Iron overload increases osteoclastogenesis and aggravates the effects of ovariectomy on bone mass

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Abstract

Postmenopausal osteoporosis is a metabolic disease associated with estrogen deficiency. The results of numerous studies have revealed the positive correlation between iron accumulation and postmenopausal osteoporotic status. Although the results of previous studies have indicated that estrogen or iron alone have an effect on bone metabolism, their combined effects are not well defined. Using an in vivo mouse model, we found that bone mass was minimally affected by an excess of iron in the presence of estrogen. Once the source of estrogen was removed (ovariectomy), iron accumulation significantly decreased bone mass. These effects were accompanied by fluctuations in the level of oxidative stress. To determine whether these effects were related to bone formation or bone resorption, primary osteoblasts (OBs), RAW264.7 cells, and bone-marrow-derived macrophages were used for in vitro experiments. We found that iron accumulation did inhibit the activity of OBs. However, estrogen had little effect on this inhibition. In contrast, iron promoted osteoclast differentiation through the production of reactive oxygen species. Estrogen, a powerful reactive oxygen scavenger, suppressed this effect in osteoclasts. Our data provided direct evidence that iron affected the bone mass only in the absence of estrogen. The inhibitory effect of estrogen on iron-induced osteopenia was particularly relevant to bone resorption rather than bone formation.

Key Words
- osteoporosis
- estrogen
- iron
- osteoclasts
- oxidative stress

Introduction

Primary type I osteoporosis, also known as postmenopausal osteoporosis (PMO), is a bone disease associated with reduced bone mineral density, disordered bone architecture and increased fragility (Chinese Orthopaedic Association 2009, Van den Bergh et al. 2012). Osteoporosis occurs when the bone mass decreases more rapidly than the body can replace it. Approximately 50% of women over 50 years of age suffer from osteoporosis, resulting in staggering financial costs for the USA (Ross 1996, Ray et al. 1997). Estrogen deficiency is regarded as the main causative factor in PMO. Withdrawal of estrogen or estrogen deficiency stimulates bone resorption by 90%, while increasing bone formation by 45%, as measured using biochemical markers (Yang et al. 2011).
This phenomenon, bone resorption outpacing bone formation, cannot be explained by estrogen deficiency alone (Huang et al. 2013). Therefore, we examined whether estrogen shortage is the sole factor in the development of PMO or if other risk factors are involved.

Iron overload has recently been linked to abnormalities in bone metabolism, including osteopenia, osteoporosis, and osteomalacia (Matsushima et al. 2001, Guggenbuhl et al. 2005). Osteoporosis occurs frequently in disorders associated with an excess of iron, such as thalassemia and hereditary hemochromatosis (Mahachoklertwattana et al. 2003, Vogiatzi et al. 2006). Most importantly, iron accumulation, a recently observed clinical phenomenon in postmenopausal women, might be involved in the pathogenesis of PMO (Kim et al. 2012). Menses is a critical pathway of iron excretion while menopause is a natural aging process that occurs as a woman passes from the reproductive to the non-reproductive years (Milman & Kirchhoff 1992, Riggs et al. 2002). Because of physiological amenorrhea, postmenopausal women experience an excess of iron. Although this iron status is considered to be within the normal physiological range, PMO has been linked to increased iron storage, a normal but not necessarily healthy process (Kim et al. 2013). Although there was no report of the relationship between iron and osteocytes, inhibition of osteoblastogenesis and stimulation of osteoclastogenesis caused by iron had been reported (Jia et al. 2012, He et al. 2013) and reactive oxygen species (ROS) were involved in iron-induced osteopenia (Li et al. 2012). ROS attenuate osteoblastogenesis, decrease osteoblast/osteocyte lifespan, and are required for osteoclast generation, function, and survival (Khosla et al. 2012). Estrogen has been shown to increase markers of oxidative stress in bone (Khosla et al. 2012). We propose the hypothesis that estrogen deficiency and increased iron levels are the main and secondary risk factors for PMO respectively. We speculate that iron accumulation aggravates PMO in the absence of estrogen. Furthermore, this deterioration would be suppressed if estrogen is restored using hormone replacement therapy.

In previous work, we established a male mouse model to observe the influence of iron on oxidative stress and bone resorption in vivo (Jia et al. 2012). In addition, we treated a murine macrophage cell line, RAW264.7 with ferric ions to test their activity (Jia et al. 2012). We found that iron promotes osteoclast differentiation and bone resorption. Using the fluorescent probe dichloro-dihydro-fluorescein diacetate (DCFH-DA), we examined oxidative stress as the underlying cause of iron-induced enhanced osteoclast function and increased osteoclast differentiation. Our results confirmed that these effects were regulated by the level of reactive oxidative species (ROS) (Li et al. 2012).

To examine the combined action of estrogen and iron on bone metabolism, especially the role of iron in PMO, we established an ovariectomized, iron-treated female mouse model. We performed experiments on primary osteoblasts (OBs), bone-marrow-derived macrophages (BMMs), and RAW264.7 cells to elucidate the mechanism underlying the development of PMO. We propose the hypothesis that the NF-κB signaling pathway may be involved in this development. This pathway is activated by virtually all stimuli that affect NF-κB including the receptor activator of NF-κB ligand (RANKL), the master osteoclastogenic cytokine and ROS (Ha et al. 2004, Lee et al. 2005). Free iron produces ROS through Fenton reactions, and estrogen is a well characterized antioxidant (Kim et al. 2006, Nakrst et al. 2011). Our analysis of the combined action of estrogen and iron may further our understanding of PMO pathogenesis.

Materials and methods

Animal experiments

All animal experiments were approved by the animal care committee of Soochow University. Two-month-old ICR female mice were divided into four groups (n=6/group). Two groups of mice were bilaterally ovariectomized and received intraperitoneal injections of normal saline (OVX) or 40 mg/kg ferric ammonium citrate (FAC) (F+OVX) three times a week for 2 months. The feed was decreased after ovariectomy for weight control. The Control (Con) and F groups, considered as sham operation groups, received identical normal saline and FAC treatments, but their ovaries were not excised. All mice were killed 2 days after the last treatment, and the body weight of all mice was measured.

Blood was collected from the dead mice and then centrifuged at 1000 g for 10 min in order to acquire the serum. Aliquots of serum were stored at −80 ºC for future analysis. BMMs were immediately extracted from one of the two femurs. The other femur from each mouse was scanned using micro-CT. The femurs and livers were then fixed in 10% buffered formalin at 4 ºC for Prussian blue staining.

Culture of OBs

Primary OBs derived from the calvaria of 1-day-old ICR mice were obtained as described previously (Suda et al. 1997). OBs were maintained in phenol-red-free α-MEM.
(Gibco) supplemented with 10% fetal bovine serum (FBS) (HyClone, Logan, UT, USA), 100 units/ml penicillin and 100 µg/ml streptomycin in a humidified atmosphere of 5% CO2 at 37 °C. The medium was replaced every 2 days. After reaching 70% confluence, the cells were detached using 0.05% trypsin and re-plated in either 55-cm² dishes or 12-well plates (3.8 cm² wells) at a density of 1×10⁴ cells/cm². The medium was replaced with serum-free medium containing 0 nM or 10 nM estradiol (E2) (Krum et al. 2009). Throughout the culture period, 25 µg/ml ascorbic acid, 10 mM sodium β-glycerolphosphate, and 100 nM dexamethasone were supplied to the cells.

**Culture of osteoclasts**

Murine RAW264.7 macrophage cells were plated in 55-cm² dishes or in 96-well plates at 3×10³ cells/well. RAW264.7 cells were cultured in phenol-red-free α-MEM/10% FBS containing 100 ng/ml RANKL (Peprotech, Rocky Hill, NJ, USA) for 4 days at 37 °C and 5% CO₂. Serum-free medium containing deionized water, E2, or FAC was supplied to cells during the last 2 days of culture. The method of intervention was the same as for the experiments on OBs.

BMMs were isolated from the femurs by flushing the bone marrow cavity with phenol-red-free α-MEM. Isolated cells were then centrifuged. The medium was replaced with ammonium-chloride-potassium buffer (pH 7.4, 0.1 mM NH₄Cl, 0.1 mM EDTA, and 1 mM K₂CO₃) for 30 s at room temperature to remove the red blood cells. Cells were cultured for 24 h in α-MEM/10%FBS. To generate osteoclast precursors, the non-adherent cells were collected and then cultured for 3 days in the presence of 30 ng/ml of macrophage colony-stimulating factor (M-CSF; PeproTech). The floating cells were discarded and adherent cells were considered to be BMMs. BMMs were cultured in α-MEM containing 30 ng/ml M-CSF and 50 ng/ml RANKL for 5 days.

**Cell proliferation assay**

Cell proliferation was analyzed using the Cell Counting kit 8 (Dojindo, Tokyo, Japan). RAW264.7 cells were detached and seeded into 96-well plates at a density of 3×10³ cells/well. After 24 h, cells were treated with FAC at various concentrations (12, 25, 50, 100, and 200 µM, 100 µl/well) or E₂ (1, 10, 10², 10³, and 10⁴ nM, 100 µl/well) and they were then incubated for another 24 h. The optical density (OD) of each group was measured at 450 nm using a BioTek microplate reader.

**Alkaline phosphatase, alizarin red staining and alkaline phosphatase activity**

OBs were treated with FAC and E₂ for 3 days and were then stained using alkaline phosphatase (ALP) following the manufacturer’s instructions (Beyotime, Shanghai, China). For alizarin red staining, cells were treated according to the experimental design. After 14 days, Cells were fixed and added with Alizarin Red-S solution (Beyotime) for 30 min. Then mineralized nodules were photographed. Cultured cells were seeded in 12-well plates and treated with FAC and E₂. After 10 days, cells were lysed with cell lysis buffer and centrifuged at 250 g for 5 min. Aliquots of supernatant were collected to measure ALP activity and protein concentration by using an ALP kit (Jiancheng, Nanjing, China) and a BCA protein assay kit (Beyotime) respectively. The OD was measured at a wavelength of 520 nm by using the BioTek microplate reader.

**Tartrate-resistant acid phosphatase staining**

To estimate the number of differentiated osteoclasts from bone marrow and RAW264.7 cells, they were stained with tartrate-resistant acid phosphatase (TRAP). Briefly, cells were fixed in 3.7% formaldehyde for 10 min and then incubated for 60 min at 37 °C in the dark with a solution containing sodium nitrite, Fast Garnet GBC, acetate, naphthol AS-BI phosphoric acid, and tartrate from the Leukocyte Acid Phosphatase Assay kit (Sigma) following the manufacturer’s instructions. Trap-positive multinucleated cells (MNCs) containing three or more nuclei were scored using light microscopy. The data are expressed as the mean ± S.D. of three samples.

**Resorption pit assay**

RAW264.7 cells were detached from the flask by using a trypsin/EDTA solution and were then resuspended as a single cell suspension. Cells were re-plated in Osteo Assay Surface 24-well plates (Corning, Corning, NY, USA) at a density of 2.5×10⁴ cells/well. α-MEM/10% FBS containing 100 ng/ml RANKL was added to each well. Plates were incubated at 37 °C in a humidified atmosphere of 5% CO₂ for 7 days. The serum-free medium was changed and reagents were added on day 4. The medium was aspirated from the well on day 7 and 400 µl of 10% bleach solution was added for 5 min. The wells were washed twice in distilled water. Individual pits were examined in dry wells by using a light microscope. The number of pits and the object area/total area were measured using Image-Pro Plus version 6.0 Software (Media Cybernetics, Warrendale, PA, USA).
Measurement of intracellular ROS level

OB and RAW264.7 cells were cultured and treated in 96-well plates. On the last day of culture, cells were incubated in the dark at 37 °C in serum-free medium containing 10 μM DCFH-DA (Beyotime) for 20 min. The cells were washed three times in serum-free medium to remove extracellular DCFH-DA. Cells were immediately examined using a fluorescent microscope. The OD was measured using a multi-detection reader at excitation and emission wavelengths of 488 and 525 nm respectively.

Micro-computed tomography analysis

The distal region of the left femur was subjected to three-dimensional micro-computed tomography (micro-CT) analysis. Bones were analyzed on a SkyScan 1172 high-resolution micro-CT scanner (SkyScan, Aartselaar, Belgium) using a 9-μm voxel size, 59 KvP, 127 μA, and 0.4° rotation step. The measured volumes of the cortical and trabecular regions of interest (ROIs) were obtained using a utility of the processing system. Trabecular ROIs extended from 50 μm proximally to the end of the distal growth plate over 1.7 mm toward the diaphysis. Cortical ROIs were 1.7 mm segments of the femoral middle-diaphysis. Cortical ROIs were defined by digitally subtracting the respective trabecular ROIs from the whole bone volume. All ROIs were drawn semiautomatically. Bone mineral density measurements were performed on cortical and trabecular volumes after segmentation of the bone voxels by using the global threshold, including only bone slices. Cone-beam CT Reconstruction Software version 2.6 was used to perform three-dimensional reconstructions and data analyses. The following trabecular and cortical parameters were analyzed: bone volume/total volume (BV/TV), trabecular number (Tb.N), trabecular separation (Tb.Sp), trabecular thickness (Tb.Th), trabecular bone pattern factor (Tb.Pf), structure model index (SMI), cortical volume, cortical surface, and cortical thickness.

Quantitative RT-PCR analysis

Total RNA was extracted from OB and RAW264.7 cells and was reverse-transcribed into cDNA by using a reverse transcription kit (Invitrogen) following the manufacturer’s instructions. The cDNA was mixed with SYBR Green Supermix (Bio-Rad) and primers for runt-related transcription factor 2 (RUNX2), transcription factor 7 (SP7), bone γ-carboxyglutamate protein (BGLAP), TRAP5, cathepsin K (CTSK), matrix metallopeptidase 9 (MMP9), or calcitonin receptor (CALCR) respectively. The primers used are listed in Table 1. Each experiment was performed in duplicate and the results were standardized to glyceraldehyde 3-phosphate dehydrogenase (GAPDH). All primers were purchased from Sangon (Shanghai, China). Data are expressed as the fold-change relative to the control.

Measurement of ferritin, oxidative stress, and markers of bone turnover in serum

Serum ferritin was measured using a mouse ferritin ELISA kit (Abnova, Taipei, China) according to the manufacturer’s protocol. Bone turnover markers including ALP, osteocalcin, C-terminal telopeptide of type 1 collagen (CTX), and TRAP5b were analyzed. ELISA kits for osteocalcin, CTX, and Trap5b were purchased from R&D, while the ALP kit has been previously described. Absorbance at 450 nm was measured for the contents of each well.

Table 1 Primers used for quantitative RT-PCR

<table>
<thead>
<tr>
<th>Gene</th>
<th>Accession no.</th>
<th>Primers (forward/reverse)</th>
</tr>
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<tbody>
<tr>
<td>RUNX2</td>
<td>NM_001145920</td>
<td>(F) 5'-AACCTCCTTGTCTCCGTCGTTGCTG-3' (R) 5'-TCCTGAACTTCGCTATCTGG-3'</td>
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<tr>
<td>SP7</td>
<td>NM_130458</td>
<td>(F) 5'-AGGAGCCAAAGAGAAGCCCTAC-3' (R) 5'-GATGCCGAGGTTGACACGAGC-3'</td>
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<tr>
<td>BGLAP</td>
<td>NM_007541</td>
<td>(F) 5'-GGACCATCTTCGCTACTCTG-3' (R) 5'-GTTCACACTTTATGCTCTCTG-3'</td>
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<tr>
<td>TRAP5</td>
<td>NM_001102405</td>
<td>(F) 5'-TACCTGGTGAGCAGATACC-3' (R) 5'-GATCCATAGTGAAACGCC-3'</td>
</tr>
<tr>
<td>CTSK</td>
<td>NM_007802</td>
<td>(F) 5'-TGTTAACCGCCAGGGCAA-3' (R) 5'-GGTTCAATATTACGTACGTCA-3'</td>
</tr>
<tr>
<td>MMP9</td>
<td>NM_013599</td>
<td>(F) 5'-TCCAGTAACACAGAAAG-3' (R) 5'-TTGACTGACCGTGG-3'</td>
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<tr>
<td>CALCR</td>
<td>NM_007588</td>
<td>(F) 5'-TCAGAACCAGGAAATCCTC-3 (R) 5'-ACATTCAAGCGATGCTTCT-3'</td>
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</table>

BGLAP, bone γ-gamma carboxyglutamate protein; CALCR, calcitonin receptor; CTSK, Cathepsin K; MMP9, matrix metalloproteidase 9; RT-PCR, real-time polymerase chain reaction; RUNX2, runt-related transcription factor 2; SP7, transcription factor SP7; TRAP5, tartrate-resistant acid phosphatase.
The plate reader was calibrated according to the manufacturer’s specifications. Oxidative stress markers, including malondialdehyde (MDA) and superoxide dismutase (SOD), were detected in the serum by using the MDA assay kit and SOD activity assay kit respectively. ROS will alter the level of SOD and MDA (Hohn et al. 2013). Both of the kits were purchased from Jiancheng. The ODs of MDA and SOD were measured at wavelengths of 532 and 450 nm respectively.

Western blot analysis

To analyze the level of NF-κB, whole-cell, cytoplasmic, and nuclear proteins were extracted. Nuclear proteins were extracted using the NucBuster Protein Extraction kit (Novagen, Rockland, MA, USA). Briefly, $2 \times 10^7$ RAW264.7 cells were lysed with 150 μl of reagent 1 to remove cytoplasmic proteins. The pellet was resuspended in 1 μl of 100× Protease Inhibitor Cocktail, 1 μl of 100 mM DTT, and 75 μl of reagent 2. The nuclear protein extracts were collected using centrifugation at 16 000 g for 5 min at 4 °C. A total of 40 μg of protein was mixed with 5× SDS–PAGE Sample loading buffer (Beyotime), boiled at 100 °C for 5 min. Proteins were transferred to PVDF membranes by electroblotting. The membranes were blocked using 5% non-fat dry milk in TRST for 1.5 h and then probed with anti-P50 (1:5000, Millipore, Billerica, MA, USA), anti-P65 (1:2000, Millipore), anti-IkBα (1:1000, Cell Signaling Technology, Danvers, MA, USA), anti-PIkBα (1:1000, Cell Signaling Technology), anti-histone 3 (1:5000, Millipore). Primary antibodies were detected using the DyLink TM800 Labeled Antibody to Rabbit/Mouse IgG (H+L) (1:10 000, KPL, Gaithersburg, MD, USA). Bound complexes were measured using the Odyssey Infrared Imaging System.

Statistical analysis

All the results are presented as the mean ± S.D. The differences between the experimental groups were evaluated using one-way ANOVA followed by Bonferroni post tests to allow for multiple comparisons. Statistical analyses were performed using the SPSS 19.0 Software. $P < 0.05$ was considered statistically significant.

Results

Osteopenia in mice is associated with iron accumulation and estrogen deficiency

As shown in Fig. 1A, ovariectomized mice fed the control diet showed little weight increase during the period of the experiment. FAC treatment (1.2 g/kg per week) over a 2-month period resulted in a marked iron accumulation in the liver, the largest ferritin reservoir. We stained the liver and bone sections by using Prussian blue, which indicated that iron was deposited in both the liver and bone marrow (Fig. 1B and C). Serum ferritin, a classic marker of iron storage, appeared to be significantly different between these groups ($F = 142.87, P < 0.001$). Ferritin increased in the F (335.30 ± 44.10 μg/l) and F+OVX (324.80 ± 38.60 μg/l) groups compared with the Con (39.40 ± 3.81 μg/l) and OVX (41.38 ± 5.65 μg/l) groups ($P < 0.001, P < 0.001$) (Fig. 1D).

To evaluate the effects of E2 and FAC on bone metabolism, micro-CT analyses of trabecular and cortical bone were performed. As expected, the excess iron in the F+OVX group (absence of estrogen) markedly reduced both trabecular and cortical bone compared with the OVX group, whereas the Con and F groups (presence of estrogen) showed no significant difference in these parameters except Tb.Sp (Table 2). No significant differences were observed for the Tb.N, SMI, and cortical indexes. These data indicated that estrogen played a key role in suppressing iron-induced bone loss. Three-dimensional reconstruction images are shown in Fig. 2A.

Oxidative stress was evaluated by measuring the levels of MDA and SOD, two commonly used indicators of ROS levels. We compared MDA and SOD in ovariectomized and sham-operated mice that had been administered FAC and normal saline. Differences were significant (MDA: $F = 19.62, P < 0.001$; SOD: $F = 22.73, P < 0.001$). FAC was shown to promote the generation of ROS in ovariectomized mice (MDA: 8.46 ± 1.91 versus 16.14 ± 2.42 nM, $P = 0.004$; SOD: 4.43 ± 0.81 versus 1.80 ± 0.67 U/ml, $P < 0.001$, OVX versus F+OVX), whereas the presence of estrogen suppressed oxidative stress (MDA: 6.22 ± 1.38 versus 8.14 ± 0.93 nM, $P = 0.182$; SOD: 5.83 ± 0.77 versus 4.73 ± 0.93 U/ml, $P = 0.081$, Con versus F) (Fig. 2B and C).

To further investigate bone turnover, sera were analyzed using ELISA kits. The levels of two factors involved in bone resorption, TRAP5b and CTX, were significantly different (TRAP5b: $F = 140.08, P < 0.001$; CTX: $F = 266.93, P < 0.001$). We observed higher levels of TRAP5b and CTX in the F+OVX group than in the OVX group (TRAP5b: 60.15 ± 3.49 versus 79.04 ± 6.47 U/l, $P < 0.001$; CTX: 9624 ± 1140 versus 14506 ± 915 pmol/l, $P < 0.001$). However, the presence of estrogen abrogated this effect. In accordance with the results of the micro-CT analysis, no significant differences were observed between the Con and F groups (TRAP5b: 23.27 ± 5.66 versus 29.51 ± 5.65 U/l, $P = 0.085$; CTX: 2588 ± 662 versus 3633 ± 125
449 pmol/l, Con versus F) (Fig. 2D and E). We also assessed ALP and osteocalcin levels in each group (ALP: F = 30.69, P < 0.001; osteocalcin: F = 69.93, P < 0.001). Iron reduced ALP and osteocalcin in both ovariectomized (ALP: 17.57 ± 2.09 versus 11.65 ± 2.02 U/100 ml, P < 0.001; osteocalcin: 98.27 ± 7.00 versus 83.06 ± 4.46 ng/ml, P = 0.002, OVX versus F + OVX) and sham-operated (ALP: 14.15 ± 1.22 versus 8.47 ± 1.30 U/100 ml, P < 0.001; osteocalcin: 73.33 ± 3.85 versus 57.75 ± 3.95 ng/ml, P < 0.001, Con versus F) mice, indicating that estrogen improved iron-induced osteopenia during the process of bone resorption, but not during bone formation (Fig. 2F and G).

Iron-induced defects in bone formation were not attenuated by estrogen

We investigated activation of varieties of deionized-water-treated and FAC-treated OBs in the presence of E₂ or vehicle. ALP staining and ALP activity assays indicated that ALP activity was significantly different (F = 23.63, P < 0.001) and the exposure of FAC-treated OBs to vehicle affected ALP activity (1.34 ± 0.08 versus 0.49 ± 0.17 U/gprot, deionized water-treated versus FAC-treated, P = 0.015), as had been reported previously (Yamasaki & Hagiwara 2009). To validate our in vivo findings, we added FAC to the culture
medium containing E2. In comparison with the cells treated with E2 alone, the cells co-incubated in FAC and E2 had lower ALP activity (1.49 ± 0.20 versus 0.76 ± 0.20 U/g prot, E2-treated versus FAC & E2-treated, P = 0.021) (Fig. 3A and B). Alizarin red staining provided us with a result that FAC inhibited OBs mineralization ability with or without estrogen. (Fig. 3C)

Quantitative RT-PCR analysis of RUNX2, SP7, and BGLAP mRNA levels confirmed the ALP analysis results, demonstrating that iron-induced changes in mRNA levels were not attenuated by E2 (RUNX2: F = 9.21, P = 0.029; SP7: F = 15.56, P = 0.011; BGLAP: F = 7.20, P = 0.043). FAC decreased RUNX2, SP7, and BGLAP expression, and the fold-decreases of these mRNAs, 0.24-, 0.69- and 0.72-fold

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Micro-CT analysis of mice trabecular and cortical bone (n = 6)a</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Con</td>
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<tr>
<td>Trabecular bone</td>
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<tr>
<td>BMD (mg/mm³)</td>
<td>0.19 ± 0.06</td>
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<tr>
<td>BV/TV (%)</td>
<td>20.37 ± 2.62</td>
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<tr>
<td>Tb.Th (µm)</td>
<td>95.94 ± 6.00</td>
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<tr>
<td>Tb.Sp (µm)</td>
<td>284.60 ± 9.74</td>
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<td>Tb.Pf (1/mm)</td>
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<td>Tb.N (1/mm)</td>
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<tr>
<td>SMI</td>
<td>1.840 ± 0.239</td>
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<td>Cortical bone</td>
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<tr>
<td>Volume (mm³)</td>
<td>1.135 ± 0.018</td>
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<tr>
<td>Surface (mm²)</td>
<td>12.102 ± 0.002</td>
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<tr>
<td>Thickness (mm)</td>
<td>0.194 ± 0.009</td>
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</table>

Con, control group; F: mice treated with FAC; OVX: mice treated with ovariotomy; F + OVX: mice treated with FAC and ovariotomy; BMD, bone mineral density; BV/TV, bone volume/total volume; Tb.Th, trabecular thickness; Tb.Sp, trabecular separation; Tb.Pf, trabecular bone pattern factor; Tb.N, trabecular number; SMI, structure model index.

*FAC markedly reduced trabecular bone mass and significantly affected other related parameters on the basis of ovariotomy. While the indices of mice without ovariotomy were rarely affected by iron.

bSignificant difference with OVX group (P < 0.05).

cSignificant difference with Con group (P < 0.05).

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Figure 2
Micro-CT three-dimensional reconstruction images and evaluation of oxidative stress, bone resorption, and bone formation markers in vivo. (A) Micro-CT images of mice belonging to the Con, F, OVX, and F + OVX groups. The trabecular bone of the distal femur and cortical bone of the mid-diaphysis femur are shown. (B and C) Levels of the oxidative stress markers superoxide dismutase (SOD) and malondialdehyde (MDA). (D and E) Serum levels of the bone resorption markers tartrate resistant acid phosphatase (Trap5b) and C-terminal telopeptide of type 1 collagen (CTX). (F and G) Serum levels of osteocalcin and alkaline phosphatase (ALP) were evaluated as markers of bone formation. The bar graphs show the means ± S.D. The asterisks (*) indicate significant differences at P < 0.05.
FAC and E2-treated, of OBs. Briefly speaking, though E2 attenuated the iron-varied from the results of the quantitative RT-PCR analysis (E2 - 0.050 respectively) (Fig. 3D, E and F). Therefore, our conclusions of the animal experiments. However, they
induced ROS level, iron-induced reduction in activation of OBs could not be improved by E2.

**Inhibitory effect of estrogen on osteoclast differentiation in the presence of iron**

Because the iron-induced defects in bone formation could not be attenuated by estrogen, we next explored the combined actions of estron and iron on bone resorption by evaluating the effects of FAC and E2 on osteoclast activation. Various concentrations of FAC (0, 12, 25, 50, 100, and 200 μM) and E2 (0, 1, 10, 10², 10³, and 10⁴ nM) were applied to RAW264.7 cells in the presence of 100 ng/ml of RANKL. As indicated by the number of Trap-positive MNCs, though higher concentrations of FAC and E2 inhibited proliferation of the cells (Fig. 4A and B), differentiation of RAW264.7 cells was stimulated with the indicated doses of FAC (50 μM) (P<0.001) and inhibited with the indicated doses of E2 (10 nM) (P=0.001) (Fig. 4C and D). These doses were used during the subsequent experiments.

The number of Trap-positive MNCs in cultures treated with both FAC and E2 was also scored (F=43.19, P<0.001). Preliminary results indicated that FAC could only promote osteoclast differentiation in the absence of

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E2 (37.80±7.22 versus 67.33±12.40, deionized water-treated versus FAC-treated, P<0.001; 17.33±6.71 versus 22.50±5.09, E2-treated versus FAC&E2-treated, P=0.299) (Fig. 5A and B), a finding consistent with the results of our resorption pit assay, including number of pits (F=316.50, P<0.001) and area ratio (F=61.13, P<0.001) (Fig. 5C, D and E). BMMs isolated from femurs were cultured and then Trap stained. The number of Trap-positive MNCs was significantly greater than that for the OVX group (12.67±1.75 versus 19.67±3.08, OVX versus F+OVX, P<0.001). However, the presence of estrogen partly abrogated iron-induced osteoclast differentiation (2.33±1.75 versus 3.17±1.47, Con versus F, P=0.501) (Fig. 5F and G). This result supports our hypothesis that estrogen ameliorates iron-induced bone loss during the process of bone resorption, but not bone formation.

We used the fluorescent probe DCFH-DA to determine whether iron or estrogen changed intracellular ROS levels. As expected, the results of this experiment were consistent with the results for detection of ROS in OBs, as shown in Fig. 6A and B (F=55.66, P<0.001). We examined whether iron and estrogen affected osteoclast differentiation by measuring TRAP, CTSK, MMP9, and CALCR expression. As shown in Fig. 6C, D, E and F (TRAP: F=7.12, P=0.044, CTSK: F=9.92, P=0.025; MMP9: F=16.00, P=0.011, CALCR: F=13.50, (Trap-positive cells) containing three or more nuclei were considered to be osteoclasts. The numbers of wine red cells were quantified. The bar graph shows the means±s.d. *P<0.05 indicates a statistically significant difference. A full colour version of this figure is available at http://dx.doi.org/10.1530/JOE-14-0657.

Postmenopausal osteoporosis, which is characterized by decreased bone mass and microarchitectural deterioration of bone tissue, represents an increasing medical and socioeconomic threat in the aged population worldwide.
In addition to estrogen deficiency, iron accumulation has recently been reported to be a secondary causative factor involved in PMO (Jia et al. 2012, Li et al. 2012, He et al. 2013, Kim et al. 2013). Postmenopausal women with abnormally high iron are reportedly more susceptible to osteoporosis (Kim et al. 2012, 2013). However, whether iron accumulation affects bone metabolism in premenopausal women has not, to our knowledge, been examined. This prompted us to ask whether an excess of iron leads to bone loss in the presence of estrogen.

Figure 5
E2 abrogated the stimulatory effect of FAC on osteoclast differentiation. (A and B) RAW264.7 cells were treated with FAC and E2 as indicated and Trap stained (200× magnification). Wine red Trap-positive cells were scored. (C, D and E) The effect of FAC and E2 on bone resorption. The eroded area and the number of pits were shown. (F and G) Bone-marrow-derived macrophages (BMMs) were cultured in the presence of 50 ng/ml RANKL and 30 ng/ml macrophage colony-stimulating factor (M-CSF) for 5 days and subjected to Trap staining to observe osteoclast formation (100× magnification). Wine red cells were considered to be mononucleated cells (MNCs). The bar graph shows the means ± S.D. *P < 0.05 indicates a statistically significant difference. A full colour version of this figure is available at http://dx.doi.org/10.1530/JOE-14-0657.
In this study, we generated an OVX mouse model with elevated iron to replicate the characteristic of postmenopausal females. Body weight had an influence on bone mass. The ratio of weight gain in OVX mice was slowed down with a control diet. Our mouse model had a pronounced bone phenotype, with defects similar to the OVX mouse model with normal levels of iron. Defects included trabecular and cortical thinning and alterations in the material properties of the bones. Considering the individual differences between mice and the tiny amounts of samples, levels of Tb.Sp in F group were increased and no significant difference was observed in Tb.N, SMI, cortical volume, surface, or thickness. In spite of these results, changes of other parameters were in accordance with theoretical analysis. In addition, a significant decrease in bone formation and resorption activities were observed in our model, recapitulating iron-induced osteopenia in ovariectomized mice. Similar results have been reported by Jia et al. (2012) and Tsay et al. (2010).

We also established an iron-accumulation mouse model that was not ovariectomized. In contrast to our OVX model, these mice experienced minimal bone loss, indicating a central role for the bioavailability of estrogen in attenuating iron-induced bone turnover. Interestingly, serum levels of bone formation markers were not consistent with the results of the micro-CT analysis, while those of bone resorption markers were consistent. Based on these findings, we investigated whether estrogen played a key role in protecting OB activity or affecting osteoclast differentiation in vitro. Expression of RUNX2 and SP7 decreased in iron-treated OBs in both the presence and absence of estrogen. In order to determine the dose of FAC and E2, we generated dose-response curves before performing experiments on osteoclasts. We found that FAC affected differentiation of osteoclasts at a concentration of 50 μM. The concentration for E2 was 10 nM. We also found that the differentiation of BMMs and RAW264.7 cells could not be stimulated using FAC when estrogen was present. Therefore, we proposed the hypothesis that estrogen protected mice from iron-induced osteopenia by influencing osteoclast differentiation, but not OB activity.

Estrogen is the predominant female steroid hormone and a pivotal regulator in bone formation and bone resorption processes (Chokalingam et al. 2012).
Menopausal women have been found to have increased ferritin levels, revealing a possible relationship between estrogen and iron (Jian et al. 2009, Ikeda et al. 2012, Yang et al. 2012). However, our results indicated that serum ferritin levels in the Con and OVX groups did not differ significantly. Thus, estrogen might not act directly on iron metabolism. As previously reported, ROS might be involved in the process of osteopenia induced by iron accumulation (Tsay et al. 2010). The mechanisms responsible, such as ROS regulation, should be further investigated. Though E2 eliminated iron-induced free radicals in OBs, FAC suppressed the activity of OBs in the presence of estrogen. This might indicate that iron-induced ROS are not a major factor mediating the inhibition of the activity of OBs.

Oxidative stress occurs in a cellular system when the production of free radical moieties exceeds the antioxidant capacity of that system. An increase in ROS levels may constitute a stress signal that stimulates specific redox-sensitive signaling pathways (Baek et al. 2010). As previously reported, the IκB/p-Tyr, Ras/mitogen-activated protein kinases/NF-κB or protein tyrosine kinase/phospholipase C/NF-κB pathways might be involved in ROS-induced NF-κB regulation (Trushin et al. 1999, Ye et al. 2000, Siomek 2012). Importantly, NF-κB controls the differentiation or activity of the major skeletal cell types, including osteoclasts, OBs, osteocytes, and chondrocytes (Alles et al. 2010, Novack 2011). NF-κB is a transcription factor that forms either homo- or heterodimers with various members of the Rel family, such as P50, P52, P65, cRel, and RelB. Together, they regulate the expression of numerous genes that are critical for the regulation of apoptosis, inflammation, viral replication, and various autoimmune diseases (Mankan et al. 2009, Hua et al. 2012). NF-κB, inhibited by its interaction with IκBa, IκBβ, IκBγ, and IκBe, is sequestered in the cytoplasm (Dyson & Komives 2012). Based on the results of this study, we speculate that iron-induced production of ROS leads to the phosphorylation of IκB and activates NF-κB (P50-P65 heterodimer), exposing a subunit of NF-κB (P50) and inducing nuclear translocation (Pereira & Oakley 2008). Nuclear P65, another subunit of NF-κB, binds to the consensus sequence of various genes, thus activating their transcription (Vermeulen et al. 2002). Therefore, the NF-κB pathway plays a fundamental role in the process of osteoclast differentiation (Soysa & Alles 2009, Novack 2011).

In summary, our data provided evidence supporting the conclusion that ROS, generated by iron, stimulated differentiation of osteoclasts, while estrogen inhibited iron-induced osteopenia by eliminating ROS in osteoclasts. Our results provided evidence indicative of a unique role of iron in pre- or post-menopausal osteoporosis. Drugs such as an iron chelator might be a potential treatment for PMO in the future. A weakness might be the size of the FAC dose, which induced a six- to sevenfold higher peak in iron levels than that of non-iron-treated mice. While in humans, the ratio is only two- to threefold (Huang et al. 2013). Species differences must be taken into account when considering the relationship between severity of iron excess and development of osteopenia. Non-canonical NF-κB signaling has not been considered. It will be examined in our follow-up experiment. A major strength of this study was that our mouse model reasonably recapitulated many features of postmenopausal women who have elevated iron levels. Furthermore, we used combined treatment with estrogen and iron to elucidate the effects of iron on bone metabolism.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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