 weeks later, phosphate supplementation is initiated with 200 mg four times daily. In the dose-adjusting phase, laboratory data are followed every one to four weeks, and phosphate supplementation is adjusted by 100 mg of increment or decrement for a time until the peak phosphate level stably settles within the lower 50% of the reference range. Most importantly, we did not attempt to adjust conventional therapy to increase trough phosphate levels within the reference range, as it is strongly associated with the development of SHPT/THPT and consequent progression of CKD [1]. Additionally, in adult XLH patients with  $\text{eGFR} < 45 \text{ ml/min/1.73 m}^2$ , a lower dosage of active vitamin D and phosphate supplementation is required to maintain the peak serum phosphate level in the target range, and often only a small dose of active vitamin D (e.g., 0.25 to 0.50  $\mu\text{g}$  of alfacalcidol) or no medication is required among adult XLH patients with  $\text{eGFR} < 30 \text{ ml/min/1.73 m}^2$ , although the change in treatment to burosumab should be recommended to prevent the further progression of CKD in case phosphate supplementation is still required (e.g., uncontrolled BAP) in patients with  $\text{eGFR} < 45 \text{ ml/min/1.73 m}^2$ . In the maintenance phase, laboratory data are followed every 3 to 6 months, and normalization of BAP is the ideal goal of treatment, as it directly reflects the ossification status of the bone. However, in treatment-naïve XLH patients with active osteomalacia, BAP initially increases in response to treatment up to 3 to 6 months, indicating the recommencement of mineralization, and then decreases to the basal value at approximately 6 to 12 months after the initiation of treatment. Thus, dose adjustment of conventional therapy targeting normalization of BAP should be considered after 12 to 18 months. Changing treatment to burosumab should be considered in patients with uncontrolled BAP.

Once again, the prioritized agenda alongside conventional treatment is the prevention of severe SHPT and THPT; therefore, the dosage of phosphate should be immediately decreased by 100 mg or more for a period of time once overadjustment of peak phosphate (e.g., within the upper 25% of the reference range or more) or severe SHPT (e.g., peak intact  $\text{PTH} \geq$  double the upper limit of the reference range) is observed. Then, if an increase in phosphate supplementation is required in patients who once develop severe SHPT, escalation of active vitamin D by 0.25  $\mu\text{g}$  for calcitriol and 0.25 to 0.5  $\mu\text{g}$  for alfacalcidol should precede 1 to 2 weeks before an increase in phosphate supplementation. In patients who develop THPT (hyperparathyroidism with the value of calcium  $>$  the middle of the reference range), conventional therapy should immediately be terminated to prevent the progression of CKD; in addition, patients should try their best to avoid dehydration, and immediate initiation of allosteric modulator of calcium sensing receptor (e.g., cinacalcet, evocalcet) is recommended until parathyroidectomy is performed after an examination with ultrasound and  $^{99\text{m}}\text{Tc}$ -sestamibi (MIBI) scintigraphy. In the cases with THPT, all recognizable parathyroid glands by ultrasound and MIBI scintigraphy should be removed as persisting THPT and recurrence of THPT after parathyroidectomy is very high (75%) among XLH patients [5]. In patients who developed THPT with contraindication for surgery (e.g., severe cardiopulmonary disorder, oldest-old patients), allosteric modulation of the calcium sensing receptor should be continued, and in these patients, additional anti-osteoporotic treatment should be considered after the surrogate marker of

osteomalacia (e.g., BAP) is well-controlled because increased intact PTH persists with allosteric modulation of the calcium sensing receptor [35]. It is recommended to change the treatment to burosumab in XLH patients with uncontrolled severe SHPT undergoing conventional therapy or who once developed THPT. Supplementation of active vitamin D or replacement therapy with recombinant 1-84 PTH is required in addition to the treatment of hypophosphatemia in patients in which all parathyroid glands have been removed due to THPT. Please refer to Table 2 for guidance for conventional treatment for adults with XLH (Table 2).

**Table 2.** Suggested guidance of conventional therapy for adult XLH patients.

Priority	Event	Action
		Start with active vitamin D
	Initiation of treatment	(0.50 µg b.i.d. for calcitriol or 1.0 µg s.i.d. for alfacalcidol) After 1 to 2 weeks, start phosphate supplementation (800 mg q.i.d.)
	Range of dosage	calcitriol: 0.50 to 0.75 µg b.i.d., alfacalcidol: 0.75 to 1.5 µg s.i.d. phosphate supplementation: 750 to 1,600 mg q.i.d.
	Initial phase	Adjust phosphate supplementation for a period of time by 100 mg Goal: peak phosphate <sup>1</sup> within lower 50% of the reference range; laboratory test: every one to four weeks
	Maintenance phase (after 12 to 18 months)	Adjust phosphate supplementation for a period of time by 100 mg Goal: BAP within the reference range Laboratory test: every 3 months
		Uncontrolled BAP: change the treatment to burosumab
High	Severe SHPT (peak intact PTH ≥ 2x upper limit of the reference range)	Immediately decrease phosphate supplementation by 100 mg or more Increase active vitamin D (0.25 µg daily for calcitriol and 0.25 to 0.5 µg daily for alfacalcidol) After 1 to 2 weeks, increase phosphate supplementation
		Uncontrolled severe SHPT: change the treatment to burosumab
		Immediately quit conventional therapy
High	Development of THPT (hyperparathyroidism with the value of calcium ≥ the middle of the reference range)	Initiate allosteric modulation of calcium sensing receptor Try to prevent dehydration Conduct parathyroidectomy for all recognizable glands, otherwise continue allosteric modulation of the calcium sensing receptor in patients with contraindication for surgery Change the treatment to burosumab afterward
High	Overadjustment of peak phosphate within upper 25% of the reference range or over	Immediately decrease phosphate supplementation by 100 mg or more
High	Progression of CKD to eGFR < 45 ml/min/1.73 m <sup>2</sup>	Decrease conventional therapy Change the treatment to burosumab in patients who still need phosphate supplementation (uncontrolled BAP with active vitamin D).

<sup>1</sup>Peak phosphate: phosphate level 1 to 2 hours after phosphate supplementation

### 6.3. Burosumab

Burosumab was reported to be associated with improvement in the persistent pseudofractures and fractures; the results of PROs in adult XLH patients show that most of them were treated with conventional treatment beforehand [36, 37, 38, 39]. Consequently, the initiation of or change to burosumab should be considered among adult XLH patients with severe symptoms (Table 1). In addition, given that burosumab repairs dysfunctional phosphate metabolism in patients with XLH into a physiologically corrected state by counteracting excess action of FGF23, burosumab should not be associated with

the development of SHPT/THPT and subsequent progression of CKD. Therefore, the initiation of burosumab should also be considered in patients with CKD (e.g., eGFR < 45 ml/min/1.73 m<sup>2</sup>), and a change in treatment from conventional therapy to burosumab should be considered among adult XLH patients with uncontrolled severe SHPT or a history of the development of THPT (Table 1). However, medical economic efficacy should always be considered independently for this kind of expensive orphan drug according to the guidelines/recommendations or the health care system of each country [2].

The initial dosage of burosumab for adult XLH patients is decided worldwide to be 1.0 mg/kg body weight (maximum dose of 90 mg) by subcutaneous injection every four weeks, and concomitant use of active vitamin D and phosphate supplementation are not recommended or prohibited unless other coexisting medical conditions require active vitamin D (e.g., hypoparathyroidism after parathyroidectomy of all parathyroid glands). In the initial phase, the response to burosumab should be confirmed 1 to 2 weeks after the injection (peak phosphate), as the phosphate level returned to the baseline four weeks after the last injection (trough phosphate) in a considerable number of patients. Trough phosphate level should be controlled within the lower half of the reference range or less, and dosage of burosumab should be decreased by 0.2 to 0.3 mg/kg in patients with trough phosphate levels within the higher half of the reference range and over. Adverse effects tightly associated with the use of burosumab are injection site reactions, and there is no report of the development of antagonizing antibodies [36, 37, 38, 39].

When no response or only a trivial response is observed at 1 to 2 weeks after injection (peak phosphate), coexisting vitamin D deficiency should be examined by measuring 25OHD. If the patient develops uncontrolled severe SHPT with conventional therapy and the treatment is changed to burosumab, the response to burosumab (peak phosphate) might improve after 3 to 6 injections as intact PTH decreases because stimulation of *FGF23* transcription by PTH is alleviated [26]. THPT is also associated with a diminished response to burosumab, and immediate initiation of allosteric modulators of calcium sensing receptors (e.g., cinacalcet, evocalcet) and subsequent parathyroidectomy should be recommended along with the careful prevention of dehydration. If no response or trivial response to burosumab continues after these problems are ruled out or adequately addressed, changing the treatment to conventional therapy should be considered. Please refer to Table 3 for guidance for burosumab treatment for adults with XLH (Table 3).

**Table 3.** Suggested guidance for the use of burosumab for adult XLH patients.

Priority	Event	Action
	Initiation of treatment	1.0 mg/kg body weight (up to 90 mg) subcutaneous injection every four weeks Response to the burosumab should be confirmed by peak phosphate <sup>1</sup> ; concomitant use of active vitamin D and phosphate supplementation are not recommended or prohibited unless another medical condition requires it
	Trough phosphate <sup>2</sup> within the higher half of the reference range or over	Decrease burosumab dose by 0.2 to 0.3 mg/kg Subsequently, fine tune burosumab dose by 0.1 mg/kg to target trough phosphate within lower half of the reference range
	No or trivial response at peak phosphate	Rule out vitamin D deficiency by measuring 25OHD severe SHPT: continue burosumab for 3 to 6 times and confirm the improvement in peak phosphate Continuing no response or trivial response after problems above are ruled out or addressed; change the treatment to conventional therapy Initiate allosteric modulation of the calcium sensing receptor
High	THPT	Try to prevent dehydration Conduct parathyroidectomy for all recognizable glands, otherwise continue allosteric modulation of the calcium sensing receptor in patients with contraindication for surgery

<sup>1</sup>Peak phosphate: phosphate level 1 to 2 weeks after the last injection of burosumab

<sup>2</sup>Trough phthalate: phosphate level 4 weeks after the last injection of burosumab

**7. Remaining problems and future research topics in adult XLH patients**

The etiology for the development of enthesopathy needs to be elucidated to develop a treatment option for it, as this complication remarkably debilitates ADL and QOL in a large number of adult XLH patients. The association among the development of severe SHPT or THPT, progression of CKD, the highest dosage, cumulative dosage of phosphate supplementation, or dosage of active vitamin D needs to be elucidated to create more detailed guidelines for conventional therapy. The mode of administration and dosage of burosumab should be reconsidered, as some of the adult XLH patients are obviously under-treated with the current dosage (maximum 1.0 mg/kg body weight) and mode of administration (once every four weeks).

**8. Conclusion**

The development of burosumab was a game changer in the treatment of adult XLH patients as conventional therapy is associated with undertreatment of osteomalacia and the risk of the developing severe SHPT or THPT and the progression of CKD. However, there are remaining problems to be addressed specifically among adults with XLH; of these, the complication of debilitating enthesopathy warrants special attention and must be addressed urgently.

**Disclosure Summary:** NI receives research support from Kyowa Kirin Co., Ltd.

**References**

1. Carpenter TO, Imel EA, Holm IA, Jan de Beur SM, Insogna KL. A clinician's guide to X-linked hypophosphatemia. J Bone Miner Res. 2011 Jul;26(7):1381-8. doi: 10.1002/jbmr.340. Epub 2011 May 2. Erratum in: J Bone Miner Res. 2015 Feb;30(2):394. PMID: 21538511; PMCID: PMC3157040.

2. Haffner D, Emma F, Eastwood DM, Duplan MB, Bacchetta J, Schnabel D, Wicart P, Bockenhauer D, Santos F, Levtschenko E, Harvengt P, Kirchhoff M, Di Rocco F, Chaussain C, Brandi ML, Savendahl L, Briot K, Kamenicky P, Rejnmark L, Linglart A. Clinical practice recommendations for the diagnosis and management of X-linked hypophosphataemia. Nat Rev Nephrol. 2019 Jul;15(7):435-455. doi: 10.1038/s41581-019-0152-5. PMID: 31068690; PMCID: PMC7136170.

3. Cheung M, Roschger P, Klaushofer K, Veilleux LN, Roughley P, Glorieux FH, Rauch F. Cortical and trabecular bone density in X-linked hypophosphatemic rickets. *J Clin Endocrinol Metab.* 2013 May;98(5):E954-61. doi: 10.1210/jc.2012-4133. Epub 2013 Mar 26. PMID: 23533226.
4. Kato H, Koga M, Kinoshita Y, Taniguchi Y, Kobayashi H, Fukumoto S, Nangaku M, Makita N, Ito N. Incidence of Complications in 25 Adult Patients With X-linked Hypophosphatemia. *J Clin Endocrinol Metab.* 2021 Aug 18;106(9):e3682-e3692. doi: 10.1210/clinem/dgab282. PMID: 33912912.
5. DeLacey S, Liu Z, Broyles A, El-Azab SA, Guandique CF, James BC, Imel EA. Hyperparathyroidism and parathyroidectomy in X-linked hypophosphatemia patients. *Bone.* 2019 Oct;127:386-392. doi: 10.1016/j.bone.2019.06.025. Epub 2019 Jul 2. PMID: 31276850; PMCID: PMC6836672.
6. Fukumoto S, Ozono K, Michigami T, Minagawa M, Okazaki R, Sugimoto T, Takeuchi Y, Matsumoto T. Pathogenesis and diagnostic criteria for rickets and osteomalacia--proposal by an expert panel supported by the Ministry of Health, Labour and Welfare, Japan, the Japanese Society for Bone and Mineral Research, and the Japan Endocrine Society. *J Bone Miner Metab.* 2015 Sep;33(5):467-73. doi: 10.1007/s00774-015-0698-7. Epub 2015 Jul 22. PMID: 26197863.
7. Ito N, Kubota T, Kitanaka S, Fujiwara I, Adachi M, Takeuchi Y, Yamagami H, Kimura T, Shinoda T, Minagawa M, Okazaki R, Ozono K, Seino Y, Fukumoto S. Clinical performance of a novel chemiluminescent enzyme immunoassay for FGF23. *J Bone Miner Metab.* 2021 Nov;39(6):1066-1075. doi: 10.1007/s00774-021-01250-1. Epub 2021 Jul 13. PMID: 34255195..
8. Kato H, Hidaka N, Koga M, Ogawa N, Takahashi S, Miyazaki H, Nangaku M, Makita N, Ito N. Performance evaluation of the new chemiluminescent intact FGF23 assay relative to the existing assay system. *J Bone Miner Metab.* 2022 Jan;40(1):101-108. doi: 10.1007/s00774-021-01258-7. Epub 2021 Aug 5. PMID: 34351500.
9. Kinoshita Y, Saito T, Shimizu Y, Hori M, Taguchi M, Igarashi T, Fukumoto S, Fujita T. Mutational analysis of patients with FGF23-related hypophosphatemic rickets. *Eur J Endocrinol.* 2012 Aug;167(2):165-72. doi: 10.1530/EJE-12-0071. Epub 2012 May 10. PMID: 22577109.
10. Minisola S, Peacock M, Fukumoto S, Cipriani C, Pepe J, Tella SH, Collins MT. Tumour-induced osteomalacia. *Nat Rev Dis Primers.* 2017 Jul 13;3:17044. doi: 10.1038/nrdp.2017.44. PMID: 28703220.
11. Shimizu Y, Tada Y, Yamauchi M, Okamoto T, Suzuki H, Ito N, Fukumoto S, Sugimoto T, Fujita T. Hypophosphatemia induced by intravenous administration of saccharated ferric oxide: another form of FGF23-related hypophosphatemia. *Bone.* 2009 Oct;45(4):814-6. doi: 10.1016/j.bone.2009.06.017. Epub 2009 Jun 23. PMID: 19555782.
12. Hidaka N, Kato H, Koga M, Katsura M, Oyama Y, Kinoshita Y, Fukumoto S, Makita N, Nangaku M, Ito N. Induction of FGF23-related hypophosphatemic osteomalacia by alcohol consumption. *Bone Rep.* 2021 Oct 16;15:101144. doi: 10.1016/j.bonr.2021.101144. PMID: 34901334; PMCID: PMC8640868.
13. Veilleux LN, Cheung MS, Glorieux FH, Rauch F. The muscle-bone relationship in X-linked hypophosphatemic rickets. *J Clin Endocrinol Metab.* 2013 May;98(5):E990-5. doi: 10.1210/jc.2012-4146. Epub 2013 Mar 22. PMID: 23526465.
14. Ito N, Fukumoto S, Takeuchi Y, Takeda S, Suzuki H, Yamashita T, Fujita T. Effect of acute changes of serum phosphate on fibroblast growth factor (FGF)23 levels in humans. *J Bone Miner Metab.* 2007;25(6):419-22. doi: 10.1007/s00774-007-0779-3. Epub 2007 Oct 25. PMID: 17968495.
15. Beck-Nielsen SS, Brusgaard K, Rasmussen LM, Brixen K, Brock-Jacobsen B, Poulsen MR, Vestergaard P, Ralston SH, Albagha OM, Poulsen S, Haubek D, Gjørup H, Hintze H, Andersen MG, Heickendorff L, Hjelmborg J, Gram J. Phenotype presentation of hypophosphatemic rickets in adults. *Calcif Tissue Int.* 2010 Aug;87(2):108-19. doi: 10.1007/s00223-010-9373-0. Epub 2010 Jun 4. PMID: 20524110.
16. Ye L, Liu R, White N, Alon US, Cobb CM. Periodontal status of patients with hypophosphatemic rickets: a case series. *J Periodontol.* 2011 Nov;82(11):1530-5. doi: 10.1902/jop.2011.100736. Epub 2011 Mar 21. PMID: 21417586.
17. Biosse Duplan M, Coyac BR, Bardet C, Zadikian C, Rothenbuhler A, Kamenicky P, Briot K, Linglart A, Chaussain C. Phosphate and Vitamin D Prevent Periodontitis in X-Linked Hypophosphatemia. *J Dent Res.* 2017 Apr;96(4):388-395. doi: 10.1177/0022034516677528. Epub 2016 Nov 13. PMID: 27821544.
18. Connor J, Olear EA, Insogna KL, Katz L, Baker S, Kaur R, Simpson CA, Sterpka J, Dubrow R, Zhang JH, Carpenter TO. Conventional Therapy in Adults With X-Linked Hypophosphatemia: Effects on Enthesopathy and Dental Disease. *J Clin Endocrinol Metab.* 2015 Oct;100(10):3625-32. doi: 10.1210/JC.2015-2199. Epub 2015 Jul 15. PMID: 26176801; PMCID: PMC4596038.
19. Steele A, Gonzalez R, Garbalosa JC, Steigbigel K, Grgurich T, Parisi EJ, Feinn RS, Tommasini SM, Macica CM. Osteoarthritis, Osteophytes, and Enthesophytes Affect Biomechanical Function in Adults With X-linked Hypophosphatemia. *J Clin Endocrinol Metab.* 2020 Apr 1;105(4):e1798-814. doi: 10.1210/clinem/dgaa064. PMID: 32047911; PMCID: PMC8416779.
20. Karaplis AC, Bai X, Falet JP, Macica CM. Mineralizing enthesopathy is a common feature of renal phosphate-wasting disorders attributed to FGF23 and is exacerbated by standard therapy in hyp mice. *Endocrinology.* 2012 Dec;153(12):5906-17. doi: 10.1210/en.2012-1551. Epub 2012 Oct 4. PMID: 23038738; PMCID: PMC3512070.
21. Saito T, Shimizu Y, Hori M, Taguchi M, Igarashi T, Fukumoto S, Fujita T. A patient with hypophosphatemic rickets and ossification of posterior longitudinal ligament caused by a novel homozygous mutation in ENPP1 gene. *Bone.* 2011 Oct;49(4):913-6. doi: 10.1016/j.bone.2011.06.029. Epub 2011 Jul 2. PMID: 21745613.
22. Kato H, Ansh AJ, Lester ER, Kinoshita Y, Hidaka N, Hoshino Y, Koga M, Taniguchi Y, Uchida T, Yamaguchi H, Niida Y, Nakazato M, Nangaku M, Makita N, Takamura T, Saito T, Braddock DT, Ito N. Identification of ENPP1 Haploinsufficiency in Patients With Diffuse Idiopathic Skeletal Hyperostosis and Early-Onset Osteoporosis. *J Bone Miner Res.* 2022 Mar 26. doi: 10.1002/jbmr.4550. Epub ahead of print. PMID: 35340077.



23. Ferreira CR, Kintzinger K, Hackbarth ME, Botschen U, Nitschke Y, Mughal MZ, Baujat G, Schnabel D, Yuen E, Gahl WA, Gafni RI, Liu Q, Huertas P, Khursigara G, Rutsch F. Ectopic Calcification and Hypophosphatemic Rickets: Natural History of ENPP1 and ABCC6 Deficiencies. *J Bone Miner Res.* 2021 Nov;36(11):2193-2202. doi: 10.1002/jbmr.4418. Epub 2021 Aug 16. PMID: 34355424; PMCID: PMC8595532.
24. Bernhard E, Nitschke Y, Khursigara G, Sabbagh Y, Wang Y, Rutsch F. A Reference Range for Plasma Levels of Inorganic Pyrophosphate in Children Using the ATP Sulfurylase Method. *J Clin Endocrinol Metab.* 2022 Jan 1;107(1):109-118. doi: 10.1210/clinem/dgab615. PMID: 34498693; PMCID: PMC8684482.
25. Maulding ND, Kavanagh D, Zimmerman K, Coppola G, Carpenter TO, Jue NK, Braddock DT. Genetic pathways disrupted by ENPP1 deficiency provide insight into mechanisms of osteoporosis, osteomalacia, and paradoxical mineralization. *Bone.* 2021 Jan;142:115656. doi: 10.1016/j.bone.2020.115656. Epub 2020 Sep 24. PMID: 32980560; PMCID: PMC7744330.
26. Ito N, Prideaux M, Wijenayaka AR, Yang D, Ormsby RT, Bonewald LF, Atkins GJ. Sclerostin Directly Stimulates Osteocyte Synthesis of Fibroblast Growth Factor-23. *Calcif Tissue Int.* 2021 Jul;109(1):66-76. doi: 10.1007/s00223-021-00823-6. Epub 2021 Feb 22. PMID: 33616712.
27. Kubota T, Fukumoto S, Cheong HI, Michigami T, Namba N, Ito N, Tokunaga S, Gibbs Y, Ozono K. Long-term outcomes for Asian patients with X-linked hypophosphataemia: rationale and design of the SUNFLOWER longitudinal, observational cohort study. *BMJ Open.* 2020 Jun 29;10(6):e036367. doi: 10.1136/bmjopen-2019-036367. PMID: 32601114; PMCID: PMC7328740.
28. Nakamura Y, Takagi M, Takeda R, Miyai K, Hasegawa Y. Hypertension is a characteristic complication of X-linked hypophosphatemia. *Endocr J.* 2017 Mar 31;64(3):283-289. doi: 10.1507/endocrj.EJ16-0199. Epub 2016 Dec 27. PMID: 28025445.
29. Skrinar A, Dvorak-Ewell M, Evins A, Macica C, Linglart A, Imel EA, Theodore-Oklota C, San Martin J. The Lifelong Impact of X-Linked Hypophosphatemia: Results From a Burden of Disease Survey. *J Endocr Soc.* 2019 May 7;3(7):1321-1334. doi: 10.1210/js.2018-00365. PMID: 31259293; PMCID: PMC6595532.
30. Seefried L, Smyth M, Keen R, Harvengt P. Burden of disease associated with X-linked hypophosphataemia in adults: a systematic literature review. *Osteoporos Int.* 2021 Jan;32(1):7-22. doi: 10.1007/s00198-020-05548-0. Epub 2020 Jul 24. PMID: 32710160; PMCID: PMC7755619.
31. Ito N, Kang HG, Nishida Y, Evins A, Skrinar A, Cheong HI. Burden of disease of X-linked hypophosphatemia in Japanese and Korean patients: a cross-sectional survey. *Endocr J.* 2022 Apr 28;69(4):373-383. doi: 10.1507/endocrj.EJ21-0386. Epub 2021 Nov 2. PMID: 34732603.
32. Dahir K, Dhaliwal R, Simmons J, Imel EA, Gottesman GS, Mahan JD, Prakasam G, Hoch AI, Ramesan P, Díaz-González de Ferris M. Health Care Transition From Pediatric- to Adult-Focused Care in X-linked Hypophosphatemia: Expert Consensus. *J Clin Endocrinol Metab.* 2022 Feb 17;107(3):599-613. doi: 10.1210/clinem/dgab796. PMID: 34741521; PMCID: PMC8852209.
33. Horn A, Wright J, Bockenhauer D, Van't Hoff W, Eastwood DM. The orthopaedic management of lower limb deformity in hypophosphataemic rickets. *J Child Orthop.* 2017 Aug 1;11(4):298-305. doi: 10.1302/1863-2548.11.170003. PMID: 28904636; PMCID: PMC5584499.
34. Bettinelli A, Bianchi ML, Mazzucchi E, Gandolini G, Appiani AC. Acute effects of calcitriol and phosphate salts on mineral metabolism in children with hypophosphatemic rickets. *J Pediatr.* 1991 Mar;118(3):372-6. doi: 10.1016/s0022-3476(05)82149-3. PMID: 1847972.
35. Peacock M, Bolognese MA, Borofsky M, Scumpia S, Sterling LR, Cheng S, Shoback D. Cinacalcet treatment of primary hyperparathyroidism: biochemical and bone densitometric outcomes in a five-year study. *J Clin Endocrinol Metab.* 2009 Dec;94(12):4860-7. doi: 10.1210/jc.2009-1472. Epub 2009 Oct 16. PMID: 19837909.
36. Insogna KL, Briot K, Imel EA, Kamenický P, Ruppe MD, Portale AA, Weber T, Pitukcheewanont P, Cheong HI, Jan de Beur S, Imanishi Y, Ito N, Lachmann RH, Tanaka H, Perwad F, Zhang L, Chen CY, Theodore-Oklota C, Mealiffe M, San Martin J, Carpenter TO; AXLES 1 Investigators. A Randomized, Double-Blind, Placebo-Controlled, Phase 3 Trial Evaluating the Efficacy of Burosumab, an Anti-FGF23 Antibody, in Adults With X-Linked Hypophosphatemia: Week 24 Primary Analysis. *J Bone Miner Res.* 2018 Aug;33(8):1383-1393. doi: 10.1002/jbmr.3475. Epub 2018 Jun 26. PMID: 29947083.
37. Insogna KL, Rauch F, Kamenický P, Ito N, Kubota T, Nakamura A, Zhang L, Mealiffe M, San Martin J, Portale AA. Burosumab Improved Histomorphometric Measures of Osteomalacia in Adults with X-Linked Hypophosphatemia: A Phase 3, Single-Arm, International Trial. *J Bone Miner Res.* 2019 Dec;34(12):2183-2191. doi: 10.1002/jbmr.3843. Epub 2019 Oct 1. PMID: 31369697; PMCID: PMC6916280.
38. Portale AA, Carpenter TO, Brandi ML, Briot K, Cheong HI, Cohen-Solal M, Crowley R, Jan De Beur S, Eastell R, Imanishi Y, Imel EA, Ing S, Ito N, Javaid M, Kamenicky P, Keen R, Kubota T, Lachmann R, Perwad F, Pitukcheewanont P, Ralston SH, Takeuchi Y, Tanaka H, Weber TJ, Yoo HW, Zhang L, Theodore-Oklota C, Mealiffe M, San Martin J, Insogna K. Continued Beneficial Effects of Burosumab in Adults with X-Linked Hypophosphatemia: Results from a 24-Week Treatment Continuation Period After a 24-Week Double-Blind Placebo-Controlled Period. *Calcif Tissue Int.* 2019 Sep;105(3):271-284. doi: 10.1007/s00223-019-00568-3. Epub 2019 Jun 4. PMID: 31165191.
39. Briot K, Portale AA, Brandi ML, Carpenter TO, Cheong HI, Cohen-Solal M, Crowley RK, Eastell R, Imanishi Y, Ing S, Insogna K, Ito N, Jan de Beur S, Javaid MK, Kamenicky P, Keen R, Kubota T, Lachmann RH, Perwad F, Pitukcheewanont P, Ralston SH, Takeuchi Y, Tanaka H, Weber TJ, Yoo HW, Nixon A, Nixon M, Sun W, Williams A, Imel EA. Burosumab treatment in adults with X-linked hypophosphataemia: 96-week patient-reported outcomes and ambulatory function from a randomised phase 3 trial and open-label extension. *RMD Open.* 2021 Sep;7(3):e001714. doi: 10.1136/rmdopen-2021-001714. PMID: 34548383; PMCID: PMC8458321.