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A distinctive patchy osteomalacia characterises Phospho1 deficient mice

Alan Boyde¹, Katherine A Staines², Behzad Javaheri³, Jose-Luis Millan⁴, Andrew A Pitsillides³, Colin Farquharson²

¹ Dental Physical Sciences, Barts and The London School of Medicine and Dentistry, Queen Mary University of London, London E1 4NS, UK, ² Roslin Institute and R(D)SVS, The University of Edinburgh, Easter Bush, Midlothian EH25 9RG, UK, ³Comparative Biomedical Sciences, Royal Veterinary College, Royal College Street, London NW1 0TU, UK, ⁴Sanford Children's Health Research Center, Sanford Burnham Prebys Medical Discovery Institute, La Jolla, CA 92037, USA

Corresponding author

Alan Boyde, Dental Physical Sciences, Barts and The London School of Medicine and Dentistry, Queen Mary University of London, London E1 4NS, UK

Email: a.boyde@qmul.ac.uk Tel: +44 20 7882 5984

Short running title: Patchy osteomalacia in the absence of Phospho1

Abstract

The phosphatase PHOSPHO1 is involved in the initiation of biomineralisation. Bones in Phospho1 KO mice show histological osteomalacia with frequent bowing of long bones and spontaneous fractures:
they contain less mineral, with smaller mineral crystals. However, the consequences of Phospho1 ablation on the microscale structure of bone are not yet fully elucidated. Tibias and femurs obtained from wild-type and Phospho1 null (KO) mice (25-32 week-old) were embedded in PMMA, cut and polished to produce near longitudinal sections. Block surfaces were studied using 20kV backscattered-electron (BSE) imaging, and again after iodine staining to reveal non-mineralised matrix and cellular components. For 3D characterisation, we used x-ray microtomography. Bones opened with carbide milling tools to expose endosteal surfaces were macerated using an alkaline bacterial pronase enzyme detergent, 5% hydrogen peroxide and 7% sodium hypochlorite solutions to produce 3D surfaces for study with 3D BSE scanning electron microscopy. Extensive regions of both compact cortical and trabecular bone matrix in Phospho1 KO mice contained no significant mineral and/or showed arrested mineralisation fronts, characterised by a failure in the fusion of the calcospherite-like, separately mineralising, individual micro-volumes within bone. Osteoclastic resorption of the uncalcified matrix in Phospho1 KO mice was attenuated compared with surrounding normally-mineralised bone. The extent and position of this aberrant biomineralisation varied considerably between animals, contralateral limbs and anatomical sites. The most frequent manifestation lay, however, in the nearly complete failure of mineralisation in the bone surrounding the numerous transverse blood vessel canals in the cortices.
In conclusion, SEM disclosed defective mineralising fronts and extensive patchy osteomalacia, which has previously not been recognised. These data further confirm the role of this phosphatase in physiological skeletal mineralisation.

**Key words:** osteomalacia, biomineralisation, PHOSPHO1, osteoid, backscattered-electron imaging

**Introduction**

Bone formation depends on an integration of events orchestrated by the capacity of osteoblasts to synthesise, secrete and mineralise bone’s extracellular matrix (ECM). This cascade of events which culminates in mineral deposition in collagen (biomineralisation), is critical to skeletal structure, integrity and function throughout life (Kawasaki et al., 2009).

PHOSPHO1, a member of the haloacid dehalogenase superfamily, is highly expressed in bone and contributes to bone mineralisation through its participation in the formation of the initial hydroxyapatite seed crystals (Stewart et al., 2003, 2006, Roberts et al., 2007, Houston et al., 1999, 2004). Treatment of developing chick embryos with lansoprazole, a small molecule that inhibits PHOSPHO1, completely inhibited mineralisation of all long bones (Macrae et al., 2010). Furthermore, bones from Phospho1 KO mice accumulate mineral poorly, leading to less stiff and more ductile bones in young juvenile mice (Huesa et al., 2011, Carriero et al., 2014, Rodriguez-
This more compliant bone is consistent with the relative protection of long bones from *Phospho1* KO mice against fracture during 3-point bending (Huesa et al., 2011). Also, the greater deformability is also consistent with the multiple skeletal abnormalities described in *Phospho1* KO mice which include osteomalacia, bowing of long bones and sporadic spontaneous fractures exhibiting a predominantly greenstick presentation (Yadav et al., 2011). Interestingly, this greater bone deformability appears to resolve in adult mice where the degree of stiffness in the *Phospho1* KO bones was corrected upon attainment of skeletal maturity (Javaheri et al., 2015). It is therefore likely that the optimal actions of PHOSPHO1 are most pronounced during rapid and timely ECM mineralisation in the early development of the skeleton, and that this contribution can be effectively mimicked by alternative mechanisms in later life (Millan, 2013, Yadav et al., 2016).

Studies by Yadav et al (2011) have examined whether overexpression of tissue non-specific alkaline phosphatase (TNAP) can effectively prevent the development of skeletal abnormalities in *Phospho1* KO mice. They found that TNAP overexpression offered no apparent protection against the deleterious bone phenotype that develops in *Phospho1* KO mice despite the correction of plasma pyrophosphate (PPi) levels - a potent inhibitor of ECM mineralisation (Meyer, 1984) - in these mice. This suggests that PHOSPHO1 and TNAP have non-redundant functional roles during skeletal mineralisation. This is emphasised by the deletion of a single allele of *Alpl* in the *Phospho1* null
background, which was found to aggravate the skeletal phenotype of Phospho1 KO mice whereas there was a complete absence of skeletal mineralisation in embryonic mice with a double ablation of Phospho1 and Alpl function (Yadav et al., 2011).

In spite of this growing awareness of the contribution made by PHOSPHO1 to biomineralisation, our knowledge is mainly limited to observations made at the whole bone level (macroscale) where valuable information on bone structure and biomechanical properties of Phospho1 KO mice have been reported. In contrast, the importance of PHOSPHO1’s contribution to bone mineralisation at the micron resolution scale has been less well addressed. The aim of this present study was therefore to perform a mainly scanning electron microscopic (SEM) based analysis of Phospho1 KO bones to further clarify the function of this intracellular/intravesicular phosphatase in biomineralisation.

Materials and methods

Animal model

Phospho1-R74X-null mutant (Phospho1 KO) mice were generated by N-ethyl-N-nitrosourea mutagenesis (ENU) as described previously (Yadav et al., 2011). Mice were housed up to 4 per cage in polypropylene cages with wood chip and paper bedding and provided standard mouse chow and water ad libitum throughout the study. Weaners up to 8 weeks of age were fed a
standard rodent breeding diet and thereafter a standard rodent maintenance
diet (Special Diet Services, South Witham, UK). All procedures complied
with the UK Animals (Scientific Procedures) Act 1986 and were reviewed and
approved by the ethics committee of The Roslin Institute, University of
Edinburgh.

**PMMA embedding**

Tibias and femurs from C57BL6 wild-type (WT) and Phospho1 KO male
mice (25 – 32 week-old) were fixed in formaldehyde for 48 hours, dehydrated
in ethanol; substituted with xylene; infiltrated in pure methyl methacrylate
monomer; then with monomer catalysed with α-azo-iso-butyronitrile at room
temperature until rubbery; then finally hardened at 37°C. The PMMA blocks
were cut and polished to produce near longitudinal sections.

**Backscattered Electron (BSE) SEM imaging (Fig 1A)**

Uncoated PMMA blocks were imaged using 20kV backscattered electron
(BSE) imaging at 50Pa pressure in a Zeiss EVO MA10 SEM (Zeiss,
Cambridge, UK). To determine whether there were obvious differences in the
degree of mineralisation in bone in WT and Phospho1 KO mice, matching
fields were imaged at identical sample-to-detector distances in quick sequence
under stable instrument operating conditions.

**BSE SEM imaging of triiodide stained PMMA block surfaces (Fig1B, 2A-D,
Fig 3A&B)**
To further characterise the histology of WT and Phospho1 KO bones, block surfaces were stained with ammonium tri-iodide to reveal non-mineralised matrix (such as osteoid) and cellular components before further BSE SEM (Boyde, 2012b).

*Maceration using Tergazyme™ and Hydrogen Peroxide (Fig 4A-E)*

To remove the cells and superficial non-mineralised collagen, we opened 70% ethanol fixed bones with carbide milling tools to expose the endosteal surfaces, which were then macerated with alkaline bacterial pronase (Tergazyme™, Alcanox, New York, NY: Boyde, 2012a) at 50°C and 5% H₂O₂ at room temperature. The samples were subsequently washed, dehydrated in ethanol and air-dried to produce uncoated surfaces for study with 3D 20kV BSE SEM, again at ~50Pa chamber pressure.

*X-ray microtomography (XMT) (Fig3C)*

For additional 3D characterisation, we used XMT (1172 x-ray microtomograph, Skyscan, Belgium). Bones from male WT and Phospho1 KO mice were fixed in 70% EtOH and stored until scanning, with X-ray tube operated at 50 kV and 200 μA1600 ms exposure time with a 0.5 mm aluminium filter, voxel size 5μm, 0.6° rotation angle, 2 frame averaging, of whole bones before embedding in PMMA. Trimmed PMMA blocks were scanned again after SEM imaging 3D, when rendering was performed using Drishti software (Australian National University, Canberra: Fig 3C).
Preparation of 10 micron ground section from SEM block face (Fig 3D-Ff)

In one instance, a ten micron thick ground section of a *Phospho1* KO block face studied by BSE SEM both before and after iodine staining and after XMT was prepared. The block face was glued to a 1mm thick glass slide with cyano-acrylate adhesive. A 1030nm laser operated in pulsed mode to give tissue ablation, was used to cut the block at ~15 µm into the tissue (‘TissueSurgeon’, Rowiak GmbH, Hannover, Germany), the surface then being finished by gentle polishing on 4000 grade silicon carbide polishing paper. The section was mounted using glycerine, studied by plane polarised light microscopy (Fig 3D), and further stained with MacNeal’s tetrachrome (Fig 3E: George T Gurr, London, UK), and acridine red (Fig 3F: Gurr) for conventional transmitted light microscopy.

Further SEM studies (Figs 5A-D & 6A,B)

After initial rounds of SEM study, many samples were re-prepared and re-examined by SEM. Some samples which had been macerated with Tergazyme and H₂O₂ were further treated with 5% available chlorine sodium hypochlorite (bleach) solutions, a more rigorous and vigorous method of removing non-mineralised collagenous matrices (Fig5A-D).

In some cases the bone was first remilled to provide better exposure of endosteal surfaces where these had been partially obscured in the initial preparation round. PMMA block surfaces were etched with an acid gel (Icon-
Etch, DMG, Hamburg, Germany) followed by removal of demineralised matrix with sodium hypochlorite solution to leave surface casts of PMMA embedded space (Figs 6A&B). Some samples which had been macerated as above were embedded in PMMA and block faces polished as above (Supplementary Figs 6C&D).

**Results**

BSE imaging of PMMA block faces revealed extensive regions of compact, cortical bone matrix in *Phospho1* KO mice which contained no significant mineral and showed arrested mineralisation fronts. This arrest - not observed in any of the WT control bones studied - was demonstrated by the failure of fusion of the separately mineralising micro-volumes (Fig. 1A).

We therefore stained the samples with iodine or tri-iodide to reveal non-mineralised matrix and cellular components, which highlighted the distribution of non-mineralised matrix (osteoid). Elliptical patches of mineralised matrix were regularly seen within the regions of non-mineralised matrix close to the otherwise mineralised matrix of *Phospho1* KO mice (Figs 1B; 2A-C; 3A,B; 6A).

Attempts to explore whether this patchy, arrested mineralisation occurred reproducibly at specific anatomical locations, demonstrated instead that the extent and position varied considerably between animals, contralateral limbs and anatomical sites. There was also great variation in the size of the non-
mineralised matrix areas, which ranged from zones approximately 50µm across that were centred around the blood vessel canals in the cortex, to the largest patches which extended to the millimetre scale (Figs 1A,B; 2A-C; 3A-F). The vascular channel associated deficiency appears to be the most frequent manifestation of defective biomineralisation in the Phospho1 KO mice (Fig 2A,B). Neither was ever observed in WT bones (Fig 2D).

In order to more precisely define the biomineralisation deficit, we selected a single location from BSE SEM imaging of iodine stained PMMA block surfaces that we could readily identify and thus examine at a range of scales, using several methodologies (Fig. 3). By XMT, we were unable to discriminate PMMA as such from PMMA embedded uncalcified bone and cartilage, or tendon and ligament, or bone marrow space. The non-mineralised matrix regions in bone therefore appeared as ‘empty’ space (Fig 3C, see also supplementary Fig7 and video submitted for supplementary documentation), but nevertheless allowed spatial equivalence to be registered in the images of the entire bone segment.

Polarised light examination of a ground section of the bone block-face prepared by dual photon laser ablation and at the boundary between non-mineralised cortex and mineralised bone in these Phospho1 KO mice revealed clear continuity of lamellae from mineralised to non-mineralised regions (Fig 3D).
Tetrachrome (blue-green in Fig 3E) or acridine red (pale purple, Fig 3F) staining of this patch of non-mineralised tissue in *Phospho1* KO mice was positive and resembled the staining colour and intensity observed in non-mineralised regions of osteoid found as expected elsewhere at forming bone surfaces. It is noteworthy, that osteocytes were present in the abnormal osteoid patches as seen both by iodine stained SEM of block faces and light microscopy (Fig 3B,E,F).

To visualise the calcified bone proper, we removed both cells and non-mineralised collagen with Tergazyme and H$_2$O$_2$ (Fig. 4). The mineralised matrices of the inorganic tissues were analysed following such ‘maceration’ by 3D BSE SEM morphological imaging (Boyde, 2012a). As some non-mineralised matrix osteoid remained after Tergazyme and H$_2$O$_2$ cleaning, we treated some samples additionally with sodium hypochlorite bleach (Fig. 5). This removed non-mineralised matrix remnants leaving a cleaner mineral front, making it easier to survey periosteal bone surfaces where fibrous periosteum, tendon or ligament residues are more difficult to remove.

Bones from *Phospho1* KO mice after Tergazyme and H$_2$O$_2$ cleaning exhibited extensive areas of osteoid and abnormal, arrested or defective mineralisation fronts on both the endosteal (Figs 4A-D) and periosteal surfaces of the cortex and on trabeculae (Suppl. Figs. 4G & H). The WT control bones revealed endosteal surface with resorbed patches and fully mineralised (smooth) regions (Fig. 4E, Suppl. Fig 4F).
The arrested or impeded collagen-fibre-bundle centred mineralisation in the *Phospho1* KO bones is indicative of osteomalacia. The observed hints of attenuation of osteoclastic resorption at the edges of uncalcified matrix patches in *Phospho1* KO bones (Fig 4D, 5A) is in line with the widely accepted understanding that osteoclasts hunt for mineral.

All macerated and bleached samples revealed very obvious areas of abnormal mineralisation fronts on trabecular bone which can be clearly demarcated from regions of osteoclastic resorption (Fig 5A, see also Supplementary Fig 6F&G). This clear demarcation was also observed on endosteal surfaces (Fig 5B) reinforcing the observation that aberrant mineralisation in Phospho1 KO mice attenuates resorption compared with surrounding normally-mineralised bone (Fig 5A&B). This treatment also accentuated the failure of mineralisation in the bone, surrounding blood vessel canals in the cortices, where separately mineralised micro-volumes were very obvious. The individual, calcospherite-like micro-volumes could be seen to be multiply-penetrated by osteocyte canaliculi (Fig 5C&D; see also Supplementary Fig 6C).

Abnormal mineralising fronts were encountered much less frequently at periosteal surfaces, in keeping with the fact that it is the endosteal surfaces in both distal femur and proximal tibia that are largely formative, whilst periosteal surfaces show a higher incidence of resorbing or resting regions (Enlow, 1962).
Finally, we have used PMMA-embedded material that had subsequently been etched with acid and bleach, to expose the non-mineralised regions ‘proud’ as a surface elevations and thus reveal the extent of the non-mineralised patches of bone in *Phospho1* KO mice in 3D. SEM of these etched PMMA blocks confirmed the extension of this non-mineralised bone into regions within the cortical bone of these *Phospho1KO* mice and also provide scope to visualise the osteocyte content of both the non-mineralised patch of bone and the normally mineralised neighbouring cortices (Fig 6A,B).

**Discussion**

Osteomalacia is often characterised by broad bands of osteoid with partial or complete failure of calcification; a phenotype that has been observed previously in *Phospho1* KO mice where the osteoid volume fraction is significantly increased (Yadav et al., 2011, 2014, 2016). The new data presented here suggests that the osteomalacia present in *Phospho1* deficiency may be locally more severe than previously reported and yet be clearly delineated with an abrupt transition. Furthermore, previous studies of the endosteal surface of many genetically modified mice with defective bone mineralisation phenotypes had, as yet, never revealed the distinctive type of defective mineralisation observed in the *Phospho1* KO mice. This suggests to us that the patchy osteomalacia noted in the *Phospho1* KO mice is not a general skeletal hallmark of hypomineralisation but is, as yet, rather a specific consequence of *Phospho1* deficiency.
The present observations were unexpected and it is difficult to fully reconcile the presence of the stochastically-distributed, relatively large patches of non-mineralised matrix with our present understanding of how PHOSPHO1 initiates the nucleation of bone mineral (Millan, 2013). Since the bone phenotype of Phospho1 KO mice was first described, efforts have been focussed on elucidating the mechanisms by which PHOSPHO1 contributes to skeletal mineralisation. Systemic levels of calcium and phosphate are normal in Phospho1 KO mice and therefore the defect in bone mineralisation is likely to be at the cellular level (Yadav et al., 2011). A possible mechanism may involve the potent calcification inhibitors, PPI, and osteopontin (OPN, encoded by the Spp1 gene) whose plasma and bone expression levels are both elevated in Phospho1 KO mice (Meyer 1984, Speer et al. 2002, Steitz et al. 2002, Yadav et al, 2011, 2014). However, reducing the PPI levels in Phospho1 KO mice by cross breeding them to a transgenic strain expressing human TNAP under control of the ApoE promoter, did not improve the skeletal phenotype of Phospho1 KO mice (Yadav et al. 2011). In contrast, the development of the skeletal phenotype of Phospho1 KO mice is prevented by the ablation of Spp1 function and establishes a primary role for OPN, rather than PPI, in the aetiology of the osteomalcia observed in the Phospho1 KO mice (Yadav et al. 2014).

Further efforts have centred on an early observation that PHOSPHO1 is present and active within matrix vesicles (MVs) where it may have a role in
scavenging inorganic phosphate from MV membrane phospholipids (phosphocholine and phosphoethanolamine) to favour intra-vesicular hydroxyapatite deposition (Roberts et al., 2004, Stewart et al., 2006). Atomic force microscopy and Raman spectroscopy analyses have shown that chondrocyte-derived MVs from Phospho1 KO mice have a decreased ability to initiate mineralisation (Yadav et al., 2016). These data extend our previous observations on mice KO in both Phospho1 and AlPl. Although embryonic lethal, the embryos show a complete absence of skeletal mineralisation and MVs devoid of mineral (Yadav et al., 2011).

Immature woven bone, like peripheral or mantle dentine, normally contains large numbers of MVs and the mineralisation pattern shows large numbers of small, separate microcalcospherites (Boyde, 1980a,b; 2012a). The slowly formed lamellar bone in large bones in adult large mammals (including man) contains few, if any, MVs: the mineralisation process occurs instead by spreading through the collagen fibre bundles of the bone matrix giving rise to the very characteristic collagen-bundled centred mineralisation front pattern seen in anorganic (deproteinised) preparations for SEM (Boyde, 1980a,b, 2012a). On the other hand, the rapidly formed lamellar bone in small rodents shows a much larger number of separate centres, and it is much more difficult to visualise the collagen layout in the matrix in the mineralising ‘front’. This would suggest that there are indeed large numbers of MVs. Then it becomes clear that MVs devoid of PHOSPHO1 will result in the observed
mineralisation pattern demonstrated here, but why the distribution of this osteomalacia is patchy is unclear. One possible explanation is that PHOSPHO1 plays a much more subtle and pivotal role at the inception phase of mineralisation than previously considered. We propose that the mineralisation deficiency conferred by PHOSPHO1 deletion is initially insurmountable, mostly at certain locations, where rapid rates of mineralisation are otherwise unconstrained, and are thus centred predominantly at endosteal and innermost perivascular surfaces. Long-term recovery from such patchy osteomalacia in the more mature adult skeleton would therefore imply that this PHOSPHO1 function can eventually be overcome by potentially diverse, slower-acting mineralisation mechanisms.

Alternative proposed mechanisms for mineral nucleation include the precipitation and propagation of mineral crystals in the gap zone of collagen fibrils (Glimcher, 1987, Landis et al., 1993, Landis and Jacquet, 2013, Nudelman et al., 2010). Alternatively, electron microscopy and X-ray diffraction studies on human cortical bone, have led others to propose that the majority of mineral is present outside of collagen fibrils and in the interfibrillar compartment, in the form of elongated mineral plate structures (McNally et al., 2012, McNally et al., 2013, Schwarcz et al., 2014). The results of ion beam thinning to prepare hard tissue sections for TEM have been interpreted as showing that mineral enwraps the collagen microfibril and is therefore within the fibril but topologically outside the (tropo)collagen
molecules, the mineral phase being present as extremely long features unrelated to the gap zone (Boyde, 1974, Boyde and Pawley, 1976). This further explains why anorganic bone samples, such as those prepared by prolonged hypochlorite treatment, do not fall apart to make a suspension of fine crystals. Instead they permit us to study the mineralisation phenomena in 3D objects, as in several of the figures in this paper.

A novel concept that mineral may enter osteoid collagen as ready-packaged globules of disordered calcium phosphate (Mahamid et al., 2011; Kerschnitzki et al., 2016a,b) - perhaps with carbonate enrichment at initial mineral cluster sites (Nitiputri et al, 2016) - does not contribute to explaining the patchy nature of the histological osteomalacia which is the focus of the present report.

PHOSPHO1 deficiency has been suggested to significantly increase osteocyte lacunae density and vascular content of cortical bone (Javaheri et al., 2015). These conclusions could, however, be explained by the present findings using higher resolution imaging as follows: the vascular canals are very frequently contained within the more numerous and smaller unmineralised matrix regions we reveal here. We show that osteoid cannot be distinguished from canal space by XMT, and the canals will therefore appear to be larger and to occupy a larger volume fraction. If there were, in addition, a thin seam of unmineralised matrix around some of the osteocyte lacunae – as is known to be the case in several instances of osteomalacia (Boyde, 1980a,b) - then the
lacunae will be more effectively detected by XMT - the more so the better the resolution and Javaheri et al. (2015) employed 0.6µm - and the count will increase, as well as the volume fraction.

The 3D extent of these non-mineralised regions can be seen in XMT image sequences (Suppl. Figure 7 and video). XMT resolution is inadequate in discriminating focal mineralisation defects from vascular/ lacunar-canalicular changes, but XMT has the advantage of maintaining the sample whole. Co-registration of images from both modalities discloses very effective co-localisation of the areas of bone matrix devoid of mineral and highlight the relatively random targeting of osteomalacia in the cortical bone. The patchy dissemination of the most severe type of lesion is a reminiscent of the mosaic distribution of lesions seen in fibrous dysplasia (Saggio et al., 2014). The present results with those shown earlier also disclose that osteocytogenesis seems to continue relatively unabated in the bones of Phospho1 KO mice, including in the totally unmineralised osteoid regions, yet some cell culture and in vivo studies have previously suggested that ECM mineralisation is an essential regulator of osteocytogenesis (Irie et al., 2008, Prideaux et al., 2012).

In conclusion, this study revealed the consequences of Phospho1 deficiency on the microscale structure of mineralised bone. SEM disclosed defective mineralising fronts and extensive histological osteomalacia which was most frequent in bone surrounding the numerous transverse blood vessel canals in the cortices. This extensive - but patchy or spotty - histological osteomalacia
found in Phospho1 KO lower limb bones has previously not been recognised and these data further confirm the necessity of this phosphatase for physiological mineralisation and the establishment of proper skeletal structure.

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**References**


Captions to illustrations.

Figure 1, 2 parts, Column width. Before and after ammonium triiodide staining.
Figure 1A. BSE of tibial cortex from Phospho1 KO mouse showing one very large and several smaller patches of non-mineralised bone matrix, i.e., osteoid. Section plane is a tangent to the external surface. All SEM images were recorded from uncoated samples, Zeiss EVO MA10 SEM, 20kV, with 50Pa chamber pressure to prevent electrostatic charging.

Figure 1B. Part of same region as Fig 1a after ammonium triiodide staining to reveal non-mineralised matrix and cellular components. Arrows show patches of mineralised matrix within the background of non-mineralised matrix osteoid and along the arrested mineralising front.

Figure 2, 4 parts, page width, i.e., a ‘Plate’. All parts ammonium triiodide stained to reveal soft tissue components.

Figure 2A. KO. Proximal tibia showing many patches of mineralised matrix within areas of non-mineralised matrix osteoid (white arrows).

Figure 2B. KO. Tibial diaphysis. White arrows show osteoid patches at endosteal, marrow surface. Black arrows show variation in the size of osteoid areas (down to zones 20 - 50µm across) centred on the blood vessel canals in the cortex.

Figure 2C. KO. Knee. Dense fibrous connective tissue of periosteum at white arrows, not to be confused with osteoid areas in cortex at black arrows.

Figure 2D. WT control knee, same scale.

Figure 3, 6 parts, page width, i.e., a ‘Plate’.
Figure 3A-F. Correlation of block face BSE SEM, XMT and PLM and LM. Asterix shows same spot in each part.

**Figure 3A.** KO tibia. BSE SEM of ammonium tri-iodide stained block face to reveal non-mineralised matrix osteoid and cellular components. Black arrow points to large osteoid patch. (After SEM studies we cut the blocks to make smaller samples for high resolution XMT).

**Figure 3B.** Higher magnification view of large and small osteoid patches. Blood vessel canals are arrowed (black in bone and white in osteoid). Asterix shows spot similarly marked in Fig 3c-f.

**Figure 3C.** Drishti reconstruction of XMT data. The large osteoid patch seen in Figs 3a & B show as void (white horizontal arrow) in the rendered XMT Image. This is in same position as black arrow in Fig 3A..

**Figure 3D.** ~10μm section of face of same block prepared by laser ablation (Rowiak ‘TissueSurgeon’) viewed between crossed polars, showing continuity between lamellae in mineralised bone and osteoid. Field width = 141μm.

**Figure 3E.** Same section, stained with MacNeal’s tetrachrome stain giving green-blue tint to osteoid. Field width = 900μm.

**Figure 3F.** Same field stained with acridine red. Field width = 900μm.

**Figure 4, 5 parts, page width, ie, a ‘Plate’**.
**Figure 4A.** Central part of shaft endosteum in a Tergazyme macerated KO femur. The dark patches are all abnormal osteoid (This scene is shown as a 3D anaglyph (red cyan) image in Supplementary Fig 4A).

**Figure 4B.** Endosteum in a Tergazyme macerated KO tibia. Dark osteoid surrounds blood vessel canals (white arrows) within an area showing a mineralising front with unfused mineralising centres. This is surrounded by resorption (labels Res).

**Figure 4C.** Endosteum in a Tergazyme macerated KO femur. Dark residual osteoid surrounds blood vessel canals (white arrows). Individual canaliculi show as small black dots (too small to label) within the unfused mineralising centres show. The fully mineralised back wall of a forming osteocyte lacuna lies near dead centre of image at point of black arrow.

**Figure 4D.** Endosteum in a Tergazyme and H$_2$O$_2$ macerated KO femur. The additional treatment removes residual osteoid from the endosteal surface and lining the blood vessel canals (black arrows). An area with abnormal mineralising front (outlined with white asterixes ‘*’) is surrounded by resorption (Res).

**Figure 4E.** Endosteum in a Tergazyme and hydrogen peroxide macerated WT tibia at same scale, showing mostly a fully mineralised, smooth surface with some shallow resorbed patches (Res: one area outlined with white asterixes ‘*’). The openings of the blood vessel canals (black arrows) show a fully
mineralised surface. Surface osteocyte lacunae above labels ‘sol’. Some osteocyte lacunae are so superficial that their canaliculi lie in the plane of the fully mineralised (smooth) surface, showing as grey linear features in areas opposite points of white arrows. See also Supplementary Fig 4F.

**Figure 5, 4 parts, page width, ie, a ‘Plate’**.

**Figure 5. KO endosteal surfaces after further cleaning with NaOCl bleach to remove all organic matrix, importantly here, any osteoid.**

**Figure 5A.** KO tibia. Osteoclastic resorption (Res) has not proceeded into the abnormal mineralisation front area on an endosteal trabeculum.

**Figure 5B.** KO tibia This is an area in which endosteal osteoclastic resorption (Res) has extended into bone with defective mineralisation surrounding blood vessel canal spaces (all the large dark features). There would have been osteoid everywhere that we see unfused mineralised patches.

**Figure 5C.** KO tibia, cut edge at top. This is an area in which endosteal osteoclastic resorption (Res) has extended into an endosteal osteoid patch (now removed by NaOCl) analogous to the regions arrowed white in Fig 2B. It again shows defective mineralisation surrounding blood vessel canal spaces (the large dark features).

**Figure 5D.** KO femur. Unmineralised osteoid would certainly have been present over the right side of the field which shows unfused mineralised regions in which canaliculi can be seen as small dark features. The left side
had been subject to osteoclastic resorption (Res). Note that the mineralisation of this region – at the border - appears to be complete, i.e., the bone was uniformly mineralised where mineralised.

**Figure 6, 2 parts, column width.**

**Figure 6. Same field before and after etching**

**Figure 6A.** KO tibia, PMMA embedded ammonium tri-iodide stained to reveal soft tissue components, showing an osteoid patch at centre. Surface normal to electron beam. (All other SEM images in this MS were also recorded at normal electron beam incidence).

**Figure 6B.** The same region after ‘DMG Icon-Etch’ HCl gel etching for 1 min, followed by 5% NaOCl for 10 mins and ammonium tri-iodide staining. PMMA within the osteoid patch (os) is left proud after the combined etch treatment. Osteocyte lacunae also appear as ‘casts’ above the etched surface. Surface tilted at 33 degrees.

**Supplementary images and information**

**Suppl Figure 4A,** anaglyph version for viewing with red (left eye) and cyan (right eye) spectacles on a computer/monitor screen. Central part of shaft endosteum in a Tergazyme macerated KO femur. The dark patches are all abnormal osteoid with unfused mineralising centres.

**Suppl Figure 4F.** Tergazyme and hydrogen peroxide macerated WT femur showing endosteal surface with resorbed patches (Res) and fully mineralised
(smooth) regions, one such outlined with white asterixes ‘*’. None of the changes found in KOs were seen in WTs. See also Figs 4D (KO) and Fig 4E (WT).

**Suppl Figure 4G & H show same field,** Tergazyme macerated KO femur, endosteum. An abnormal mineralising front indicating arrested or impeded collagen-fibre-bundle centred [NB NOT matrix vesicle centred!] is seen both at endosteal surfaces of the cortex and on trabeculae: this is indicative of osteomalacias in general. Focus in **Suppl Fig 4G** is on trabeculum in foreground and in **Suppl Fig 4H** is on endosteum in background.

**Post-maceration embedding.**

PMMA blocks of samples which had been previously macerated provided evidence that the maceration procedures had cleared out the cell contents of marrow spaces and blood vessel canals, but had left the abnormal osteoid in place.

**Suppl Figure 6C.** A Tergazyme and H$_2$O$_2$ macerated KO tibia was embedded in PMMA, the block surface polished and stained with ammonium triiodide solution. Methacrylate has filled the blood vessel canal spaces, including those within the abnormal osteoid patches (arrows), This proves that the maceration procedure did successfully removed cell remnants from within the blood vessel canal spaces.
**Suppl Figure 6D.** Tergazyme and H₂O₂ macerated KO tibia embedded in PMMA in the fused growth plate region, showing unfused calcospherites within the calcified cartilage (Cs), which is normal, and the unfused mineralised regions within the osteoid (os), which is an abnormal phenomenon (osteomalacia).

**Suppl Figure 7.** Sequence of XMT slices, every tenth slice = every 50µm horizontally, every 250µm vertically, showing distribution of osteoid patched within cortex from KO femur. Original slice numbers outside field at edges.

**Phospho1KO_XMT_movie.ppt or pptx or GIF**

Viewing XMT movies showing a series of cross sections is the best way to show the remarkable 3D extent of the totally non-mineralised regions.

Same data set as Suppl Fig 7 but using every 5µm section in the range.
Figure 2
Figure 5
Figure 6

A

marrow

20 μm

B

marrow

20 μm

os