Skeletogenesis occurs through endochondral and intramembranous ossification. During intramembranous ossification, mesenchymal stem cells (MSC) directly differentiate into osteoblasts. In endochondral ossification, MSCs first differentiate to chondrocytes forming the cartilage, which is subsequently replaced by bone. Invading vasculature brings bone forming osteoblasts to endochondral bone, where they replace the cartilage by bone1–3. Osteoclasts are bone resorbing cells and together with osteoblasts they are responsible for bone remodeling4. Bone vasculature brings bone forming osteoblasts to endochondral bone, where they replace the cartilage by bone1–3. Osteoclasts are bone resorbing cells and together with osteoblasts they are responsible for bone remodeling4. Bone forming osteoblasts to endochondral bone, where they replace the cartilage by bone1–3. Osteoclasts are bone resorbing cells and together with osteoblasts they are responsible for bone remodeling4.

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Cilia-related protein SPEF2 regulates osteoblast differentiation

Mari S. Lehti1,2, Henna Henriksson2, Petri Rummukainen1,2, Fan Wang1, Liina Uusitalo-Kylmälä2, Riku Kiviranta2,3, Terhi J. Heino2, Noora Kotaja2 & Anu Sironen1

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In the targeted allele, mice in the testis and trachea and at low level in the lung, pituitary gland and kidney (Supplemental Fig. S2A). In the targeted allele, Dsred was introduced after Spef2 exons 1–2 in order to produce a SpeF2(exon1–2)/DsRed fusion protein. However, we were unable to detect the DsRed protein expression in KO tissues (data not shown).

Although the exon 3 is, as a rule, not included in the SpeF2(exon1–2)/Dsred hybrid transcript due to the transcription termination site after the Dsred sequence, we were able to detect a low level of the longer SpeF2/Dsred hybrid transcript in the trachea with exons 1–3 amplifying primers when we used a high number of cycles and a long elongation time in RT-PCR (Supplemental Fig. S2B). The sequencing of this PCR product revealed that the Dsred sequence was in fact not in frame with the Spef2 coding sequence and an additional stop codon was introduced after exon 2 of Spef2 (Supplemental Fig. S2B). Therefore, the DsRed protein expression could not be used to monitor the tissue distribution of Spef2 expression.

Spef2 KO mice suffered from severe hydrocephalus (Supplemental Fig. S3A) and mice survived a maximum of five weeks. The hydrocephalus was present in mice with 129 and B6 background indicating that the bgh mutation in exon 28. Bgh mice have similar spermatogenetic phenotype as the ISTS pigs, which is most likely caused by the nonsense mutation in exon 28. In addition, the bgh mice show primary ciliary dyskinesia (PCD)-like symptoms including sinusitis and hydrocephalus29, which are most likely caused by the missense mutation in exon 3 that affects several Spef2 isoforms. SPEF2 has been shown to interact with IFT-related protein IFT20, suggesting the involvement of SPEF2 in IFT30. In this study, we generated a mouse model with a stop codon located after exon 2 of Spef2 gene to further investigate the role of SPEF2 in ciliated tissues in mice.

Results

Disruption of Spef2 gene causes hydrocephalus and growth retardation. For identification of the role of SPEF2 in mice, we have generated a Spef2 KO mouse model. The targeting construct was designed to produce two mouse lines; conventional full KO model by introducing a DsRed reporter and transcription termination sequence after the Spef2 exon 2 and a conditional KO model by introducing loxp sites to surround exons 3–5 (Supplemental Fig. S1A). To generate the conditional male germ cell-specific Spef2 KO mouse line, the Dsred construct was removed using FLP-FRT recombination, and subsequently, the mice with floxed Spef2 gene were crossed with transgenic mice expressing Cre under the Neurogenin3 (Ngn3) promoter41. Here we report the results from the conventional KO mouse model, in which the Spef2 gene was inactivated in all tissues examined. The position of the introduced construct in relation to known Spef2 transcript variants and other reported Spef2 mutants is shown in Supplemental Fig. S1B. The genotype of pups was confirmed by PCR using primers flanking the LoxP site after exon 5 to amplify 478 bp product from mutated allele and 379 bp product from wild type (WT) allele (Supplemental Fig. S1A).

In control mice, the 5′ end of the Spef2 messenger RNA (mRNA) was expressed in the testis, epididymis, brain, trachea, pituitary gland, lung, kidney and eye as demonstrated by RT-PCR using primers amplifying exons 1–3 (Supplemental Fig. S2A). These primers failed to amplify the WT product in any of the Spef2 KO tissues, therefore confirming the successful abolishment of Spef2 gene expression (Supplemental Fig. S2A). The expression of the 3′ end of Spef2 (exons 37–43) was also shown to be absent in KO tissues, while the expression was detected in WT mice in the testis and trachea and at low level in the lung, pituitary gland and kidney (Supplemental Fig. S2A). In the targeted allele, Dsred was introduced after Spef2 exons 1–2 in order to produce a SpeF2(exon1–2)/DsRed fusion protein. However, we were unable to detect the DsRed protein expression in KO tissues (data not shown).

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Spef2 KO mice suffered from severe hydrocephalus (Supplemental Fig. S3A) and mice survived a maximum of five weeks. The hydrocephalus was present in mice with 129 and B6 background indicating that Spef2 is a causative gene for PCD46,47. The severe hydrocephalus developed between P15 and P30 probably causing the death of KO tissues, therefore confirming the successful abolishment of KO mouse line, the exon 2 and a conditional KO model by introducing loxp sites to surround the role of SPEF2 in mice, we have generated a Spef2 KO mouse model. The targeting construct was designed to

Ngn3 were crossed with transgenic mice expressing Cre under the Neurogenin3 (Ngn3) promoter (Supplemental Fig. S1A).

The LoxP site after exon 5 to amplify 478 bp product from mutated allele and 379 bp product from wild type (WT) allele (Supplemental Fig. S1A).

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SPEF2 has been reported to play an essential role during spermatogenesis48,49. Because Spef2 KO mice died before reaching sexual maturity, we were unable to study the progress of spermatogenesis and fertility in Spef2 KO mice. The analysis of Spef2 heterozygous (HEZ) mice revealed that the haploinsufficiency of Spef2 gene did not affect male fertility. Spef2 HEZ males produced average litter sizes (average number pups/litter 9.5 ± 2.13) and the distribution of genotypes of pups followed Mendelian ratio (Supplemental Fig. S3F). We did not find major defects in the overall organization of the seminiferous epithelium and spermatogenesis (Supplemental Fig. S3G). Furthermore, the motility of Spef2 HEZ spermatozoa was normal and equal numbers of progressive motile (PR), non-progressive motile (NP) and immotile (IM) spermatozoa were counted from both WT and Spef2 HEZ sperm samples (Supplemental Fig. S3H). Furthermore, the HEZ mice appeared viable and no phenotypic changes were observed compared to WT. Thus, a single copy of the Spef2 gene appears to be sufficient for the function of the gene.

Spef2 is expressed in the bone and cartilage. The size difference between WT and Spef2 KO mice prompted us to further investigate the possible effects of Spef2 depletion on bone formation and structure. To study the Spef2 gene expression in skeletal tissues, we isolated trabecular bone (P31) and cartilage (P3, P15) from proximal tibia. Spef2 was shown to be expressed in these tissues, and the expression was markedly decreased in both the bone and cartilage of Spef2 KO mice compared to WT (Fig. 1). To confirm Spef2 expression in osteoblasts and chondrocytes, we isolated and cultured osteoblasts and chondrocytes from the WT mouse calvaria and ribcages, respectively. Spef2 gene expression increased during osteoblast differentiation (Fig. 1B) and expression was also detected in the cultured chondrocytes (Fig. 1B).
Disruption of Spef2 affects the skeletal growth, structure and strength. We analyzed macroscopic organization and composition of the cartilage and bone tissue between WT and Spef2 KO mice using whole-mount staining. Alcian blue was used to stain the cartilage and Alizarin red the bone tissue. Staining appeared similar in WT and Spef2 KO mice at P31 and no differences were observed in ossification between WT and Spef2 KO, although the size difference of the animals was obvious (Fig. 2A). It has to be noted that the staining was performed for only one KO and WT animal. To investigate the bone tissue content in more detail, X-ray images were obtained from hind limbs and spines at P15 and P31 (Fig. 2B,C). X-ray images showed a reduced mineral content in Spef2 KO bone at both vertebral column and in long bones at both timepoints (P15 and P31) compared to WT. Bone strength was further evaluated using a 3-point bending test, where mechanical load was subjected to the WT and Spef2 KO femora midshaft at P31 (Fig. 2D). Spef2 KO bones were weaker and broke down with significantly lower maximal load than the WT femurs (Fig. 2D). To investigate whether Spef2 has a role in long bone growth we measured tibia and femora lengths at P15 and P31 and demonstrated that in Spef2 KO mice, both bones were significantly shorter at P15 with a (non-significant) growth delay also at P31 (Fig. 2E,F). We also measured the skull bone thickness at P15 and P31 and observed significantly thinner bones at P15 and P31 in Spef2 KO mice (Fig. 2G). However, since the hydrocephalus was more severe at P31, it may explain the more profound difference between WT and Spef2 KO at P31 compared to P15.

Disruption of Spef2 impairs the cortical and trabecular bone volume and structure. Micro-computed tomography (µCT) was used for microstructural analysis of trabecular and cortical bone in Th10 and L2 vertebrae (Fig. 3A). Trabecular bone volume density (BV/TV) was decreased at P31 in both Th10 and L2 vertebrae (Table 1). Trabecular number (Nb.Th, Fig. 3B) and thickness (Tb.Th, Fig. 3C) were significantly decreased and open porosity (Po(op), Fig. 3D) was increased in Spef2 KO mice vertebrae at P31. Cortical bone volume (BV, Fig. 3E), surface (BS, Fig. 3F) and connectivity (conn, Fig. 3G) appeared reduced in Spef2 KO mice in both studied vertebrae and timepoints, suggesting more loose cortical bone in the spine.

Distal femur cortical and trabecular bone parameters were also analyzed using µCT (Fig. 4A). BV/TV was significantly decreased at P15 in distal femur trabecular bone (Table 2). In addition, Tb.N (Fig. 4B) and Tb.Th. (Fig. 4C) were also significantly decreased at P15. Open porosity was increased and difference was significant at P15 (Fig. 4D). Parameters for distal femur cortical bone supported the data acquired from the vertebrae: BV (Fig. 4E), BS (Fig. 4F) and connectivity (Fig. 4G) were all decreased in Spef2 KO mice.

Osteoblast differentiation is impaired in Spef2 KO mouse. To investigate the effects of Spef2 depletion on osteoblast differentiation we first analyzed several osteoblast markers of the proximal tibia trabecular bone: the expression of Alp, Ocn, collagen 1 (Col1), Runx2 and Oss was studied by RT-qPCR. Although Alp expression appeared unchanged between WT and Spef2 KO bone tissue, Runx2, Ocn and Col1 were reduced in Spef2 KO bone indicating possible defects in osteoblast activity (Fig. 5A). To study the direct role of Spef2 in osteoblast differentiation, calvarial osteoblasts were isolated from WT and Spef2 KO mice and cultured for 21 days in vitro. Samples were collected at D0, 7, 14 and 21 for Oss, Alp, and Ocn gene expression analysis. Interestingly, the expression of all studied genes was decreased in Spef2 KO osteoblasts during the in vitro differentiation (Fig. 5B–D). Decreased Oss expression (Fig. 5B) suggests a defect in early osteoblast differentiation and the decreases in Alp (Fig. 5C) and Ocn (Fig. 5D) indicate lower mineralization capacity of the Spef2 KO osteoblasts. These results were supported by the decreased von Kossa staining in Spef2 KO osteoblast cultures (Fig. 5E,F).

We also analyzed the expression of cathepsin K and tartrate-resistant acid phosphatase in WT and Spef2 KO bones and did not detect any significant differences suggesting that the differentiation and function of osteoclasts was not affected by Spef2 depletion (data not shown). This was further supported by similar expression levels of

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Figure 1. Spef2 is expressed in the bone and cartilage. (A) Spef2 RNA expression was detected in the bone and cartilage tissues and significantly decreased in Spef2 KO mice. RNA was extracted from the proximal tibia and cartilage and the same exposure was used for different gels. (B) Spef2 expression was increasing during the osteoblast differentiation in vitro and also present in chondrocytes. Primers used for Spef2 RT-PCR (A) and qPCR (B) are located in exon 1 and exon 3. Error bars are presented as ± SD.
receptor activator of nuclear factor kappa-B ligand and osteoprotegerin in calvarial osteoblasts isolated from WT and Spef2 KO (data not shown). Altogether these results suggest that the defective bone formation in Spef2 KO mice is not caused by impaired osteoclast differentiation.

Since bone growth was impaired in Spef2 KO mice, we analyzed the expression levels of several markers for chondrocyte differentiation. The expression of mineralization marker, Ocn, as well as the expression of Collagen 2a1 (Col2a1), Sox9, Runx2 and Col1 was studied in the growth plate cartilage of proximal tibia at P3 and P15, but no significant differences were observed between WT and Spef2 KO cartilage (Supplemental Fig. S4). However, a slight decrease in Ocn expression was detected in Spef2 KO mice compared to WT at P15. Although we did not observe any major defects in chondrocyte function, it has to be noted that Spef2 mRNA was shown to be expressed in cultured chondrocytes (Fig. 1B). Thus, more careful analysis is required to conclude about the potential role of Spef2 in chondrocytes.

**Discussion**

Previous studies have indicated the importance of SPEF2 in ciliated tissues, especially during sperm development, tracheal cilia beating and in ependymal cilia function. In this study, we show for the first time that SPEF2 is also required for bone formation. In our mouse model with the full inactivation of Spef2 gene caused by the...
introduction of a stop codon after exon 2, we observed severe postnatal growth retardation in addition to previously identified PCD phenotypes. The phenotype of the Spef2 KO is more pronounced than the phenotype of bgh mice that is caused by a missense and nonsense mutation in Spef2 gene. SPEF2 protein is known to have

**Figure 3.** μCT analysis of the trabecular and cortical bone in Th10 and L2 vertebrae. (A) Regions of Interest (ROIs) of the trabecular (tbr) and cortical (cb) vertebrae are marked. (B–D) Trabecular number (Tb.N) (B), thickness (Tb.Th) (C) and open porosity (Po(op)) (D) indicate that depletion of Spef2 affects the trabecular bone content and structure. (E–G) Cortical bone volume (BV) (E), surface (BS) (F) and connectivity (conn) (G) suggests decreased bone volume and structure in the Spef2 KO vertebra. Error bars ± SD; * = p < 0.05; ** = p < 0.01; *** = p < 0.001.

### Table 1. μCT data from vertebra Th10 and L2 trabecular bone at P15 and P31. Error bars ± SD, * = p < 0.05; ** = p < 0.01; *** = p < 0.001.

|       | WT, P15 | KO, P15 | WT, P31 | KO, P31 |
|-------|---------|---------|---------|---------|
|       | Tissue volume (TV), mm³ | 0.581 ± 0.0237 | 0.328 ± 0.0466<sup>a</sup> | 0.087 ± 0.2623 | 0.565 ± 0.1641 |
|       | Bone volume (BV), mm³ | 0.109 ± 0.0235 | 0.066 ± 0.0098<sup>b</sup> | 0.341 ± 0.1493 | 0.158 ± 0.0872 |
|       | Bone volume density (BV/TV), % | 18,735 ± 3,5152 | 20,055 ± 2,3983 | 37,800 ± 7,6047 | 26,283 ± 9,5989 |
|       | Tissue volume (TV), mm³ | 1,1436 ± 0.091 | 0.698 ± 0.1369<sup>ab</sup> | 1.637 ± 0.6347 | 0.917 ± 0.113 |
|       | Bone volume (BV), mm³ | 0.204 ± 0.0317 | 0.147 ± 0.0389 | 0.648 ± 0.2709 | 0.199 ± 0.0591<sup>a</sup> |
|       | Bone volume density (BV/TV), % | 18,010 ± 3,8619 | 20,958 ± 1,9395 | 39,191 ± 4,1460 | 21,428 ± 4,5726<sup>ab</sup> |
various splicing variants (Supplemental Fig. S1B), of which all N-terminal splicing variants are eliminated in Spef2 KO mice. This may explain the phenotypic differences between the different Spef2 mutant mouse models, and it is likely that the depletion of short N-terminal variants in addition to the full length Spef2 causes the bone phenotype. The sperm tail phenotype appears to be identical in all Spef2 mutant animal models and is most probably caused by the lack of the full length isoform of SPEF2 protein. This study underlines the importance of the N-terminal part of SPEF2 in various tissues including tissues with primary cilia. Although depletion of SPEF2 appears to have a clear effect only on the structure of the sperm tail axoneme, it obviously does have a role in cilia motility and potentially in signaling pathways in primary cilia.

The clear changes in various skeletal parameters in Spef2 KO mice suggest that SPEF2 has a role in bone formation. Although we cannot rule out the possibility that some of the growth defects originate as secondary effects of severe PCD symptoms such as hydrocephalus, several findings suggest that SPEF2 is directly involved in the regulation of bone formation. First, we showed that Spef2 is expressed in the bone and cartilage, and Spef2

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**Figure 4.** μCT analysis of the distal femur trabecular and cortical bone. (A) ROI for the trabecular bone (tbr) was drawn starting from the distal growth plate of the femur, extending for 1 mm towards diaphysis, excluding the cortical bone material. ROI for the cortical bone (cr) was drawn starting from 1 mm from the distal growth plate to the diaphysis, extending a longitudinal distance of 1 mm. (B–D) Trabecular number (Tb.N) (B), thickness (Tb.Th) (C) and open porosity (Po(op)) (D) indicated reduced trabecular bone content and impaired structure. (E–G). Cortical bone volume (BV) (E), surface (BS) (F) and connectivity (conn) (G) showed decreased cortical bone volume and structure in Spef2 KO mice. Error bars ± SD; * = p < 0.05; ** = p < 0.01; *** = p < 0.001.

|                | WT, P15      | KO, P15      | WT, P31      | KO, P31      |
|----------------|--------------|--------------|--------------|--------------|
| Tissue volume (TV), mm³ | 1,751 ± 0,1775  | 1,670 ± 0,0802² | 2,022 ± 0,3248  | 2,973 ± 1,6425 |
| Bone volume (BV), mm³ | 0,217 ± 0,0507  | 0,031 ± 0,0179² | 0,472 ± 0,2936  | 0,661 ± 0,5184 |
| Bone density volume (BV/TV), % | 12,386 ± 2,3503 | 2,805 ± 1,4952² | 21,635 ± 10,7071 | 15,532 ± 4,8941 |

Table 2. μCT data from distal femur trabecular bone at P15 and P31. Error bars ± SD, * = p < 0.05; ** = p < 0.01; *** = p < 0.001.
expression in osteoblasts increases during in vitro differentiation. The direct role of SPEF2 is further supported by defective in vitro differentiation of Spef2 KO osteoblasts. The osteoclast function appeared unaffected indicating that the observed phenotype was not due to imbalanced bone remodeling. Furthermore, although Spef2 was expressed in the cartilage we were unable to observe any significant differences in the expression of specific markers for chondrocyte differentiation. This suggests that the mineralization defect observed in Spef2 KO mice originates mostly from the defective osteoblast function rather than impaired chondrocyte differentiation and function. However, additional studies are required to confirm the exact role of SPEF2 in cartilage formation.

The effect of SPEF2 depletion in osteoblasts is evident, but the mechanism how SPEF2 regulates osteoblast function and the process of bone formation requires further studies. The existing evidence supports the hypothesis, that the SPEF2 function is associated with cilia-related processes. We have shown that the cilia structure appears intact in Spef2 mutant models, and therefore, the phenotype is not caused by the absence of cilia (Supplemental Fig. S3)39. On the other hand, cilia motility was altered in Spef2 mutant mice, suggesting that SPEF2 is required for the normal function of cilia39. Interestingly, recent data strongly suggests that SPEF2 functions in the intracellular transport of proteins or vesicles via microtubules. SPEF2 interacts with intraflagellar transport protein IFT20 and colocalizes with IFT20 in differentiating male germ cells30. Furthermore, we have shown that SPEF2 interacts with a motor protein Dynein 1, and Dynein activity is required for the correct localization of SPEF2 in elongating spermatids31. Thus, SPEF2 may have a role in central protein transport or cilia-related signaling pathways required for osteoblast differentiation.
In addition to the canonical role the SPEF2 interaction partner IFT20 in cilia-related transport, it has been shown to function in intracellular trafficking between different cellular compartments.\textsuperscript{20-35} For example, during craniofacial skeletal development, IFT20 has been shown to be involved in the intracellular trafficking of procollagen from the endoplasmic reticulum (ER) to the Golgi complex\textsuperscript{25}. IFT20 depletion caused a severe delay in the exocytosis of the matrix protein collagen 1 from the osteoblasts leading to osteopenia\textsuperscript{25}. Moreover, IFT20 has been shown to mediate polycystin-2 (PKD2) trafficking to cilia from the ER through the Golgi complex\textsuperscript{38}. Osteoblast specific depletion of \textit{Pkd2} has been shown to result in similar defects in bone mineral density, trabecular bone volume and cortical thickness, and expression of osteoblast related genes e.g. \textit{Ocn} and \textit{osteopontin}\textsuperscript{38} that were observed in \textit{Spef2} KO mice. On the basis of these studies, it is possible that SPEF2 and other cilia- and IFT-related genes have diverse roles in osteoblasts, both in cilia and non-cilia associated sites. Even though the detailed molecular mechanisms of SPEF2 function in osteoblasts remain to be characterized, our results highlight a novel important role of SPEF2 in the bone formation and mineralization in mice.

**Material and Methods**

**Generation of \textit{Spef2} full knockout mouse model.** Mice BAC clone (RP23-340E4) containing exons 3-5 (including 3' and 5' -flanking regions) from \textit{Spef2} gene (chromosome 15) was purchased from Children's Hospital Oakland Research Institute (Oakland, CA, USA). All primers used for targeting construct are listed in Supplemental Table S1. Shaving part containing ampicillin resistance and 50 bp homology arms were cloned from pACYC177 plasmid. Shaving part was electroporated into electrocompetent \textit{E. coli} cells containing BAC clone and pRedET for recombination. LoxP-PGK-tn5-neo-loxP – cassette was cloned from pgKneo10xp plasmid and recombined into Shaved BAC clone after exon 5. Neo cassette was removed from the shaved BAC clone using 294-Cre \textit{E. coli} cells leaving only one LoxP site after exon 5. LoxP, Frt and DsRed2 were cloned from pRES2-DsRed2 plasmid. Targeting vector was digested with NheI, SpeI, and BgIII (Promega Corporation, Madison, WI, USA) according to manufacturer's instructions to confirm the correct orientation and insertion of all cloned inserts. \textit{Spef2} targeting vector was linearized using Sac II restriction enzyme (Promega, Madison, WI, USA). Linearized \textit{Spef2} targeting vector was electroporated into hybrid mouse embryonic stem cells (G4, 129S6B6F1) and homologous recombination was screened by PCR. To generate chimeric mice, ES cells were injected into blastocysts of C57BL/N6 mice and targeting vector integration to the genome was detected from the DNA samples of pups isolated from ear marks. Primer sequences for \textit{Spef2} genotyping PCR are listed in Supplemental Table S1. Three chimeric male mice were born and were bred with C57BL/6NShsd females. Two founders transferred the transgene into the next generation. Mice used for further analysis originated from heterozygous breedings (N1) and were maintained as inbred for several generations. Control mice for the experiments were used WT mice originating from \textit{Spef2} heterozygous breedings. WT and \textit{Spef2} HEZ mice presented normal health status while \textit{Spef2} KO mice presented the phenotype caused by depletion of the \textit{Spef2} gene.

**Ethical statement.** Mice were sacrificed with CO\textsubscript{2} or cervical dislocation and thereafter tissues were collected for all experiments. All mice were maintained in a specific pathogen-free stage at the Central Animal Laboratory of the University of Turku and handled in accordance with international guidelines on the care and use of laboratory animals. Studies were approved by the Finnish ethical committee for experimental animals (license 315/041003/2011).

**Animal material.** For all experiments male mice were used except for \textit{in vitro} studies, where collected material was pooled from animals with a specific genotype. Age matched WT mice were used as controls from the same litter, when possible. All experiments were repeated (number of animals used is indicated for each experiment) except the whole-mount preparation that was done only for one animal per genotype.

**Sperm motility analysis.** The cauda epididymis of WT (n = 8) and \textit{Spef2} HEZ (n = 10) mice (8–9 week old) was dissected and placed in +37 °C KSOM medium with amino acids (Merck Millipore). Sperm was collected by making small incision to cauda and let them swim out at +37 °C for 30 min. Sperm motility was investigated under microscope, where 100 spermatozoa were counted in every sample and classified as progressive motility (PR), non-progressive motility (NP) or immotile (IM). The average of duplicates was used for analysis.

**Histology of the testis.** Testis of adult WT and \textit{Spef2} HEZ mice was dissected and fixed with Bouin's solution, washed several times with 70% ethanol and embedded in paraffin. Paraffin embedded testis were cut into sections, deparaffinized, rehydrated and stained with Mayer's Hematoxylin (Histolab, Västra Frölunda, Sweden) and eosin. Sections were mounted using Pertex (Gibco, Thermo Fischer Scientific, Waltham, MA, USA) mounting media.

**Whole-mount preparations.** For whole-mount preparations skeletons of P31 male mice (n = 1 for WT and \textit{Spef2} KO) were dissected as whole and fixed in 95% ethanol for four days. The skeletons were stained overnight with Alcian Blue 8 G, washed with ethanol, cleared in 1% KOH for 8 h, stained with Alizarin red for 8 h, and cleared in 2% KOH until the staining was easily observable. The preparations were brought to glycerol in ascending concentrations and photographed on a projection table.

**X-ray analysis and micro-computed tomography.** X-ray images of formalin fixed hind limbs (n = 4 for WT, n = 3 for \textit{Spef2} KO) and spinal columns (n = 4 for WT, n = 3 for \textit{Spef2} KO) were obtained by using Faxitron LX-60X-ray imaging device (X-Ray LLC, Lincolnshire, IL, USA). To determine the three-dimensional
structure of the bones, femora, thoracic vertebral bone Th10 and lumbar bone L2 vertebrae were imaged by micro-computed tomography (µCT, SkyScan 1072 Micro-CT device, Kontich, Belgium) and the data was reconstructed by using the Nrecon 1.6.9.4 software and modeled and analyzed by CTan 1.13.5.1 software (SkyScan). In µCT analysis image pixel size of 4.18467 µm (P15 group) or 7.32622 µm (P31 group) for vertebral bones, and 9.76926 µm for femora was applied; X-ray tube potential of 70 kV and current of 148 µA were used with integration time of 3900 ms and rotational step of 0.45 degrees. During reconstruction smoothing level of 3, beam-hardening reduction of 85%; and ring artifact reduction level of 7 were used with attenuation coefficient value range of 0.006 to 0.15.

Mechanical testing. Bone mechanical properties were evaluated by applying 3-point bending test by using Nexygen program (Lloyd Instruments, West Sussex, United Kingdom). Left femora of P31 male mice (n = 4 for WT, n = 3 for Spef2 KO) were placed on two holders located at a range of 3 mm. The bending force was applied at a crosshead speed of 5 mm/min to the middle until fracture occurred. Maximum load (N) was obtained directly from the load-deformation curve.

RNA analysis. For mRNA analysis tissues were collected, snap frozen in liquid nitrogen and stored at −80°C. For gene expression analysis of the proximal tibiae cartilage and trabecular bone mass (n= 2 for WT, n= 2–3 for Spef2 KO) were collected in RNAlater stabilization solution (RNAlater RNA stabilization reagent, Qiagen, Hilden, Germany) and stored at −80°C. Total RNA was extracted by applying the RNeasy Midi kit (Qiagen) following manufacturer’s instructions.

Real-time PCR (RT-PCR). For analysis of gene expression with RT-PCR the total RNA was reverse transcribed with random primers and an RT-PCR kit (ImProm-II Reverse Transcription System, Promega) according to the manufacturer’s instructions. Produced cDNA was amplified by using gene specific primers listed in Supplemental Table S1. Housekeeping gene Rpl13a or Eef2 was used as a reference gene to calculate the relative expression. The qPCR was performed with a ViiA™ 7 Real-Time PCR System in 96-well microtiter plates using Absolute qPCR SYBR Green ROX Mix (VWR, Radnor, PA, USA). Amplification by qPCR contained 12.5 µl of Absolute qPCR SYBR Green Mix, 100 ng of cDNA, and 70 nM of each primer in a final volume of 25 µl. Amplifications were initiated with 15 min enzyme activation at 95°C followed by 40 cycles of denaturation at 95°C for 15 s, primer annealing at 60°C for 1 min, and extension at 72°C for 30 s. All samples were amplified in triplicate, and the mean value was used for further calculations. Raw data were analyzed with the sequence detection software (Applied Biosystems, Foster City, CA, USA) and relative quantitation was performed with GenEx software (MultiD, Göteborg, Sweden). Ratios between the target and reference gene were calculated by using the mean of these measurements. A standard curve for each primer pair was produced by serially diluting a control cDNA and used to correct the differences in amplification. A melting curve analysis was performed allowing single product-specific melting temperatures to be determined. No primer–dimer formations were generated during the application of 40 real-time PCR amplification cycles.

Sanger sequencing. For Sanger sequencing of the Spef2 KO hybrid transcript the cDNA fragment was amplified with Spef2 specific primers in exons 1 and 3 (Supplemental Table S1) and PCR amplicons were purified using ExoSAP-IT (Amersham Biosciences). PCR fragments were sequenced in both directions with amplification primers and Dsred primers within the PCR amplicon (Supplemental Table S1). Sequencing was performed on a MegaBace 500 capillary DNA sequencer (Amersham Biosciences) using DYEnamic ET Terminator Kits supplied by the manufacturer with a MegaBace 500 capillary DNA sequencer (Amersham Biosciences) using DYEnamic ET Terminator Kits supplied by the manufacturer. The data was analyzed using the Variant Reporter v1.0 program (Applied Biosystems) and Sequencer 5.2.3 (Gene Codes Corporation).

Calvarial osteoblast culture. Calvarias were collected from three-day old WT and Spef2 KO mice. After decapitation the calvarias were removed and cells were released from the matrix using 1 ml digestion medium [0.1% Collagenase A (Roche Diagnostics, Germany), 0.2% Dispase II (Roche Diagnostics, Germany) diluted in α-MEM] at 37°C with shaking. After 10 min digestion first fraction was collected and discarded and four subsequent fractions were collected in every 20 min and pooled. Cells were plated in α-MEM containing 10% FBS and antibiotics (proliferation medium) in 10 cm dishes. Cells were allowed to proliferate until 80–90% confluency before pooling the same genotypes and seeding into six-well plates cultured until confluency (day 0 timepoint). Osteoblast differentiation was induced with 5 mM sodium beta-glycerophosphate, 10−8 M dexamethasone and 50 µg/ml ascorbic acid in proliferation medium. Differentiation medium was changed every 2–3 days. Samples for RNA extraction were collected at 0, 7, 14 and 21 days by rinsing the wells with PBS, scraping cells off the bottom of the well with a cell scraper and stored in −20°C in RNAlater (Invitrogen, USA).

Cytological stainings and quantification of stained areas in calvarial osteoblast cultures. For cytological stainings, osteoblast cultures were rinsed with PBS and fixed with 3.7% formalin for 10 min. After dH2O washes formalin-fixed wells were stained for ALP with Naphthol AS MX-PO4 (Sigma, USA) dissolved in DMF (Sigma USA) mixed with Fast Blue RR salt (Sigma, USA) in 0.1 M Tris-HCl (pH 8.3). ALP stained wells were stained for von Kossa with 2.5% Silver nitrate (Fisher Scientific, UK) for 30 min exposed to direct light, and washed with dH2O for three times. ALP and von Kossa stained six-well plates were scanned using a flatbed scanner with a transparency adaptor (HP ScanJet 5370 C) and saved as 24-bit color images in TIFF format. Transparency exposure adjustments were maintained constant to create images of equal intensity. Positively stained areas were quantified using Imaging Software ImageJ. RGB images were split into three 8-bit grayscale images containing the red, green and blue components. Threshold and Region of Interest (ROI) were adjusted and kept the same to maintain standard measuring conditions.
Chondrocyte cultures. Primary chondrocytes were isolated from three-day old mouse using standard protocols. Briefly, chondrocytes were digested from mouse ribcages during 2 h digestion in 3 mg/ml collagenase D diluted in Dulbecco's Modified Eagle Medium (D-MEM) at 37 °C. Digestion solution was changed to 0.5 mg/ml collagenase D in D-MEM and ribcages were incubated at 37 °C for overnight. After filtering through 100 μm strainer, chondrocytes were seeded (3 × 10⁶ cells/well) on six-well plates and cultured in D-MEM supplemented with 100 U/ml penicillin-streptomycin, 10% FBS (Gibco, USA), 2 mM L-glutamine and Insulin-Transferrin-Selenium (ITS, Gibco). The culture medium was changed every 2 days and cells were harvested in RNALater at day 6 for RNA isolation.

Electron microscopy. Trachea were isolated from WT and Spef2 KO mice and fixed with 5% glutaraldehyde. Samples were treated with potassium ferrocyanide-osmium fixative and embedded in epoxy resin. Sectioned samples were stained using 1% uranyl acetate and 0, 3% lead citrate. Samples were visualized with JEM-1400 Plus (JEOL).

Immunohistochernistry of the bone and cartilage. Tibia of WT and Spef2 KO (n = 3) mice were collected and fixed in 10% buffered formalin overnight, decalcified in 5% formic acid, embedded in paraffin and cut into 5 μm-thick sections. Tibia sections were digested using ficin (Digest-All 1, Thermo Fischer Scientific) at +37 °C for 10 min after paraffin removal and rehydration. Sections were blocked with 10% normal goat serum and 3% bovine serum albumin diluted in 0,01% Triton X-100 PBS. Primary antibody (anti-acetylated α-tubulin (Sigma-Aldrich, St Louis, MO, USA) 1:1000) was diluted in 3% normal goat serum, 1% bovine serum albumin and 3% bovine serum albumin and 0, 01% Triton-X 100 in PBS and incubated at +4 °C overnight. After washes with 0, 1% Triton-X 100 in PBS sections were incubated with secondary antibody (1:500, goat anti-mouse Alexa 488 (Molecular probes, Eugene, OR, USA)) at room temperature for 1 hour. Sections were mounted with Prolong® Diamond Antifade Mountant (Molecular Probes) and imaged using Leica DMRBE microscope and DFC320 camera.

Statistical analysis of data. Data is presented as average values with standard deviation (SD). Statistical differences were calculated by paired Student's t-test was used for pairwise comparisons between groups. P-values of 0.05 or less were considered statistically significant.

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Author Contributions
M.S.L., H.H., R.K., T.J.H., N.K. and A.S. designed research; M.S.L., H.H., R.K., T.J.H., N.K. and A.S. analyzed data; M.S.L., H.H., P.R., F.W. and L.U.-K. performed research; M.S.L., H.H., R.K., T.J.H., N.K. and A.S. wrote the paper.

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