CNC-bZIP protein NFE2L1 regulates osteoclast differentiation in antioxidant-dependent and independent manners

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ABSTRACT

Fine-tuning of osteoclast differentiation (OD) and bone remodeling is crucial for bone homeostasis. Dissecting the mechanisms regulating osteoclastogenesis is fundamental to understanding the pathogenesis of various bone disorders including osteoporosis and arthritis. Nuclear factor erythroid 2-related factor 1 (NFE2L1, also known as NRF1), which belongs to the CNC-bZIP family of transcription factors, orchestrates a variety of physiological processes and stress responses. While Nfe2l1 gene may be transcribed into multiple alternatively spliced isoforms, the biological function of the different isoforms of NFE2L1 in bone metabolism, osteoclastogenesis in particular, has not been reported. Here we demonstrate that knockout of all isoforms of Nfe2l1 transcripts specifically in the myeloid lineage in mice [Nfe2l1(M-KO)] results in increased activity of osteoclasts, decreased bone mass and worsening of osteoporosis induced by ovariectomy and aging. In comparison, LysM-Cre-mediated Nfe2l1 deletion has no significant effect on the osteoblast and osteocytes. Mechanistic investigations using bone marrow cells and RAW 264.7 cells revealed that deficiency of Nfe2l1 leads to accelerated and elevated OD, which is attributed, at least in part, to enhanced accumulation of ROS in the early stage of OD and expression of nuclear factor of activated T cells, cytoplasmic, calcineurin dependent 1α (Nfatc1/α). In addition, NFE2L1 regulates the transcription of multiple antioxidant genes and Nfatc1/α and OD in an isoform-specific manner. While long isoforms of NFE2L1 function as accelerators of induction of Nfatc1/α and antioxidant genes and OD, the short isoform NFE2L1-453 serves as a brake that keeps the long isoforms’ accelerator effects in check. These findings provide a novel insight into the regulatory roles of NFE2L1 in osteoclastogenesis and highlight that NFE2L1 is essential in regulating bone remodeling and thus may be a valuable therapeutic target for bone disorders.

1. Introduction

Bone is a highly dynamic tissue that is constantly being remodeled. The remodeling of bone is a complex process involving a delicate balance between the activities of osteoblasts and osteoclasts, which are regulated by various factors, including cytokines/chemokines, hormones and mechanical stimuli. Interference with this balance results in a variety of disorders that affect bone integrity. Tipping the balance in favor of osteoclasts leads to pathological bone resorption, which is observed in many bone diseases, including osteoporosis and rheumatoid arthritis [1]. Osteoclasts, the bone-resorbing cells, are multinucleated giant cells derived from mononuclear precursor cells in the monocyte/macrophage hematopoietic lineage [2, 3]. Osteoclastogenesis is a tightly regulated multiple steps process comprising osteoclast
progenitor commitment, macrophage colony-stimulating factor (M-CSF)-induced osteoclast precursor proliferation and receptor activator of nuclear factor-κB ligand (RANKL)-mediated osteoclast differentiation (OD) [4–8]. The binding of RANKL and RANK recruits TRAF6 (tumor necrosis factor receptor-associated factor 6), which activates the downstream signaling molecules including reactive oxygen species (ROS), fos proto-oncogene (c-Fos), nuclear factor kappa-B (NF-κB) and mitogen-activated protein kinases (MAPKs). Physiological ROS is an important second messenger in regulating the differentiation of osteoclasts, and the nuclear factor of activated T cells 1 (NFATc1, also known as NFAT2) signaling could be the crucial downstream signaling event in RANKL-mediated ROS signaling [9]. RANK activation and immunoreceptor tyrosine-based activation motif (ITAM)-mediated calcium signals synergistically regulate the expression of NFATc1 and subsequent signaling mediating OD [10]. NFATc1 regulates a number of osteoclast-specific genes, such as tartrate-resistant acid phosphatase (TRAP), cathepsin K, calcitonin receptor, and osteoclast-associated receptor. Although NFATc1 has been recognized as an orchestrator of OD, the molecular details on NFATc1 transcription are still incompletely understood.

Nuclear factor-erythroid 2-related factor 1 (NFE2L1, commonly symbolized as Nrf1; human analogs are also termed as TCFT11 or LCR-F1) is a member of the CNC-BZIP transcription factor family. Human and mouse NFE2L1 genes can be transcribed into alternatively-spliced forms, resulting in multiple protein isoforms [11,12]. NFE2L1 exerts its transcriptional control on genes harboring the NF-E2/AP1-like ARE/EppRE site by forming heterodimers with small musculoaponeurotic fibrosarcoma (sMAF) proteins, subunits of AP-1 or other bZIP proteins [13,14]. NFE2L1 plays crucial regulatory roles in the transcriptional control of multiple essential biological processes, including antioxidant response, proteasome homeostasis, lipid metabolism and cell differentiation [15–20]. Based on the latest version of Ensembl Genome Browser (Ensembl release 104 - May 2021), the mouse Nfe2l1 gene spans 12,554 bp DNA and is divided into 10 exons. The Nfe2l1 gene may be transcribed into multiple spliced forms due to alternative transcription start and termination sites, resulting in at least two long isoforms (L-NFE2L1) containing 741 and 742 amino acids (aa) and two short isoforms (S-NFE2L1) containing 453 and 583 aa, respectively. The human NFE2L1 can also be transcribed into alternatively spliced forms, resulting in multiple protein isoforms containing 772, 761, 742, 616, 584 and 573 aa, respectively [21–24]. Posttranslational regulation, including glycosylation and proteolytic processing, plays critical roles in the transactivation and stabilization of human L-NFE2L1 [25,26]. The studies using a variety of cell-specific Nfe2l1-knockout mice illustrated that NFE2L1 plays essential regulatory roles in many types of cells, including neurons, hypertrophic, pancreatic β-cells, adipocytes, osteoblasts, yet the specific contribution of each individual isoform of NFE2L1 in these cells still remains to be elucidated [27–32]. With regard to bone metabolism, in vitro studies found that human L-NFE2L1 interacts with CGAAT enhancer-binding protein β (C/EBPβ) to repress the expression of dentin sialophosphoprotein (DSP) during odontoblast differentiation [19]. In response to ascorbic acid induction, NFE2L1 upregulates the expression of ostein in mouse osteoblasts [20]. In addition, targeted deletion of all isoforms of Nfe2l1 in osteoblasts in mice resulted in decreases in bone size, bone formation, trabecular bone, peak bone mass and mechanical strength [32], suggesting that NFE2L1 is involved in regulating osteoblast differentiation and bone formation. Intriguingly, multiple single nucleotide polymorphisms (SNPs) of the human and mouse Nfe2l1 gene appears to be associated with bone density in humans [33] and bone metabolism in mice [34]. However, the isoform-specific biological function of NFE2L1 in bone metabolism, osteoclastogenesis in particular, has not been reported. In the present study, we developed myeloid cell lineage-specific Nfe2l1-knockout (Nfe2l1(M)-KO) mice and various lines of cells to investigate the isoform-specific roles of NFE2L1 in osteoclastogenesis and bone homeostasis.

2. Material and methods

Mice. Nfe2l1(M)-KO mice were generated by crossing the mice bearing a Nfe2l1 allele flanked by LoxP sites and LysM-Cre allele (J004781, Jackson Laboratory, Bar Harbor, ME) as detailed previously [35–37]. Nfe2l1(M)-KO mice with the genotype of Nfe2l1<LoxP/LoxP> LysM-Cre<LoxP> and their littermates with the genotype of Nfe2l1<LoxP/LoxP> LysM-Cre<+/+> were used as the primary controls in this study. Skeletal phenotype of a group of mice with the genotype of LysM-Cre<+/+> was also examined to investigate the effect of LysM-Cre transgenic alone on bone mass and structure. Genotyping was performed using genomic DNA that was isolated from tail snips as previously described [38]. The sequences of genotyping PCR primers are shown in Table S1. The mice were housed up to four per cage in virus-free facilities on a 12-h light/12-h dark cycle and fed NIH07 chow diet (Jiangsu Xietong BioTech, Nanjing, China) and provided reverse osmosis water ad libitum. All protocols for animal use were approved by the Institutional Animal Care and Use committee of China Medical University (Shenyang, China), which is in accordance with the guidelines of US National Institutes of Health.

Bilateral ovariectomy (OVX) to induce osteoporosis in mice. Bilateral OVX was performed as reported previously [39]. In brief, the female mice at age of 12 weeks were first weighed and anesthetized using an intraperitoneal injection of pentobarbital sodium (70 mg/kg). Then bilateral OVX was carried out in the OVX group to remove ovaries and fallopian tubes. The mice in sham control group were subjected to laparotomy but ovaries and fallopian tubes were not removed. Surgically removed ovaries were examined histologically to confirm successful removal of bilateral ovaries.

Analysis of bone volume and mineral density. Following euthanization via CO2 asphyxia, the femurs of the mice were removed immediately and kept wrapped in gauzes soaked in PBS at 4 °C. Microcomputed tomography (Micro-CT, Skyscan 1176 and 1276; Bruker-MicroCT, Kontich, Belgium) was used to measure trabecular bone volume within the metaphysis of the femurs as detailed previously [40]. The trabecular bone volume fraction (bone volume/total volume (BV/TV)), trabecular number (Tb.N), trabecular separation (Tb.Sp) and trabecular thickness (Tb.Th) were measured and analyzed by Skyscan CTAn software (v.1.1.7, Skyscan CTAn, Kontich, Belgium). A dual-energy X-ray absorptiometry (DXA, Hologic Discovery DXA system, Hologic Inc., Waltham, MA) was used to determine BMD according to the manufacturer’s instructions.

Histological analysis. The femurs isolated were fixed in 10% paraformaldehyde for 1–2 days, followed by decalcification in 10% EDTA (pH = 7.0) for 6 weeks prior to embedding in paraffin. Serial paraffin sections (5 μm) were stained for H&E and TRAP using a Leukocyte Acid Phosphatase Kit (387A, Sigma, St. Louis, MO) according to the manufacturer’s instructions. Histomorphometric analyses of osteoclast and osteoblast numbers were performed using the Image J (NIH Image, Bethesda, MD).

Cell culture of BM-derived OPCs and RAW264.7 cells. The BMCs flushed from femurs and tibias of 8–12 week-old male Nfe2l1(M)-KO or Flox control mice (n = 4–6 mice each condition) were pooled and cultured in minimum essential medium alpha (α-MEM) containing 10% heat-inactivated fetal bovine serum (HI-FBS), 100 units/ml penicillin and 100 μg/ml streptomycin for 24 h. Then the suspended cells were harvested and further cultured with 30 ng/ml M-CSF in α-MEM containing 10% (v/v) of HI-FBS. After 3 days of culture, the adherent cells were used as BM-derived OPCs for osteoclast differentiation studies. The mouse leukemic monocyte/macrophage cell RAW264.7 (RAW) were purchased from the American Type Culture Collection (ATCC, Manassas, VA) and cultured in Dulbecco’s modified essential media (DMEM) supplemented with 10% (v/v) HI-FBS, 100 units/ml penicillin, 100 μg/ml streptomycin and 10 mmol/L HEPES at 37 °C in a 5% CO2 incubator. Phosphate buffered saline (PBS, pH 7.4) and supplements for cell culture were purchased from Life Technologies (Grand Island, NY). All other
reagents including M-CSF (416-ML-010) and RANKL (462-TEC-010) were purchased from R&D Systems (Minneapolis, MN).

**Induction of OD in vitro and TRAP staining.** The BM-derived OPCs and RAW cells were plated onto 24-well plates at a density of 8 × 10^3 and 2 × 10^5 cells/well, respectively. Then the cells were cultured in α-MEM containing 10% (v/v) FBS, 100 units/ml penicillin, 100 μg/ml streptomycin and 10 mmol/L HEPES in the presence of 50 ng/ml recombinant RANKL and 30 ng/ml M-CSF. The culture media were replaced every 2 days. Following 5–7 days of culture, the cells were washed twice with PBS, fixed with 4% paraformaldehyde and stained for TRAP using a Leukocyte Acid Phosphatase Kit (387A, Sigma, St. Louis, MO) according to the manufacturer’s instructions. TRAP-positive red-colored cells with 3 or more nuclei were considered as mature osteoclasts.

**Bone resorption assay.** BM-derived OPCs (8 × 10^4 cells/cm²) and RAW cells (2 × 10^6 cells/cm²) were seeded on the Osteo assay plate (3987, Corning, Kennebunk ME) and cultured in α-MEM containing 10% (v/v) FBS, 100 units/ml penicillin, 100 μg/ml streptomycin, 10 mmol/L HEPES, 30 ng/ml M-CSF and 100 ng/ml RANKL. Following 7 days of culture, the cells were removed from the plates with bleach and the calcium phosphate substrate was washed with 5% sodium hypochlorite. The pit formation was observed and imaged under an optic microscope (Axio Lab.A1 pol, Leica, Germany). The resorbed area was also measured by image J and expressed as percentage of the control values.

**Detection of F-ACTIN ring formation.** Differentiated osteoclast cells were fixed with 4% PFA for 15 min, permeabilized with 0.1% Triton X-100 for 5 min and stained with Rhodamine-Phalloidin (KGP001, KeyGEN BioTECH, China) (20 min) and DAPI (KGR0001, KeyGEN BioTECH, China) (5 min) subsequently. Representative images of F-ACTIN belts formation were observed with fluorescence microscope (DMi8, Leica, Germany).

**Analysis of bone turnover markers in plasma.** The plasma levels of TRAP5b, C-terminal telopeptides of type 1 (CTX1), RANKL, osteoproterin (OPG) and N-terminal propeptide of type 1 procollagen (P1NP) were measured using the enzyme-linked immunoassay (ELISA) kits from Elabscience (Wuhan, China) according to the manufacturer’s protocols. The absorbance was determined with a Flex Station 3 MultiMode microplate reader ( Molecular Devices, Sunnyvale, CA) within the linear range of the assay, and the accuracy of the analysis was verified by the controls provided in each assay kit.

**Lentiviral-based short-hairpin RNA (shRNA) transduction.** RAW cells were transfected with lentivirus containing shRNA against all isoforms of Nfe2l1 (SHVRSNM_010902, Sigma), L-Nfe2l1 (SHCLUD-NM_008686) or Scramble non-target negative control (SHC002V, Sigma) as described previously [41]. Scramble shRNA does not target any known mRNA. The packaging of lentiviral particles was performed based on the manufacturer’s protocol as supplied. After transfection, the cells were selected and maintained in culture medium containing 2.5 μg/ml puromycin.

**Lentiviral-based Nfe2l1 overexpression.** The lentiviral transfer vectors encoding mouse Nfe2l1-741, 741, 583, and 453 isoforms were developed as detailed previously [18]. The lentiviral transfer vectors encoding mouse Nfe2l1-741-V5 and 453-V5 were constructed by cloning PCR-generated fragments into the lentivector pLVX-IRE6-NEO (FP2374, St. Louis, MO) and pTK6-42, a gift from Dr. Tal Kafri, University of North Carolina at Chapel Hill, respectively. Following transfection, the Nfe2l1-741 and 453-V5 were selected and maintained in culture medium containing 2 μg/ml blastocidin and 500 μg/ml G418, respectively. In the current study, we define cells transfected with negative control lentiviral vectors as control (Cont) in contrast to the cells overexpressing an isoform of NFE2L1.

**RT-qPCR.** Total RNA was extracted using TRizol reagent (Life Technologies, Carlsbad, CA) according to manufacturer’s instruction and followed by running through an RNase-free Recombinant DNase I set (TaKaRa) to clean up potential DNA contamination. The resultant RNA was dissolved in RNase-free H₂O and quantified by Nanodrop (Thermo, Wilmington, DE) at 260 nm. RNA samples were stored at –80 °C until use. The reverse transcription was conducted with iScript cDNA Supermix (TaKaRa). Real time PCR was performed with SYBR mix (TaKaRa, Dalian, China) using a Q6 instrument (Applied Biosystems, Foster City, CA). The PCR results were analyzed using a delta CT method, where β-actin was employed as loading control for normalization. The primers were designed using Primer Express (Applied Biosystems) and synthesized by Life Technologies (Shanghai, China). The primer sequences are listed in Supplementary Table S2.

**Western blot analysis.** Western blot was performed as detailed previously [42]. The nuclear and cytoplasmic extracts were prepared using a nuclear extraction kit (P0027, Beyotime, China). The protein concentrations of cell samples were measured with BCA Protein Assay Reagent (TaKaRa). The molecular weight (MW) of each protein shown on immunoblots was estimated based on the SeeBlue® Plus2 Stained Standard (Life Technologies) or ColorMixed Protein Marker (11-245KD) (PR1920, Solarbio, China). The images of immunoblots were quantified with Image J software. The relative levels of target proteins were normalized by β-ACTIN, LAMIN B or β-TUBULIN. The antibodies used in the current study were listed in Supplementary Table S3.

**Intracellular ROS assay.** Intracellular ROS levels were detected using the 2’,7’-dichlorofluorescin diacetate (DCFH-DA, S0033, Beyotime Biotechnology, China). In brief, BM-derived OPCs and RAW 264.7 cells were seeded onto 6-well plates or 24-well plates. Following a 24-h incubation, the cells were preloaded with 5 μM DCFH-DA for 30 min in the dark, and then were treated with RANKL and/or NAC for indicated time. The DCF fluorescence were either visualized using a fluorescence microscope (DMi8, Leica, Germany) or measured by flow cytometry (Becton Dickinson FACSCanto II, Becton Dickinson, San Jose, CA) using excitation/emission wavelengths of 488/525 nm. To avoid photooxidation of DCF, the fluorescence image was collected by a single rapid scan (4-line average, total scan time of 3 s).

**Chromatin immunoprecipitation (CHIP) assay.** A CHIP assay was performed with a ChIP kit (57976s, Cell Signaling Technology, Danvers, MA) according to the manufacturer’s instructions using antibodies against anti-V5 (ab9116, Abcam, MA) and control IgG (A7016, Beyotime, China). The precipitated DNA was subjected to PCR amplification with primers specific for the P1-promoter region of Nfatc1. The primers used in the assay are listed in Supplementary Table S4.

**Nfatc1 promoter-luciferase reporter assay.** Nfatc1 promoter-driven luciferase reporters were constructed by cloning 1073, 711, 520, 265, 111 and 79 bp segments upstream of transcription start site into the pGL4.10 [luc 2] vector (Promega, Madison, WI). In addition, point mutations targeting the core of CREB/Fos/ATF2 binding sites were introduced into various vectors using the TaKaRa MutanBEST Kit (R401, TaKaRa). Transfections were carried out using a FuGENE kit (Roche, Basel, Switzerland) according to the manufacturer’s instructions.

**Statistics.** All statistical analyses were performed using Graphpad Prism 6 (GraphPad Software, San Diego, CA), with p < 0.05 considered as significant. Data were expressed as mean ± standard deviation. For comparisons between groups, a Student’s t-test was performed. For comparisons among multiple groups, one-way or two-way ANOVA with Bonferroni post hoc testing was performed.

**3. Results**

**Nfe2l1(M)-KO mice exhibit reduced bone mass and elevated osteoclast activity.** The BM-derived OPCs from Nfe2l1(M)-KO mice exhibited 56.8% reduction of Nfe2l1 mRNA expression relative to their littermate Flox controls. In contrast, there were no significant changes in other tissues, including the liver, lung, heart, skeletal muscle and white adipose (Supplementary Fig. 1A). Immunoblotting of the OPCs using an antibody against all isoforms of NFE2L1 (sc-13031) showed that multiple bands with molecular weights (MW) ranging between 25–140 kDa were substantially diminished in Nfe2l1(M)-KO mice compared to their
littermate Flox controls (Supplementary Fig. 1B). The reduction of NFE2L1 at both mRNA and protein levels in the OPCs, but not in other tissues, demonstrate that the myeloid lineage-specific knockout of Nfe2l1 is successful.

As shown in Fig. 1A and B, female Nfe2l1(M)-KO mice at age of 20 weeks exhibited substantially reduced cortical and trabecular bone volumes compared to their littermate Flox controls. The Nfe2l1(M)-KO mice displayed a significantly reduced trabecular bone volume (BV/TV) by 15.5% and trabecular number (Th.N) by 16.9% and increased trabecular space (Th.Sp) by 14.9% compared to the Flox controls, whereas the trabecular thickness (Th.Th) showed no significant differences between the genotypes. To exclude the potential influence of littermate Flox controls (Supplementary Fig. 1B). The reduction of Nfe2l1(M)-KO mice exhibited substantially reduced cortical and trabecular bone volume compared to 20 weeks old female mice, as evidenced by significantly reduced BV/TV by 46.2%, Th.N by 26.9% and Th.Th by 25.9%, and increased Th.Sp by 14.6% in comparison with the Flox controls. In addition, the significantly lowered BMD of Nfe2l1(M)-KO mice was further verified by dual x-ray absorptiometry measurements (Supplementary Figs. 1G and H).

To further assess the role of NFE2L1 in aging-induced bone loss, the bone mass of aged mice was examined. As shown in Fig. 1E and F, 110 weeks old female Flox control mice displayed a dramatic reduction in cortical and trabecular bone volume compared to 20 weeks old female Flox mice (see Fig. 1A and B). Both female and male aged Nfe2l1(M)-KO mice showed markedly reduced cortical and trabecular bone volume compared to their littermate Flox controls (Fig. 1E–H).

Histomorphometric analysis, TRAP staining, showed significant increases in the number of osteoclasts and osteoclast surface in 20 weeks old female Nfe2l1(M)-KO mice compared to their littermate Flox controls (Fig. 1I and K). Consistent with the histomorphometric data, Nfe2l1(M)-KO female mice presented elevated levels of TRAP5b and C-terminal telopeptide (CTX) in plasma (Figure 1L), indicating that the mice have enhanced osteoclast activity. To further verify that Nfe2l1 deficiency induces osteoclast activation, we performed TRAP and H&E staining in 110-week-old female mice. As shown in Fig. 1J, M and N, the aged Nfe2l1(M)-KO mice also exhibited significant increases in the number of osteoclasts and osteoclast surface compared to the age-matched Flox controls. To assess the effect of LysM-Cre-mediated Nfe2l1 deletion on osteoblasts, the osteoblasts number and function were determined. Furthermore, osteostogenic cytokines RANKL and OPG, produced by osteoblasts and osteocytes, were determined in plasma. As shown in Figure 1L, N and Supplementary Fig. 2, the number of osteoclasts counted on the sections with H&E staining and plasma levels of RANKL, OPG and P1NP showed no significant differences between Flox and KO groups, indicating that LysM-Cre-mediated deficiency of Nfe2l1 does not affect the number and function of osteoblasts and osteocytes.

Lack of Nfe2l1 in myeloid lineage cells results in increased OD and osteoclast activation. As shown in Fig. 2A and B, the BM-derived OPCs from Nfe2l1(M)-KO mice showed dramatically accelerated and elevated OD compared to the cells from Flox controls. Following 5 days of induction, the number and size of TARP cells in Nfe2l1(M)-KO mice were substantially increased compared to Flox control cells. In addition, the differentiated osteoclasts derived from Nfe2l1(M)-KO mice contained much more nuclei and expressed higher mRNAs levels of Cathepsin K, H'-Apase and Atp6v0d2, which encode proteins crucial for extracellular matrix degradation and bone resorption (Fig. 2C).

To further validate the findings in primary OPCs, we developed a line of RAW264.7 cells with stable knockdown of Nfe2l1 by using a shRNA against all the spliced isoforms of Nfe2l1 (Nfe2l1-KD) (Supplementary Fig. 3A). Compared with the RAW cells expressing non-specific control shRNA (Scramble), Nfe2l1-KD cells had significantly reduced mRNA expression of Nfe2l1, as determined by RT-qPCR with a primer set targeting a common sequence of all 4 isoforms of Nfe2l1 (Fig. 2H).

Immunoblotting analysis using the antibody against all isoforms of NFE2L1 showed that multiple bands with molecular weights (MW) approximately at 65 and 120 KDa were substantially induced by acute oxidative stressor asenite in Scramble RAW cells, whereas the protein levels of NFE2L1 under non-stressed condition was almost undetectable (Fig. 2I). Importantly, lentiviral shRNA-induced knockdown of Nfe2l1 blocked the induction of NFE2L1 by asenite exposure (Fig. 2I), indicating that Nfe2l1 was silenced by Nfe2l1 shRNA. Similar to the OPCs isolated from Nfe2l1(M)-KO mice, accelerated and elevated OD was found in Nfe2l1-KD RAW cells. Compared to Scramble RAW cells, Nfe2l1-KD cells formed more multinucleated giant cells and expressed higher levels of Cathepsin K, H'-Apase and Atp6v0d2 (Fig. 2J-L). The giant osteoclasts derived from Nfe2l1-KD cells also exhibited drastically higher lytic activity than Scramble cells when cultured in calcium phosphate-coated plates (Figure 2M and N). The number of F-ACTIN ring of osteoclast in Nfe2l1-KD cells was significantly higher than that in Scramble cells (Fig. 2F and G).

Nfe2l1 deficiency–induced augmentation of OD in BM-derived OPCs and RAW 246.7 cells are associated with ROS accumulation. The BM-derived OPCs from Nfe2l1(M)-KO mice exhibited reduced mRNA expression of multiple antioxidant genes, including Gclc, Gclm, Cat and Gsr, relative to the cells from Flox controls (Fig. 3A). In line with the downregulation of antioxidant genes, intracellular ROS levels in the OPCs of Nfe2l1(M)-KO mice were significantly higher than those in control cells under basal and RANKL-stimulated conditions. In addition, the elevated accumulation of ROS in Nfe2l1(M)-KO cells were dramatically blocked by the addition of a ROS scavenger N-acetylcysteine (NAC) (Fig. 3B). To understand the effect of increased ROS accumulation in Nfe2l1(M)-KO cells on OD, a set of in vitro OD assays were conducted. As shown in Fig. 3C and D, NAC treatment significantly decreased the number of TARP cells in Nfe2l1(M)-KO and Flox control cells following OD induction, which was strongly supported by the mRNA expression of Cathepsin K, H'-Apase and Atp6v0d2 (Fig. 3E). To validate the key findings in primary OPCs, the same set of measurements, including antioxidant gene expression and key markers of OD, were performed in RAW 246.7 cells. As shown in Fig. 3F-N, Nfe2l1-KD RAW 246.7 cells
Fig. 2. Deficiency of Nfe2l1 induces elevated osteoclast differentiation in BM-derived OPCs and RAW 264.7 cells. (A–C) BM-derived OPCs isolated from Nfe2l1(M)-KO and Flox mice were treated with RANKL (50 ng/ml) and M-CSF (30 ng/ml) for 7 days, followed by TRAP staining (A), counting of TRAP-positive multinucleated cells containing more than 3 nuclei (B), and measurement of mRNA expression of Cathepsin K, H⁺-Atpase and Atp6v0d2 (C). Scale bars: 200 µm. (D–G) Bone resorption assay and detection of F- ACTIN ring formation. The OPCs were treatment with RANKL (50 ng/ml) and M-CSF (30 ng/ml) for 7 days. The resorption pits on the dentine discs were visualized and quantified (D and E). The F- ACTIN ring formation was detected by phalloidin (green) and DAPI (blue) staining (F and G). Scale bar, 200 µm. Values are mean ± SD. n = 3, *p < 0.05 vs Flox. (H) mRNA expression of Nfe2l1 in RAW cells. n = 5. *p < 0.05 vs Scr. Scr, Scramble; KD, Nfe2l1-knockdown. (I) Representative image of immunoblots of NFE2L1 protein expression. Arsenite, the RAW cells were exposed to NFE2L1 activator sodium arsenite (10 µmol/L) for 6 h. (J–L) Raw cells were treated with RANKL (50 ng/ml) and M-CSF (30 ng/ml) for indicated times, followed by TRAP staining (J). Scale bars: 200 µm. (K) TRAP-positive multinucleated cells containing more than 3 nuclei were counted. n = 3. *p < 0.05 vs Scr with the same treatment; #p < 0.05 vs Scr of Day 2. (L) mRNA expression of Cathepsin K, H⁺-Atpase and Atp6v0d2 were analyzed. Values are mean ± SD. n = 3. *p < 0.05 vs Scr with the same treatment; #p < 0.05 vs Scr with Veh. (M–P) Bone resorption assay and F- ACTIN ring formation in RAW 264.7 cells. The cells were treatment with RANKL (50 ng/ml) and M-CSF (30 ng/ml) for 5 days, followed by measurements of resorption pits on the dentine discs (M and N, n = 4) and phalloidin (green) and DAPI (blue) double staining (O and P, n = 3). Scale bar, 200 µm. Values are mean ± SD. *p < 0.05 vs Scr. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
displayed similar cellular phenotypes as Nfe2l1 (M)-KO cells, showing reduced mRNA expression of antioxidant genes, increased ROS levels and augmented OD in response to RANKL stimulation and/or NAC treatment comparing Scramble cells. In addition, NAC treatment significantly decreased the protein levels of total Nfatc1 in Scramble and Nfe2l1-KD cells (Fig. 3O and P). Taken together, the findings above reveal that the augmentation of OD in Nfe2l1-deficient myeloid lineage cells is partially attributed to their reduced antioxidant expression and subsequent elevation in ROS accumulation in response to RANKL stimulation.

**NFE2L1 regulates the expression of Nfatc1/α and OD in an isoform-specific manner.** Using the primer sets targeting Nfatc1/α and Nfatc1/β (Supplementary Fig. 3B), we found that the OPGs from Nfe2l1 (M)-KO mice displayed elevated mRNA expression of Nfatc1/α, but not Nfatc1/β, following the initiation of OD induced by M-CSF and RANKL stimulation (Fig. 4A). In agreement with mRNA expression, immunoblotting assay revealed that the OPGs from Nfe2l1 (M)-KO mice have higher protein level of Nfatc1 than that in Flod control cells (Fig. 4B and C). Consistent with the findings in primary cultures, Nfe2l1-KD RAW cells also showed significantly elevated mRNA expression of Nfatc1/α (Fig. 4D) and protein levels of Nfatc1, which are mainly localized in the nuclear fractions (Fig. 4F), whereas no differences in mRNA levels of Rank, Csf1r and Traf6 and protein levels of TRAF6 were found (Supplementary Fig. 4).

To understand the regulatory roles of different isoforms of NFE2L1 in Nfatc1 expression, we developed multiple lines of RAW cells specifically overexpressing individual isoform of NFE2L1 or in a combination of L-NFE2L1 and S-NFE2L1. As shown in Fig. 4E, H, J and K, forced expression of NFE2L1-742, 741, 583 and/or 453 increased the protein levels of the corresponding isoforms in NFE2L1 in the cells. Overexpression of L-NFE2L1-742 or 741 substantially increased the mRNA and protein levels of Nfatc1/α. In contrast, overexpression of S-NFE2L1-453, but not 583, substantially reduced the mRNA and protein expression of Nfatc1/α (Fig. 4E, I, J and L). Moreover, overexpression of S-NFE2L1-453 dramatically blocked the expression of Nfatc1/α induced by overexpression of L-NFE2L1-742 or 741 (Fig. 4J-L).

To further validate the findings, we tried to deliver different isoform(s) of NFE2L1 back into Nfe2l1-KD RAW cells to rescue their phenotypes. As shown in Fig. 5A and B, Supplementary Figs. 6A, 6B, 7A, 7B, 8A and 8B, overexpression of NFE2L1-742, 741, 583 and/or 453 in Scramble and Nfe2l1-KD RAW cells substantially increased the protein levels of the corresponding isoforms of NFE2L1 in the cells. The protein levels of various isoforms of NFE2L1 overexpressed were generally slightly lower in the Nfe2l1-KD cells than those in Scramble RAW cells. In line with the findings in normal RAW cells, overexpression of L-NFE2L1-742 or 741 in Scramble cells substantially increased the mRNA and protein levels of Nfatc1 and subsequent OD, which showed elevated number of TRAP+ multinucleated cells and enhanced mRNA levels of Cathepsin K, H+-Atpase, Atp6v0d2 and Oscar (Fig. 5 and Supplementary Fig. 6). In contrast, overexpression of S-NFE2L1-453, but not 583, substantially reduced the mRNA and protein levels of Nfatc1 as well as OD (Fig. 5 and Supplementary Fig. 7). Moreover, overexpression of S-NFE2L1-453 even blocked the L-NFE2L1-induced expression of NFATc1 and augmented OD in Scramble cells (Fig. 5 and Supplementary Fig. 8). In Nfe2l1-KD RAW cells, overexpression of L-NFE2L1-742 or 741 further exacerbated the augmented expression of Nfatc1 at mRNA and protein levels and subsequent OD caused by Nfe2l1 silencing (Fig. 5 and Supplementary Fig. 6). However, silencing L-NFE2L1-742 or 741 had no effect on Nfatc1 expression at mRNA and protein levels (Supplementary Fig. 5). In contrast, overexpression of S-NFE2L1-453 in RAW cells reduced mRNA and protein expression of Nfatc1 and OD caused by either Nfe2l1 silencing or overexpression of L-NFE2L1-742 and 741 (Fig. 5 and Supplementary Fig. 8). Furthermore, the mRNA levels of antioxidant genes (Gclc, Gclm, Cat and Gsr) were detected in the cells overexpressing different isoforms of Nfe2l1. Forced expression of L-NFE2L1-742 or 741 had a tendency to increase the mRNA levels of the antioxidant genes. In contrast, overexpression of S-NFE2L1-453, substantially reduced the antioxidant gene expression under basal and L-NFE2L1-overexpressed conditions (Fig. 4M, 4N and 5E).

L-NFE2L1s and S-NFE2L1s oppositely modulate Nfatc1/α transcription. The transcription of Nfatc1/α is directed by two promoters, P1 and P2. The P1 spans approximately 800 bp, which are highly conserved between mouse and human, and contains a variety of binding sites and plays crucial roles in Nfatc1 induction in the process of OD. A CHIP assay showed that L-NFE2L1-742 and S-NFE2L1-453 may bind to the P1 (Fig. 6A and B). Based on the target sequences of CHIP-PCR, we found that L-NFE2L1-742 and S-NFE2L1-453 bind to the regions of −714 to −81 bp and −519 to −61 bp of P1, respectively. To further clarify the regulatory roles of different isoforms of NFE2L1 on the P1, we overexpressed L-NFE2L1-453 and S-NFE2L1 individually or in a combination with a P1-driven luciferase reporter. As shown in Fig. 6C, the activity of luciferase reporter driven by −1073 bp of P1 was significantly enhanced by overexpression of L-NFE2L1-742 or 742, whereas S-NFE2L1-583 had no effect on the reporter activity. In contrast, overexpression of S-NFE2L1-453 not only inhibited the reporter activity under basal condition, but also repressed the elevated reporter activity induced by overexpression of L-NFE2L1-742 or 742, suggesting that S-NFE2L1-453 may compete with L-NFE2L1s to suppress the transcription of Nfatc1/α.

To locate the binding site(s) of different isoforms of NFE2L1 on the P1 of Nfatc1/α, we developed a set of luciferase reporters driven by different lengths of P1 (Fig. 6D), and determined the effects of overexpression of L-NFE2L1-742, 741, 583 or 453 on their activities. As shown in Fig. 6E, L-NFE2L1-742 or 741 significantly increased the activity of luciferase reporters driven by different length of P1, with the highest induction in the reporters driven by −520, −265 and −111 bp of P1. Compared with the reporters driven by those short promoters, the basal and L-NFE2L1-742 or 741-stimulated activities of the reporters driven by −1073 or −711 bp of P1 were relatively low, revealing an enhancer and suppression site(s) may exist at the regions of −111 −1 bp and −711 −520 bp, respectively, where two CREB/Fos/ATF2 binding sites were described [43]. While S-NFE2L1-583 showed no significant effect on any of the reporters, S-NFE2L1-453 displayed dramatic suppression on all these reporters, suggesting that the major binding site(s) of S-NFE2L1-453 locates between −111 and −1 bp.

To verify whether the two CREB/Fos/ATF2 binding sites in P1 are
critical in controlling the transcription of *Nfatc1/α* by NFE2L1, we developed luciferase reporters driven by −711 bp of P1 with point mutation(s), which are critical for potential binding of NFE2L1 to the sites (Fig. 6F). As shown in Fig. 6G, the reporters with a T-to-C point mutation on the CREB/Fos/ATF2 binding site located between −711~−520 bp (Mut-1) displayed significantly elevated activities under the conditions of control and overexpression of various isoforms of NFE2L1. In contrast, an A-to-C point mutation on the CREB/Fos/ATF2 binding site located between −111~−1 bp (Mut-2) resulted in dramatic reduction in reporter activities. In addition, the reporters driven by the −711 bp of P1 with double point mutations (Mut-1/2) also showed decreased activities under conditions of control and overexpression of various isoforms of NFE2L1 (Fig. 6G). These findings reveal that the two CREB/Fos/ATF2 binding sites are critical in controlling the transcription

Fig. 4. NFE2L1 regulates *Nfatc1/α* expression in an isoform-specific manner. (A–C) Relative expression of NFATc1 in BM-derived OPCs of *Nfe2l1*(M)-KO mice. The cells were treated with RANKL (50 ng/ml) and M-CSF (30 ng/ml) for 1 day (A) or 2 days (B and C). mRNA (A) and protein expression (B and C) of NFATc1 were determined. Representative images (B) and quantitation (C) of NFATc1 immunoblots are shown. Flox, *Nfe2l1*LoxP/LoxP; KO, *Nfe2l1*(M)-KO; c1/α, *Nfatc1/α*; c1/β, *Nfat1/β*. (D) mRNA expression of *Nfatc1* isoforms in RAW 264.7 cells. Scr, Scramble; KD, *Nfe2l1*-knockdown. (F and G) Representative images (F) and quantitation (G) of NFATc1 immunoblots using total cell lysates and cell fractions of RAW 264.7 cells. Relative band intensity of total cell lysates and nuclear fractions was normalized to the β-ACTIN and LAMIN B, respectively. Values are mean ± SD. n = 3. *p < 0.05 vs Flox. (E and J) Immunoblots of NFE2L1 and NFATc1 in overexpression RAW 264.7 cells. Cont, 742, 741, 583, and 453 represent the cells overexpressed with control vector or different isoforms of *Nfe2l1*, individually or in combination. (H and K) Quantitation of NFE2L1 and NFATc1. Values are mean ± SD. n = 3. *p < 0.05 vs Control. **p < 0.05 vs 742 or 741 alone. (I, L, M and N) mRNA expression of *Nfatc1/α*, *Gclc*, *Gclm*, *Cat* and *Gsr* in the RAW 264.7 cells overexpressed with different isoforms of *Nfe2l1*. Values are mean ± SD. n = 3. *p < 0.05 vs Control.
Fig. 5. Effects of forced expression of different isoforms of NFE2L1 on NFATc1 expression and osteoclastogenesis in Scramble and Nfe2l1-KD RAW 264.7 cells. (A) Representative immunoblots of NFE2L1 and NFATc1 in Scramble and Nfe2l1-KD cells with overexpression of various isoforms of NFE2L1. Scr, Scramble; KD, Nfe2l1-knockdown; Cont, blank vector control; 742, 453, 742 + 453, cells were overexpressed with NFE2L1-742, 453 and 742 + 453, respectively. (B and D) Quantitation of NFE2L1 (B) and NFATc1 (D) immunoblots. (C and E) mRNA expression of Nfatc1/α, Gclc, Gclm, Cat and Gsr. (F) Representative images of TRAP staining. Scale bars: 200 μm. The cells were treated with RANKL (50 ng/ml) and M-CSF (30 ng/ml) for 5 days. (G) TRAP-positive multinucleated cells containing more than 3 nuclei were counted in the cells treated with RANKL (50 ng/ml) and M-CSF (30 ng/ml) for 5 days. (H) mRNA expression of Cathepsin K, H⁺-Atpase, Atp6v0d2 and Oscar in the cells treated as in (F). Values are mean ± SD. n = 3. *p < 0.05 vs Scr with the same overexpression; #p < 0.05 vs Cont of the same subtype of cells. $p < 0.05 vs the same cell subtype with 742 overexpression.
of Nfatc1/α, and NFE2L1 may bind to the CREB/Fos/ATF2 located between −111 ~ −1 bp sites to modulate Nfatc1/α transcription.

To further confirm the importance of the CREB/Fos/ATF2 binding site located between −111 ~ −1 bp of P1 promoter in the transcription of Nfatc1/α, we generated a luciferase reporter driven by the promoter with an A-to-C mutation on the CREB/Fos/ATF2 binding site (Fig. 6H). As shown in Fig. 6I, the reporter with an A-to-C mutation on the CREB/Fos/ATF2 binding site displayed significantly reduced activities under the conditions of control and overexpression of various isoforms of NFE2L1. In line with the findings using the reporter driven by −711 bp of P1, overexpression of L-NFE2L1-742 or 741 and S-NFE2L1-453 significantly enhanced and decreased the activities of luciferase reporter driven by wild type −111 bp of P1, respectively. In contrast, S-NFE2L1-583 had no significant effect on the reporter activity.

4. Discussion

We here demonstrate that ablation of all isoforms of Nfe2l1 specifically in the myeloid cell lineage of mice results in reduced bone mass and an increased number and activity of osteoclasts. A series of in vitro studies reveal that deficiency of Nfe2l1 results in accelerated OD and augmented osteoclast function. The increased OD resulting from Nfe2l1 deficiency is partially attributed to the increased sensitivity of ROS accumulation and induction of Nfatc1/α. Mechanistic studies show that lack of Nfe2l1 results in downregulation of multiple antioxidant genes and subsequent ROS accumulation in response to RANKL stimulation. Importantly, NFE2L1 was found to regulate the transcription of Nfatc1/α and OD in an isoform-specific manner; while L-NFE2L1s positively regulate the transcription of Nfatc1/α during OD, the S-NFE2L1-453
In the present study, we found, for the first time, addition, targeted deletion of all isoforms of Nfe2l1 and elevated osteoclast activity. These findings highlight that particular, physiological conditions, including estrogen deficiency and aging. In ing that NFE2L1 is involved in regulating osteoblast differentiation and lipogenesis, multiple physiological roles of NFE2L1 have been elucidated. NFE2L1 is ubiquitously expressed in a wide range of tissues including bone [44], hepatocytes [45], and pancreatic β-cells and adipocytes [15,30,31]. In bone metabolism, L-NFE2L1 has been found to interact with C/EBPβ to repress the expression of DSPP during odontoblast differentiation [19]. In response to ascorbic acid treatment, NFE2L1 regulates the transcription of multiple antioxidant genes and inflammation response and adiopogenesis [11,17,18,55-57]. Stable silencing of L-Nfe2l1 in RAW cells resulted in M1 polarization, indicating that L-NFE2L1 functions as a negative regulator of M1 polarization and pro-inflammatory response in macrophages [17]. In contrast, a S-NFE2L1 isoform, which migrates on SDS page generating a band at 65 kDa, was found to be a negative regulator of L-NFE2L1-mediated antioxidant response [52]. In the present study, we found that deletion of all isoforms of Nfe2l1 either in OPGs or in RAW cells resulted in an increase in expression of Nfatc1α/β at mRNA and protein levels and subsequently accelerated OD, indicating that at least one isoform of NFE2L1 plays a dominant negative role in the regulation. Given that silencing L-NFE2L1 had no significant effect on Nfatc1α/β expression and overexpression of S-NFE2L1-453 attenuated expression of antioxidant genes at mRNA and protein levels not only in Scramble control cells, but also in Nfe2l1-KD RAW cells, it is reasonable to conclude that the S-NFE2L1-453 may suppress the basal and inducible expression of Nfatc1α/β in L-NFE2L1-dependent and independent ways. With regard to the molecular mechanism underlying transcriptional regulation of Nfatc1α/β by NFE2L1, our in vitro data indicate that L-NFE2L1 and S-NFE2L1 may bind to Nfatc1α/β promoter to stimulate and suppress the transcription, respectively. In addition, L-NFE2L1 and S-NFE2L1 appear to compete for the same DNA binding site(s) to participate in the transcription of Nfatc1α/β. However, it is still unclear what the binding partners of L-NFE2L1 and S-NFE2L1 are and how they work together. We also do not know how NFE2L1 senses the stimulation and what are the signal transduction mediators relaying external osteoclastogenic stimuli to L-NFE2L1 and subsequently to S-NFE2L1 and how those distinct isoforms of NFE2L1 respond to RANKL–RANK signaling. In addition, one of the most intriguing questions is how L-NFE2L1 and S-NFE2L1 oppose each other to regulate the transcription of Nfatc1α/β. All the questions listed above need to be addressed in future studies to elucidate the molecular mechanisms of action.

To date, NFE2L1 has been found in a wide range of tissues and extensively investigated its roles in antioxidant response, inflammation, apoptosis, metabolic homeostasis, mitochondrial function, prostate, cellular differentiation and lipid metabolism [15–19,37]. However, its characteristics and functions in osteoclastogenesis and bone remodeling have not been reported. The current study found that L-NFE2L1 and S-NFE2L1 play essential roles in fine-tuning ROS accumulation and Nfatc1α/β transcription and thus regulates osteoclastogenesis and bone homeostasis. While L-NFE2L1 functions as an accelerator of osteoclast differentiation, in which multiple factors are coordinately involved, S-NFE2L1-453 serves as a dominant suppressor in the complex transcriptional process. The KO mice are deficient in all isoforms of Nfe2l1 transcripts specifically in the myeloid lineage and thus are totally absent of the control from L-NFE2L1 and S-NFE2L1-453 on the transcription of Nfatc1α/β and antioxidant genes. Because the withholding effect of S-NFE2L1-453 on Nfatc1α/β transcription is primary and dominant over the stimulation by L-NFE2L1 and other regulators, removal of all the isoforms of NFE2L1 results in a phenotype that Nfatc1α is upregulated, leading to elevated osteoclastogenesis.

Fig. 7. A schematic illustration of the isoform-specific regulatory roles of NFE2L1 in osteoclastogenesis. NFE2L1 plays essential roles in fine-tuning ROS accumulation and Nfatc1α/β transcription and regulation of osteoclastogenesis and bone homeostasis. In L-NFE2L1 functions as an accelerator of transcription of multiple antioxidant genes and Nfatc1α/β during osteoclast differentiation, in which multiple factors are coordinately involved, S-NFE2L1-453 serves as a dominant suppressor in the complex transcriptional process. The KO mice are deficient in all isoforms of Nfe2l1 transcripts specifically in the myeloid lineage and thus are totally absent of the control from L-NFE2L1 and S-NFE2L1-453 on the transcription of Nfatc1α/β and antioxidant genes. Because the withholding effect of S-NFE2L1-453 on Nfatc1α/β transcription is primary and dominant over the stimulation by L-NFE2L1 and other regulators, removal of all the isoforms of NFE2L1 results in a phenotype that Nfatc1α is upregulated, leading to elevated osteoclastogenesis.
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Appendix A. Supplementary data

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