Impact of Early Conventional Treatment on Adult Bone and Joints in a Murine Model of X-Linked Hypophosphatemia

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X-linked hypophosphatemia (XLH) is the most common form of genetic rickets. Mainly diagnosed during childhood because of growth retardation and deformities of the lower limbs, the disease affects adults with early enthesopathies and joint structural damage that significantly alter patient quality of life. The conventional treatment, based on phosphorus supplementation and active vitamin D analogs, is commonly administered from early childhood to the end of growth; unfortunately, it does not allow complete recovery from skeletal damage. Despite adequate treatment during childhood, bone and joint complications occur in adults and become a dominant feature in the natural history of the disease. Our previous data showed that the Hyp mouse is a relevant model of XLH for studying early enthesophytes and joint structural damage. Here, we studied the effect of conventional treatment on the development of bone and joint alterations in this mouse model during growth and young adulthood. Mice were supplemented with oral phosphorus and calcitriol injections, following two timelines: (i) from weaning to 3 months of age and (ii) from 2 to 3 months to evaluate the effects of treatment on the development of early enthesophytes and joint alterations, and on changes in bone and joint deformities already present, respectively. We showed that early conventional treatment improved bone microarchitecture, and partially prevented bone and joint complications, but with no noticeable improvement in enthesophytes. In contrast, later administration had limited efficacy in ameliorating bone and joint alterations. Despite the improvement in bone microarchitecture, the conventional treatment, early or late, had no effect on osteoid accumulation. Our data underline the usefulness of the Hyp murine model for preclinical studies on skeletal and extraskeletal lesions. Although the early
INTRODUCTION

X-linked hypophosphatemia (XLH) is the most common form of genetic rickets. This rare disease is caused by inactivating mutations in the phosphate-regulating endopeptidase homolog X-linked (PHEX) gene, characterized by chronic hypophosphatemia. Impaired function of PHEX leads to elevated levels of phosphaturic fibroblast growth factor 23 (FGF23) resulting in renal phosphate-wasting hypophosphatemia and low levels of calcitriol \([1,25(OH)\text{D}_3]\) via the inhibition of 1α-hydroxylase and the activation of 24-hydroxylase (Kinoshita and Fukumoto, 2018).

Clinically, children with XLH are characterized by progressive skeletal deformities (leg bowing, waddling gait, poor growth, and disproportional short stature), dental abscesses, and craniosynostosis. Adult patients present various symptoms of osteomalacia such as bone pain, insufficiency fractures, and myopathy. In addition, adults may develop hearing loss, odontomalaia, mineralizing enthesopathy, and osteoarthropathy (Linglart et al., 2014). In adult patients with XLH, the aforementioned manifestations significantly reduce quality of life (Che et al., 2016; Steele et al., 2020). We showed that Hyp mice, a murine model of XLH, developed early osteoarticular lesions and the severity of these lesions gradually increased over 12 months, demonstrating the relevance of this murine model for osteoarticular preclinical studies (Faraji-Bellee et al., 2020).

Current medical treatment of XLH consists of oral active vitamin D [calcitriol or 1α-(OH)\text{D}_3] and multiple daily doses of phosphate supplements. To optimize the final outcomes (recovery of rickets, normalization of elevated alkaline phosphatase (ALP) levels, growth improvement, restoration of leg deformities, and dental mineralization), treatment should be started as soon as the diagnosis of XLH is made. Supplementation is commonly prescribed from early childhood to the end of growth but is also essential in certain periods of adult life such as pregnancy or breastfeeding, before planned surgical interventions, and in all symptomatic patients with XLH (recurrent dental abscesses, fractures, etc.) (Linglart et al., 2014). Nonetheless, despite this treatment during growth, musculoskeletal symptoms due to enthesopathies and osteoarthritis remain the major manifestations in the clinical progression of XLH. Further, there is a paucity of data on the effects of conventional treatment (i) if started early, on the abnormal bone phenotype in XLH, beyond the traditional goals, and (ii) if started early, on prevention of and/or recovery from osteoarticular manifestations of XLH such as osteoarthritis and enthesopathies. The main limitations of the studies on the effect of conventional treatment in XLH performed so far are small sample sizes and their retrospective cross-sectional observational design that does not take into account the age when the treatment is started.

Therefore, we designed a prospective study in a murine model of XLH (Hyp mice) aiming to evaluate whether, if started early in life, conventional treatment is capable of preventing and/or ameliorating the skeletal and extraskeletal manifestations of hypophosphatemia.

MATERIALS AND METHODS

Mice

The Hyp mouse model B6.Cg-Phex Hyp/1 was used in this study. Heterozygous breeding was carried out and tail snips were collected for genotyping. DNA was extracted from the snips using DNeasy Blood and Tissue Kit (Qiagen, France) and the genotype was determined by PCR using primers for the Phex gene. Wildtype (WT) and Hyp littermate male mice were used in experimental procedures. All experiments were performed with a protocol approved by the Animal Care Committee of the Université de Paris (project agreement 20-008, APAFiS #27827 N°202001171429974). Animals were maintained in accordance with the ethical protocol approved by the Animal Care Committee of French Veterinary Services (DPP Haut de Seine, France: agreement number D9204901). All mice were housed under standard conditions of temperature (23 ± 2°C) with a 12:12 h light-dark cycle and unlimited access to water and standard pelleted food (1.20% calcium and 0.83% phosphorus, rodent diet 3800PMS10, Proviimi Kliwa, Kaiseraugst, Switzerland).

Two groups of treated Hyp mice were carried out in this study \((n = 6\text{ mice per group})\): (1) to study the effect of long-term treatment on skeletal/extraskeletal manifestations if started early during growth, Hyp mice received the conventional treatment during the whole study period from the juvenile stage starting at weaning, which occurs at 3 weeks (W), to the beginning of the mature adult stage which occurs at 3 months (M) of life and (2) Hyp mice which received the conventional treatment from M2 (corresponding to the end of juvenile stage) to M3 to study the effect of the conventional treatment on skeletal/extraskeletal manifestations if started later during growth. Both treated groups of Hyp mice were compared to control WT and Hyp mice, which were not given conventional treatment \((n = 6\text{ per group})\).

The conventional treatment consisted of intraperitoneal injections of calcitriol 175 pg/g every other day \([1,25(OH)\text{D}_3]\), Cayman Laboratory] and phosphate supplementation (phosphate-enriched water 1.93 g phosphate element per
liter of beverage, Phosphoneuro). Doses of calcitriol were adjusted once a week according to the animals' weight.

**In vivo Study**

**The Growth Parameters**

Growth parameters (body weight and total length) were measured once a week. Precision scales and a graduated ruler were used for weight and length measurements, respectively. Additionally, the length of the rachis was measured on X-ray micro-computed tomography (Micro-CT) images.

**X-Ray Micro-computed Tomography Analysis**

Wildtype and *Hyp* mice were scanned at W3, M2, and M3 using a high-resolution X-ray micro-CT system (Quantum FX Caliper, Life Sciences, Perkin Elmer, Waltham, MA, United States) hosted by the PIV Platform (UR2496, Montrouge, France). Standard acquisition settings were applied (setting the voltage at 90 kV and intensity at 160 mA), and scans were performed with a field of view alternatively focused on the right paw (scan time of 180 s and voxel size of 20 μm3), focused on the hip (120 s and 50 μm3), or covering the full body (36 s and 236 μm3). Micro-CT datasets were analyzed using the built-in multilplanar reconstruction tool, OsiriX 5.8 (Pixmeo, Switzerland), to obtain time series of images aligned anatomically for each region of each animal.

Axial and coronal images of the sacroiliac and hip joints, sagittal images of the spine, and axial and sagittal images of the hind paw were reconstructed. The following were evaluated: hip osteoarthritis (defined as the presence of osteophytes on joint margins, narrowing of the joint space or altered shape of the bone ends); enthesopathies (defined as new bone formation at enthesis sites) on the iliac bone, spine or paw; erosion of the sacroiliac joint and periarticular calcification. The analysis was focused on these areas because they are the most frequent sites of structural involvement in adults with XLH. Erosion of the sacroiliac joints was assessed following Faraji-Belle et al. (2020) protocol, which was developed by rheumatologists specialized in the field of rare bone diseases and bone inflammatory diseases. The reader was blind to the status of the mouse (*Hyp* vs WT) but was aware of the different analysis time points (W3, M2, M3). A semi-quantitative score was established, ranging from 0 (normal) to 3 (most severe feature assessed) for sacroiliac erosions (see Supplementary Table 1).

The angle of dorsolumbar kyphosis of mice was defined for each mouse at W3, M2, and M3. Using sagittal images of mice spines from full-body CT scans, endplate orientations of thoracic and lumbar vertebrae have been marked using ImageJ (Rasband, W.S., ImageJ, U.S. National Institutes of Health, Bethesda, MD, United States1 1997–2016). The apical thoracic vertebrae of the rachis were identified and the angle of kyphosis defined by the means of (1) the tangent to the lower vertebral endplate of the fourth lower vertebral and (2) the tangent of the upper vertebral endplate of the fourth upper vertebral, from the apical vertebra. A script in MATLAB (MATLAB R2012b, The MathWorks Inc., Natick, MA, United States, 2000) was used to measure these angles at each age in each of the mice.

The trabecular bone was analyzed at the distal metaphysis of the femur. The following parameters were used: bone volume/total volume (BV/TV) ratio, trabecular number (TbN), trabecular separation (TbSp), trabecular thickness (TbTh), and trabecular pattern factor (TbPf) (Bouxsein et al., 2010).

**Ex vivo Study**

**Murine Bone Tissue Preparation**

Bones were fixed overnight at 4°C in 70% ethanol solution and dehydrated in a graded ethanol series. Undecalcified samples were embedded in methyl methacrylate (Merck, Rahway, NJ, United States). Serial sections, 4 μm thick, were cut on a microtome (Polycut E microtome, Leica, Wetzlar, Germany). Series of consecutive sections representative of micro-CT images were stained with von Kossa (5% silver nitrate solution, Sigma–Aldrich, St Louis, MO, United States) and counterstained with toluidine blue (pH 3.8), stained with Masson’s Trichrome (Sigma–Aldrich), Safranin O Lillie’s Trichrome (Sigma–Aldrich).

**Enzyme Histochemistry**

Tartrate-resistant acid phosphatase (TRAP) was used to evaluate osteoclasts by using 2.5 mM naphthol-ASTR-phosphate (Sigma–Aldrich), 0.36 M N-N-dimethyl-formamide (Sigma–Aldrich), and 4 mM salt in pH 5.2 acetate buffer. Non-osteoclastic acid phosphatase activity was inhibited with 100 mM L(+)-tartaric acid (Sigma–Aldrich) added to the substrate solution.

Alkaline phosphatase was used to reveal the layer of osteogenic cells by incubating the sections with naphthol ASTR phosphate (Sigma–Aldrich) and diazonium fast blue RR salt (Sigma–Aldrich) for 30 min at 37°C (pH 9) in the presence of MgCl2.

**Immunohistochemistry**

Sections embedded in methyl methacrylate were deplasticized in methyl glycol acetate. After rehydration in a graded ethanol series to pure distilled water, non-specific peroxidoxes were blocked for 15 min with ortho-peroxicid acid and background activity was blocked at room temperature using 5% bovine serum albumin (BSA). Sections were then incubated in a humid atmosphere for 12 h at room temperature in a dark chamber with primary antibody against sclerostin (SOST) (R&D Systems, Minneapolis, MN, United States) diluted at 5 μg/mL. Sections were washed and then incubated with polyclonal anti-antigoat immunoglobulin (Dakocytomation) diluted at 1/200 for 1 h at room temperature in a dark chamber. Peroxidase activity was detected using diaminobenzidine (DAB) substrate (Sigma–Aldrich). Control incubations to assess non-specific staining consisted of the same procedure except that the primary antibody was replaced by non-immune serum.

**Statistical Analysis**

Statistical analysis was carried out and graphs plotted with GraphPad Prism for Windows, version 7.0. The distribution of variables was tested with Kolmogorov–Smirnov test. The results are expressed as the mean ± SD for continuous variables and comparisons being performed using ANOVA. Data are expressed as median with interquartile range in Figures 4B,C, and mean ± SD in Figure 4D using Mann–Whitney and Student’s
RESULTS

Effect of Conventional Treatment on Growth and Dorsolumbar Kyphosis

Seeking to understand the effect of the conventional treatment on parameters such as growth and dorsolumbar kyphosis, we analyzed changes in spine length and changes in dorsolumbar kyphosis (expressed in degrees of curvature) by group (Figure 1).

Early Start of Conventional Treatment

At baseline, Hyp mice had significantly shorter spine lengths than WT animals (Figure 1B). After starting the treatment, Hyp mice in the early treatment group showed much larger gains in length than WT or untreated Hyp mice (Figure 1B). On the other hand, micro-CT analysis showed that at M3, WT mice were significantly longer than treated Hyp mice, and there were no significant length differences between treated and untreated Hyp mice (Figure 1B).

Regarding dorsolumbar kyphosis, spinal kyphosis increased between baseline and the end of the study in untreated Hyp mice whereas it decreased over time in the WT group. No significant difference in spine curvature was observed at M3, i.e., at the end of the study, between the Hyp mice treated early and WT groups (Figure 1B).

Late Start of Conventional Treatment

Hyp mice did not show a significant increase in length after initiating the treatment at M2, and no differences in spine length were seen between treated and untreated Hyp mice at M3 (Figure 1C). Stature growth was not statistically enhanced by conventional treatment when initiated as an adult. Nevertheless, at M3, Hyp mice with treatment showed a gain in length comparable to that in WT animals (Figure 1C).

Regarding dorsolumbar kyphosis, though not statistically significant, spinal kyphosis tended to decrease in Hyp mice after initiating treatment, following the same pattern as in the WT group. In contrast, in untreated Hyp mice, kyphosis increased over time (Figure 1C).
**Hyp** mice with late treatment showed a spine length and curvature comparable to **Hyp** mice with early treatment (Figure 1D).

### Effect of Conventional Treatment on Bone Microarchitecture

#### Early Start of Conventional Treatment

3D reconstructed images demonstrated that treated **Hyp** mice had bone microstructure very similar to that in the WT group by the end of study (M3) (Figure 2A). Regarding the parameters of bone quantity, both BV/TV and TbN were significantly higher ($p < 0.05$) by M2 in **Hyp** mice with early treatment than in untreated animals, and this difference was maintained at M3 (Figure 2B). There were no differences in TbTh and TbPf between statistical significance was not reached ($p < 0.05$) in treated than untreated **Hyp** mice, though this difference was only observed at M3 (Supplementary Figure 1A).

#### Late Start of Conventional Treatment

3D reconstructed images demonstrated a slightly greater bone mass in treated than untreated **Hyp** mice, although the bone microstructure in treated **Hyp** mice was far from that in the WT group (Figure 2A). The study of bone microarchitecture showed that there were no statistically significant differences in BV/TV, TbN, TbTh, or TbPf between treated and untreated **Hyp** mice by the end of study (M3) (Figure 2C and Supplementary Figure 1B).

Compared to **Hyp** mice in the early treatment group, **Hyp** mice treated late showed lower BV/TV and TbN at M3, although the statistical significance was not reached (Figure 2D). There were no differences in TbTh and TbPf between **Hyp** mice with early or late treatment groups (Supplementary Figure 1C).

### Effect of Conventional Treatment on Bone and Joint Structural Damages

#### Bone and Joint Alterations in the Axial Skeleton (Sacroiliac Joint)

**Early start of conventional treatment**

Micro-CT images, performed at baseline (W3), showed alterations in the sacroiliac joint in **Hyp** mice, in comparison to WT (Figure 3A). Two out of six untreated **Hyp** mice and two out of six **Hyp** mice in the early treatment group already displayed a high sacroiliac joint score for erosion at baseline (score between 2 and 2.5 out of 3 which means $<25\%$ and $\geq 25\%$ to $<50\%$ of the articular surface area affected, respectively) (Supplementary Table 2). At M2 and M3, multiple erosions and an irregular and blurred appearance of the cortical margins were noted in untreated **Hyp** mice, whereas **Hyp** mice started on the conventional treatment early showed fewer erosions and a more regular appearance of the sacroiliac joint, compared to that observed in WT animals (Figure 3A). These results were confirmed by the sacroiliac joint scores for erosion at M3 which were lower in **Hyp** mice with early treatment compared to untreated **Hyp** mice. At M3, three of six **Hyp** mice with early treatment had a mean score of 0, whereas none of untreated **Hyp** mice had such a mean score (Figures 3B-D and Supplementary Table 2).

**Late start of conventional treatment**

Multiple erosions and irregular and blurred cortical margins of sacroiliac joints were noticed on micro-CT either at M2 or M3 in untreated **Hyp** mice, in comparison to WT animals. **Hyp** mice given treatment, even when initiated late, showed a slight trend of amelioration of these alterations present at M2 before treatment, nonetheless, there were no differences in sacroiliac joint score for erosion between untreated **Hyp** mice and **Hyp** mice with late treatment (Figures 3A,C and Supplementary Table 3).

Compared to **Hyp** mice in the early treatment group, **Hyp** mice treated late had a significant higher score at M3 (Figure 4D and Supplementary Tables 2, 3).

#### Effect of conventional treatment on bone markers

To confirm the micro-CT results and study the pathophysiological mechanism, we performed histological analyses of sacroiliac joints of the 3-month-old mice (Figures 4, 5).

Von Kossa and Masson’s Trichrome staining of the sacroiliac joint confirmed altered mineralization with accumulation of osteoid in untreated **Hyp** mice, compared to WT mice. The accumulation of osteoid was still evident in **Hyp** mice with early or late treatment (Figure 4). As expected, **Hyp** mice showed strong ALP staining (a marker of ALP activity), especially at the periphery of the osteoid. In contrast, in untreated **Hyp** mice, TRAP staining indicated that osteoclasts were present but concentrated in large “clusters” in the peripheral zone of sacroiliac joint. Both early and late start of conventional treatment in **Hyp** mice hardly modified ALP/TRAP activity (Figure 5).

We further studied the expression of sclerostin as a marker of differentiated osteocytes and bone turnover. Immunohistochemistry showed sclerostin expression in osteocytes of subchondral bone in WT mice (Supplementary Figure 2). Interestingly, untreated **Hyp** mice and **Hyp** mice treated late showed only faint sclerostin expression. In contrast, in the **Hyp** mice given early treatment, sclerostin expression was somewhat higher, and this finding is suggestive of improved regulation of bone turnover by terminally differentiated osteocytes.

#### Peripheral Enthesophytes (Calcaneus)

**Early start of conventional treatment**

Micro-CT images of hind paws showed similar features in 3-month-old **Hyp** mice of both treated and untreated groups (Supplementary Figure 3). Histological sections revealed mineralizing fibrochondrocytes expanding into both Achilles tendon and plantar fascia ligament insertions of calcaneal tuberosity in **Hyp** mice (Figure 6).

**Late start of conventional treatment**

The multiple calcaneal enthesophytes present in untreated **Hyp** mice were also seen in **Hyp** mice on conventional treatment started late (Figure 6).
**FIGURE 2** | Parameters of bone microarchitecture evaluated with micro-CT in WT mice, untreated Hyp mice, and Hyp mice on conventional treatment started early and late. (A) 3D-images of bone microarchitecture at baseline (W3), M2 and M3 in WT mice, untreated Hyp mice, and Hyp mice on conventional treatment started early, and at M2 and M3 in WT mice, untreated Hyp mice, and Hyp mice on conventional treatment started late. (B) BV/TV and TbN at baseline (W3), M2 and M3 in WT mice, untreated Hyp mice, and Hyp mice on conventional treatment started early. (C) BV/TV and TbN at baseline (M2) and M3 in WT mice, untreated Hyp mice, and Hyp mice on conventional treatment started late. (D) BV/TV and TbN at M2 and M3 in Hyp mice on conventional treatment started early compared to Hyp mice on conventional treatment started late. BV/TV, bone volume/total volume ratio; TbN, trabecular number. W3: 3 weeks; M2: 2 months; M3: 3 months; *p < 0.05; **p < 0.01; ***p < 0.001.

**Effect of conventional treatment on bone markers**
Alkaline phosphatase and TRAP staining in the area of the enthesophytes at the insertion of the Achilles tendon and plantar fascia ligament was similar in Hyp mice that received treatment, either early or late, and in both cases more than that observed in WT mice (Figure 6). That is, these alterations persisted in Hyp mice regardless of early conventional treatment. Neither early nor late conventional treatment seems able to prevent excessive bone mineralization at the Achilles tendon (enthesopathies) and restore ALP and TRAP activity (Figure 6).

In WT mice, sclerostin labeling was not observed in the Achilles tendon. In contrast, labeling was observed in both untreated and treated (early or late) Hyp mice at the insertion of the tendon, corresponding to the area where enthesopathies develop (Supplementary Figure 2).
DISCUSSION

The main focus of conventional treatment (phosphate supplements and active vitamin D) is growth restoration, and the impact of this type of treatment on XLH manifestations has been poorly investigated after the end of growth. In this context, we studied its effect on the main manifestations of XLH, in particular, skeletal features. Our study is the first to demonstrate that conventional treatment of XLH started early significantly improves bone microarchitecture and sacroiliac joint lesions but has little effect on enthesopathies assessed at the calcaneus. Empirically, it is assumed that current medical treatment of XLH should be started as early as possible to optimize final clinical outcomes in children (Linglart et al., 2014; Haffner et al., 2019). Nonetheless, this has yet to be proven in prospective studies. We have demonstrated that early conventional treatment has a significantly positive effect on impaired skeletal development. Micro-CT analysis showed that Hyp mice with early treatment showed a progressive increase in parameters of bone quantity (BV/TV, TbN) and structure (TbPf) resulting in a bone microarchitecture similar to WT mice. Interestingly, patients with XLH show compromised trabecular microarchitecture despite conventional treatment received since childhood (Cheung et al., 2013; Shanbhogue et al., 2015; Colares Neto et al., 2017). The major limitation of these studies is their retrospective design without taking into account age at the start or the duration of conventional treatment.

Importantly, our histological analyses showed a persistence of osteoid accumulation in treated Hyp mice by Von Kossa staining. Regardless of improvement of bone microarchitecture, neither early nor late treatment cured osteomalacia. However, restoration of sclerostin expression at the sacroiliac joint in Hyp mice in the early treatment group suggested that terminally differentiated osteocytes may regulate bone turn over, even if ALP and TRAP
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FIGURE 4 | Masson’s Trichrome and Von Kossa staining of sacroiliac joint of 3-month-old WT mice, untreated Hyp mice, and Hyp mice on conventional treatment started early or late. Masson’s Trichrome and Von Kossa staining showed a weaker mineralization (in dark) of the collagen matrix revealing an enlarged osteoid (red arrows) characteristic of features of osteomalacia bone in Hyp mice compared to WT. Despite the conventional treatment, Hyp mice treated early or late show an accumulation of osteoid (red arrows).

activities presented similar features in treated and untreated Hyp groups.

The effect of conventional treatment on skeletal manifestations depends on the bone involved. A significant reduction in the sacroiliac alterations was demonstrated in Hyp mice treated early, compared to Hyp mice treated late. Regarding the peripheral skeleton, conventional treatment, both early and late, has little effect on enthesopathies. Hyp mice in both treatment groups had persistent heel enthesophytes at the end of the study. The ALP expression in Achilles tendon in treated Hyp mice ultimately confirmed the expansion of mineralizing fibrochondrocytes in ligaments. Interestingly, we noticed sclerostin expression by fibrochondrocytes at the tendon of Hyp mice treated early. These results are concordant with Karaplis et al. (2012) findings and also the clinical study by Connor et al. (2015), which also demonstrated the limited effect of conventional treatment on enthesopathies in patients with XLH. From a pathophysiological point of view, hyperplastic fibrocartilaginous chondrocytes in the tendon are independent of improved mineralization of the bony part at the calcaneus. In this condition, newly mineralized bone is likely not strong enough to prevent the expansion of mineralizing fibrocartilaginous chondrocytes within the enthesis at the points of mechanical strain, a process that may be a compensatory mechanism to respond to biomechanical properties of poorly mineralized bone.

Nowadays, the mechanism of early osteoarthritis and enthesopathies in XLH is poorly understood. The degenerative osteoarthropathy principally depends on the accumulation of unmineralized immature bone (Liang et al., 2011; Wei and Bai, 2016). Additionally, high levels of FGF23 may promote the Wnt/β-catenin pathway in chondrocytes which activates the genes responsible for increased chondrocyte differentiation and osteoarthritis progression. Consequently, low expression of inhibitors of the Wnt/β-catenin pathway such as sclerostin is usually associated with the promotion of cartilage degradation (Meo Burt et al., 2018). Indeed, the results of our study confirm that accumulation of osteoid in the subchondral bone and lack of sclerostin expression are associated with the appearance and progression of numerous erosions at the sacroiliac joint in non-treated Hyp mice. On the contrary, sclerostin expression was restored in Hyp mice treated early, suggesting a link with the reduction of sacroiliac bone erosion in this group compared to Hyp mice with late treatment.

Early conventional treatment did not completely restore spine length in Hyp mice. Conventional treatment significantly increased the gain in length in Hyp mice treated early or late, though it did not have a significant impact on final length. In fact, we found no significant differences in length as assessed by micro-CT between non-treated and treated Hyp mice (early or late) at the end of the experiment. These results are in line with other studies performed in children with XLH. Specifically, healing active rickets promotes growth and after 2 years of successful treatment growth velocity is restored to its maximal potential in the majority of patients; however, 25–40% of patients with well-controlled XLH show linear growth failure despite optimized treatment (Linglart et al., 2014). It may be explained by several reasons. First, linear growth depends on bone mineralization and also on the cartilage growth plate...
FIGURE 5 | ALP and TRAP enzyme histochemistry of sacroiliac joint of 3-month-old WT mice, untreated Hyp mice, and Hyp mice on conventional treatment started early or late. The TRAP reaction stained for osteoclast cell activity and the osteogenic layer stained with ALP reaction indicate that both early and late start of conventional treatment in Hyp mice hardly changed ALP/TRAP activity. ALP: alkaline phosphatase; TRAP: tartrate-resistant alkaline phosphatase.

(Fuente et al., 2018). Partial growth restoration might result from a poor effect of the conventional treatment on growth plate maturation. Second, the conventional treatment, being a supplementary treatment, does not have influence on other players involved in XLH pathogenesis such as high levels of FGF23 (Carpenter et al., 2010; Zhukouskaya et al., 2020) or accumulation in the extracellular matrix of other proteins or peptides (osteopontin, ASARM peptides, etc.) (Salmon et al., 2013, 2014; Coyac et al., 2018).

The major strength of our study lies in its clinical implications. This is the first preclinical research demonstrating the beneficial effects of early conventional treatment in a well-designed prospective study, the findings underlining the importance of this treatment in the management of XLH. Our results were confirmed by several different methods, from clinical manifestations to radiological and histological analysis. Overall, the data provided highlight the usefulness of the Hyp murine model for preclinical studies on skeletal and osteoarticular lesions.

CONCLUSION

We have shown that conventional treatment given since an early stage improves bone microarchitecture and prevents joint erosions, though it does not have a notable effect on the formation of enthesophytes. Despite the improvement of bone microarchitecture, the persistence of osteomalacia
implies seeking new therapeutic strategies. Further studies are needed to compare these outcomes with the potential benefits of new therapies such as anti-FGF23 to improve the treatment of XLH.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

**ETHICS STATEMENT**

All experiments were performed with a protocol approved by the Animal Care Committee of the Université de Paris (project agreement 20-008, APAFiS #27827 N°202001171429974). Animals were maintained in accordance with the ethical protocol approved by the Animal Care Committee of French Veterinary Services (DPP Haut de Seine, France: agreement number D9204901).

**AUTHOR CONTRIBUTIONS**

AL, CC, and CB contributed to the design of the experiments. AC, VVZ, SH, JS, LS, CM-R, KB, AL, CC, and CB performed and analyzed the experiments. VVZ and CB wrote the manuscript with contributions from all authors. All authors reviewed and approved the final version of the manuscript.
**FUNDING**

This study was supported by the Agence Nationale de la Recherche (Grant 18-CE14-0018-01) and Fondation pour la Recherche Médicale (Grant DGE20111123012). AC was supported by the Société Française d’Endocrinologie et de Diabétologie Pédiatrique.

**ACKNOWLEDGMENTS**

We are grateful to Annie Llorens, Coralie Torrens, Brigitte Baroukh, and Sandy Ribes for their kind assistance with the histological techniques, and Thorsten Schinke for the Hyp mouse line. *In vivo* imaging was performed at the Life Imaging Facility of Paris Descartes University (Plateforme Imageries du Vivant), supported by France Life Imaging (Grant ANR-11-INBS-0006) and Infrastructures Biologie-Santé.

**REFERENCES**

Boussen, M. L., Boyd, S. K., Christiansen, B. A., Guldberg, R. E., Jepsen, K. I., and Muller, R. (2010). Guidelines for assessment of bone microstructure in rodents using micro-computed tomography. *J. Bone Miner. Res.* 25, 1468–1486. doi: 10.1002/jbmr.141

Carpenter, T. O., Insogna, K. L., Zhang, J. H., Ellis, B., Nieman, S., Simpson, C., et al. (2010). Circulating levels of soluble klotho and FGF23 in X-linked hypophosphatemia: circadian variance, effects of treatment, and relationship to parathyroid status. *J. Clin. Endocrinol. Metab.* 95, E352–E357. doi: 10.1210/jc.2010-0589

Che, H., Roux, C., Etcheto, A., Rothenbuhler, A., Kamenicky, P., Linglart, A., et al. (2016). Impaired quality of life in adults with X-linked hypophosphatemia and skeletal symptoms. *Eur. J. Endocrinol.* 174, 325–333. doi: 10.1530/eje-15-0661

Cheung, M., Roscher, P., Klaushofer, K., Veilleux, L. N., Roughley, P., Glorieux, F. H., et al. (2013). Cortical and trabecular bone density in X-linked hypophosphatemic rickets. *J. Clin. Endocrinol. Metab.* 98, E954–E961. doi: 10.1210/jc.2012-4133

Colares Neto, G. P., Pereira, R. M. R., Alvarenga, J. C., Takayama, L., Funari, M. F. A., and Martin, R. M. (2017). Evaluation of bone mineral density and microarchitectural parameters by DXA and HR-pQCT in 37 children and adults with X-linked hypophosphatemic rickets. *Osteoporosis Int.* 28, 1685–1692. doi: 10.1007/s00198-017-3949-8

Connor, J., Olear, E. A., Insogna, K. L., Katz, L., Baker, S., Kaur, R., et al. (2015). Conventional therapy in adults with X-linked hypophosphatemia: effects on entheseopathy and dental disease. *J. Clin. Endocrinol. Metab.* 100, 3625–3632. doi: 10.1210/jc.2015-2199

Coyac, B. R., Hoac, B., Chaefy, P., Falgayrac, G., Slimani, L., Rowe, P. S., et al. (2018). Defective mineralization in X-linked hypophosphatemia dental pulp cell cultures. *J. Dent. Res.* 97, 184–191. doi: 10.1177/0022034517728497

Faraji-Belle, C.-A., Cauzel, A., Salmon, B., Fogel, O., Zhukouskaya, V., Benoit, A., et al. (2020). Development of entheseopathies and joint structural damage in a murine model of X-linked hypophosphatemia. *Front. Cell Dev. Biol.* 8:854. doi: 10.3389/fcell.2020.00854

Fuente, R., Gil-Peña, H., Claramunt-Taberner, D., Hernandez-Frias, O., Fernandez-Iglesias, A., Hermida-Prado, F., et al. (2018). Marked alterations in the structure, dynamics and maturation of growth plate likely explain growth retardation and bone deformities of young Hyp mice. *Bone* 116, 187–195. doi: 10.1016/j.bone.2018.08.004

Haffner, D., Emma, F., Eastwood, D. M., Duplan, M. B., Bacchetta, J., Schnabel, D., et al. (2019). Clinical practice recommendations for the diagnosis and management of X-linked hypophosphatemia. *Nat. Rev. Nephrol.* 15, 435–455. doi: 10.1038/s41561-019-0152-5

Karaplis, A. C., Bai, X., Falet, J.-P., and Macica, C. M. (2012). Mineralizing entheseopathy is a common feature of renal phosphate-wasting disorders attributed to FGF23 and is exacerbated by standard therapy in hyp mice. *Endocrinology* 153, 5906–5917. doi: 10.1210/en.2012-1551

Kinoshita, Y., and Fukumoto, S. (2018). X-linked hypophosphatemia and FGF23-related hypophosphatemia diseases: prospect for new treatment. *Endocrine Rev.* 39, 274–291. doi: 10.1210/er.2017-00220

Liang, G., Vanhouten, J., and Macica, C. M. (2011). An atypical degenerative osteoarthropathy in Hyp mice is characterized by a loss in the mineralized zone of articular cartilage. *Calciified Tissue Int.* 89, 151–162. doi: 10.1007/s00223-011-9502-4

Linglart, A., Biosse-Duplan, M., Briot, K., Chaussain, C., Estelle, L., Guillaume-Citrom, S., et al. (2014). Therapeutic management of hypophosphatemic rickets from infancy to adulthood. *Endocrine Connect.* 3, R13–R30. doi: 10.1530/EC-13-0103

Meo Burt, P., Xiao, L., and Hurley, M. M. (2018). FGF23 regulates Wnt/beta-catenin signaling-mediated osteoarthritis in mice overexpressing high-molecular-weight FGF2. *Endocrinology* 159, 2386–2396. doi: 10.1210/en.2018-00184

Salmon, B., Bardet, C., Coyac, B. R., Baroukh, B., Naji, J., Rowe, P. S., et al. (2014). Abnormal osteosponin and matrix extracellular phosphoglycoprotein localization, and odontoblast differentiation, in X-linked hypophosphatemic teeth. *Connect. Tissue Res.* 55(Suppl. 1), 79–82. doi: 10.3109/03008207.2014.923864

Salmon, B., Bardet, C., Khaddam, M., Naji, J., Coyac, B. R., Baroukh, B., et al. (2013). MEPE-derived ASARM peptide inhibits odontogenic differentiation of dental pulp stem cells and impairs mineralization in tooth models of X-linked hypophosphatemia. *PLoS One* 8:e56749. doi: 10.1371/journal.pone.0056749

Shanbhogue, V. V., Hansen, S., Folkestad, L., Brixen, K., and Beck-Nielsen, S. S. (2015). Bone geometry, volumetric density, microarchitectute, and estimated bone strength assessed by HR-pQCT in adult patients with hypophosphatemic rickets. *J. Bone Miner. Res.* 30, 176–183. doi: 10.1002/jbmr.2310

Steele, A., Gonzalez, R., Garbalosa, J. C., Steigbigel, K., Grgurich, T., Parisi, E. J., et al. (2020). Osteoarthritis, osteophytes, and entheseophytes affect biomechanical function in adults with X-linked hypophosphatemia. *J. Clin. Endocrinol. Metab.* 105(Suppl. 1), 79–82. doi: 10.1210/jc.2019-00870

Wei, Y., and Bai, L. (2016). Recent advances in the understanding of molecular mechanisms of cartilage degeneration, synovitis and subchondral bone changes in osteoarthritis. *Connect. Tissue Res.* 57, 245–261. doi: 10.1080/03008207.2016.1177636

Zhukouskaya, V. V., Rothenbuhler, A., Colas, A., Di Somma, C., Kamenicky, P., Trabado, S., et al. (2020). Increased prevalence of overweight and obesity in children with X-linked hypophosphatemia. *Endocrine Connect.* 9, 144–153.

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcell.2020.591417/full#supplementary-material