Research Article

Auranofin Inhibits RANKL-Induced Osteoclastogenesis by Suppressing Inhibitors of \( \kappa B \) Kinase and Inflammasome-Mediated Interleukin-1\( \beta \) Secretion

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Osteoporosis is a degenerative metabolic disease caused by an imbalance between osteogenesis and osteoclastogenesis. Increased levels of proinflammatory cytokines combined with decreased estrogen levels, which are commonly seen in postmenopausal women, can lead to overactivation of osteoclasts. Therefore, targeting osteoclast maturation may represent a novel strategy for both treating and preventing osteoporosis. Auranofin is a gold-based compound first approved in 1985 for the treatment of rheumatic diseases. Here, we examined whether auranofin suppresses osteoclast differentiation in vitro and in vivo. Auranofin was shown to suppress receptor activator of NF-\( \kappa B \) ligand- (RANKL-) induced osteoclastogenesis in mouse bone marrow macrophages (BMMs) and Raw264.7 macrophages. Cotreatment of macrophages with auranofin blocked the RANKL-induced inhibitors of \( \kappa B \) kinase (IKK) phosphorylation, resulting in inhibition of nuclear translocation of p65. The pan-caspase inhibitor nivocasan potently reduced not only inflammasome-mediated interleukin-1\( \beta \) (IL-1\( \beta \)) secretion but also osteoclast differentiation in BMMs. Auranofin suppressed inflammasome activation, as evidenced by decreased production of cleaved IL-1\( \beta \) in both bone marrow-derived macrophages (BMDMs) and J774.A1 cells. Loss of both bone mass in ovariectomized mice was significantly recovered by oral administration of auranofin. Taken together, these data strongly support the use of auranofin for the prevention of osteoclast-related osteoporosis.

1. Introduction

Bone is a hard and rigid organ, though one that remains highly dynamic in nature, undergoing constant remodeling according to the balance between osteoblast and osteoclast activity. Osteoblasts make and fill the bone matrix, while osteoclasts absorb bone to aid new bone formation [1]. When the activity of osteoclasts exceeds that of osteoblasts, total bone mass in the body decreases, increasing the susceptibility to bone breaks in response to even small external impacts. Osteoporosis is a degenerative metabolic disease caused by an imbalance in skeletal remodeling [2]. Therapeutic agents capable of inhibiting osteoclast activity and differentiation have been proposed as a potential first-line treatment option for osteoporosis and may help increase existing bone mass.

In osteoclastogenesis, bone marrow cells migrate to the surface of the bone and differentiate into osteoclasts. Binding of receptor activator of nuclear factor-\( \kappa B \) (NF-\( \kappa B \)) ligand (RANKL) to receptor activator of NF-\( \kappa B \) (RANK) on the surface of osteoclasts plays a key role in this differentiation [3]. RANK signaling is triggered by cytoplasmic factors, leading to activation of at least five distinct signaling cascades, including inhibitor of NF-\( \kappa B \) kinase (IKK), c-Jun N-terminal kinase (JNK), p38 kinase, extracellular signal-regulated kinase (ERK), and Src, all of which are known to be activated in the course of osteoclast formation. Of those
signaling cascades, only the role of NF-κB is fully understood in osteoclast formation [3].

Denosumab is the first FDA-approved monoclonal antibody capable of directly targeting RANKL, resulting in decreased bone turnover and increased bone mineral density (BMD) [4]. Despite a higher annual treatment cost compared to other medications for osteoporosis, denosumab was found to be a cost-effective option in older osteoporotic male patients in the United States [5]. However, denosumab remains available only by injection, with no orally effective inhibitors of RANKL-dependent osteoclastogenesis yet available.

Auranofin is an orally available anti-rheumatoid arthritis (RA) agent composed of an elemental gold particle, Au(I), in complex with a sulfur-containing ligand. Administration of auranofin to RA patients has been shown to reduce levels of inflammatory cytokines, including interleukin-1 (IL-1) and tumor necrosis factor-α (TNF-α) [6]. Although the pharmacological mechanism underlying auranofin activity is not fully understood, the anti-inflammatory effects of auranofin are postulated to be related to inhibition of NF-κB [7]. Because NF-κB activity is a prerequisite not only for the transcription of proinflammatory genes but also differentiation of bone marrow cells to osteoclasts, we hypothesized that the anti-RA agent auranofin can prevent RANKL-induced osteoclast formation. Moreover, it has been reported that auranofin decreases osteoclastic bone resorption in bone slice assay [8]. In this study, we demonstrate that auranofin inhibits differentiation of macrophages to osteoclasts and prevents bone resorption both in vivo and in vitro. Our data further reveal that the antiosteoclastogenesis effect of auranofin is mediated through the inhibition of both IKK-mediated NF-κB activation and inflammasome-dependent IL-1β secretion in bone marrow-derived macrophages (BMDMs).

2. Materials and Methods

2.1. Materials. Auranofin (BML-E1206-0100) was purchased from Enzo Life Sciences (Lausen, Switzerland). Primary antibodies against the following proteins were used in this study: cathepsin K (E-7), NFATc1 (7A6), and p65 were from Santa Cruz Biotechnology (Dallas, TX, USA). Specific antibodies against lamin A/C, IKK, IκBα, and phospho-IκBα (Ser32) were from Cell Signaling Technology (Danvers, MA, USA), and the antibody against Iκkβ was purchased from Abcam (Cambridge, MA, USA). Tartrate-resistant acid phosphatase (TRAP) staining kit (386A-1KT) and 3-(4,5-dimethylthiazole-2-yl)-2,5-diphenyltetrazolium bromide (MTT) were from Sigma-Aldrich (St. Louis, MO, USA). Tartrate-resistant acid phosphatase (TRAP) staining kit (386A-1KT) and 3-(4,5-dimethylthiazole-2-yl)-2,5-diphenyltetrazolium bromide (MTT) were from Sigma-Aldrich (St. Louis, MO, USA). Bone resorption activities were measured by bone resorption assay kit (Cosmo Bio, Tokyo, Japan). To avoid the interruption of fluorescence measurement, phenol red free MEM (Thermo, Rochester, USA) containing 10% fetal bovine serum was added. BMMs were incubated in conditioned medium containing RANKL 200 ng/mL and M-CSF 30 ng/mL for 6 days without medium change. The fluorescence of medium was evaluated by a fluorometric plate reader (SpectraMax i3, Molecular Device, CA, USA). Pit area was measured after the plate was washed with 5% sodium hypochlorite (Sigma-Aldrich, St. Louis, MO, USA). Cells were counterstained with prolong gold antifade mountant with 4′,6-diamidino-2-phenylindole (DAPI, Thermo Scientific, Rochester, USA).

2.2. In Vitro Osteoclastogenesis Assay. Murine osteoclasts were prepared from bone marrow cells of C57BL/6 mice. Bone marrow cells were plated in petri dishes containing α-minimum essential medium, 10% fetal bovine serum with macrophage colony-stimulating factor (M-CSF; 30 ng/mL) for 3 days. Floating cells were removed, and those that became adherent were used as osteoclast precursor (BMMs). For generation of osteoclasts, the BMMs were seeded 10^6 cells/well in 6-well plates with M-CSF (30 ng/mL) and 100 ng/mL RANKL followed by medium change every 3 days. After incubation for 7 days, differentiated BMMs were fixed with 4% paraformaldehyde and stained for tartrate-resistant acid phosphatase (TRAP). TRAP-positive multinucleate cells (TRAP+MNCs) containing more than three nuclei were counted as osteoclasts.

2.3. Alkaline Phosphatase (ALP) Staining. The differentiated C2C12 cells were washed with phosphate-buffered saline (PBS) and fixed with 10% formaldehyde for 15 min. After being rinsed with PBS again, cells were stained with the BCIP/NBT color development substrate (Sigma-Aldrich, St. Louis, MO, USA) for 30 min at RT to evaluate ALP activity. The ALP-positive cells were stained as purple or blue color [9].

2.4. Pit Formation Assay. Bone resorption activities were measured by bone resorption assay kit (Cosmo Bio, Tokyo, Japan). To avoid the interruption of fluorescence measurement, phenol red free MEM (Thermo, Rochester, USA) containing 10% fetal bovine serum was added. BMMs were incubated in conditioned medium containing RANKL 200 ng/mL and M-CSF 30 ng/mL for 6 days without medium change. The fluorescence of medium was evaluated by a fluorometric plate reader (SpectraMax i3, Molecular Device, CA, USA). Pit area was measured after the plate was washed with 5% sodium hypochlorite (Sigma-Aldrich, St. Louis, MO, USA).

2.5. Immunocytochemistry. Immunocytochemistry was performed to assess the effects of auranofin on the nuclear translocation of p65 in the Raw264.7 macrophage cell line. The Raw264.7 cells treated with vehicle or 3 μM auranofin were fixed with 4% paraformaldehyde for 15 min, washed three times with 0.2% Triton X-100 in PBS for 10 min, blocked with 1% horse serum in PBS, and incubated with monoclonal anti-p65 antibody followed by Alexa-488 conjugated goat anti-mouse IgG antibody (A-21202, Invitrogen, Carlsbad, CA). Cells were counterstained with prolong gold antifade mountant with 4′,6-diamidino-2-phenylindole (DAPI, Thermo Scientific, Rochester, USA).

2.6. Animals and Experimental Design. 8-week-old ICR female mice were obtained from Orient Bio Co. (Sungnam, Korea). The mice were acclimatized for at least 2 weeks prior to use and were housed in an air-conditioned room with a 12 h light/dark cycle at a temperature of 21 ± 2°C and 50 ± 5% humidity. The mice were anesthetized with tribromoethanol (Avertin) and either sham-operated (SHAM, controls; n = 6), or surgically bilaterally ovariectomized (OVX; n = 12). Thirteen weeks after surgery, the OVX mice were divided into two groups with 6 mice per group as follows: OVX control group and auranofin group. 10 mg/kg body weight auranofin suspended in 60% polyethylene glycol (PEG) was orally administered for 43 days. Mice were sacrificed at the end of treatment, and blood samples were obtained for serum biochemistry analysis. The collected
distal femurs were fixed overnight in 75% ethanol prior to bone mass measurement [10]. The animal handlings and experimental procedures were performed in accordance with the regulations and rules of the Animal Ethics Committee of Sungkyunkwan University.

2.7. Serum Biochemistry Analysis. As a representative biochemical parameter relating to bone turnover, activity of bone alkaline phosphatase (BALP) was measured using commercial kit (MB703336, MyBioSource, San Diego, CA) in accordance with the manufacturer’s instructions.

2.8. Microcomputed Tomography Analysis. Bone morphometric parameters and microarchitectural properties were analyzed by microcomputed tomography (mCT) of the left femur using a SkyScan 1172 micro-CT scanner (Kontich, Belgium). μCT images were scanned by using a high-resolution SkyScan 1172 system (SkyScan, Kontich, Belgium) at 50 kV and 201 μA with a 0.5 mm aluminum filter and a resolution of 11 μm pixel⁻¹. Images were captured every 0.7° over an angular range of 180°. The bone mineral density (BMD), trabecular number (Tb.N), bone volume/total volume ratio (BV/TV), trabecular thickness (Tb.Th), trabecular pattern factor (Tb.Pf), and structure model index (SMI) were used to produce quantitative data in the SkyScan NRecon program and analyzed using SkyScan CTAN software.

2.9. Bone Histomorphometric Analysis. Collected tibias were embedded in paraformaldehyde and decalcified in 10% EDTA buffer for 2 weeks at 4°C. Samples were gradually dehydrated and embedded in paraffin and cut into 4 mm-thick longitudinal sections. After the sections were deparaffinized with xylene, they were stained with hematoxylin and eosin (H&E) [11].

2.10. Western Blot Analysis. Proteins were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), and the fractionated proteins were then transferred electrophoretically to nitrocellulose paper. Immune complexes were detected by using the Immobilon Western Chemiluminescent HRP Substrate (CB1001, Merck Millipore, Billerica, MA). Densitometric protein level quantities were determined by the image software, Multi Gauge, V3.0 (Fujifilm, Tokyo, Japan).

2.11. Measurement of Cell Viability. MTT assay was used to assess the cell viability. Briefly, 100 μL MTT (1 mg/mL diluted in phosphate-buffered saline) was added to each well of 96-well plates and the cells were incubated for 2 h. The culture media were removed by careful pipetting, and the resident cells were dissolved in 100 μL DMSO. The absorbance was measured at 490 nm with a TriStar microplate reader (Berthold Technologies, Bad Wildbad, Germany).

2.12. Reporter Gene Assay. Lipofectamine 2000 (Invitrogen, Carlsbad, CA) was used for transient transfection. Raw264.7 cells were plated on 48-well plates 1 day prior to transfection. 70% confluent Raw264.7 cells were transfected with reporter plasmids (1 μg pNF-κB-luciferase and 0.3 μg of pRL-SV40). The transfected cells were treated with 1 and 3 μM auranoïn for 24 h and then lysed with passive lysis buffer. Luciferase activities were determined by dual-luciferase reporter assay kit (E1960, Promega, Madison, MI), and firefly NF-κB promoter activity was normalized to the corresponding renilla activity.

2.13. RNA Preparation and Real-Time Quantitative Polymerase Chain Reaction (qPCR). Total RNA was isolated using TRIzol® reagent (Thermo Fisher Scientific, Waltham, MA, USA), and Oligo (dT) primers and reverse transcriptase (iNTRON Biotechnology, Seongnam, Korea) were used to synthesize cDNA. The mRNA expression levels of osteoclast markers, TRAP, and cathepsin K (CSTK) were quantified by real-time qPCR using a SYBR Select Master Mix (Applied Biosystems, Foster City, California) and Bio-Rad CFX Manager™ Software (Bio-Rad, Hercules, CA, USA). 18s rRNA protein mRNA was employed as a reference gene for normalizing mRNA levels. PCR was performed using selective primers for mouse TRAP (sense primer, 5′-GCAGTA TCTTCAG GAGGAGAC-3′; antisense primer, 5′-TCCA TAGTGAAACCGCAAGTAG-3′), CSTK (sense primer, 5′-CGAAAGAGGCTAGC GAACA-3′; antisense primer, 5′-TGGTAGCAGCAGAAACTTG-3′), and S18 ribosomal protein (S18r) genes (sense primer, 5′-TGAACC CGTTGAACCCCAT3′; antisense primer, 5′-CCATCC AATCGGTAGTAGCG-3′).

2.14. Enzyme-Linked Immunosorbent Assay (ELISA). Mouse interleukin-1β (IL-1β) ELISA kit (R&D Systems, Minneapolis, MN, USA) was used to measure concentration of IL-1β in culture media.

2.15. Statistical Analysis. Statistical significance was determined using a Student t-test, setting P < 0.05 as the level of significance. All experiments were carried out in triplicate at least.

3. Results

3.1. Inhibition of RANKL-Induced Osteoclast Differentiation by Auranoïn. Murine BMMs were used to test the antosteoclastogenic activity of auranoïn. BMMs were incubated with 100 ng/mL RANKL and 30 ng/mL M-CSF for 7 days, after which the cells differentiated into osteoclast-like giant and multinucleated tartrate-resistant acidic phosphatase-(TRAP-) positive cells (Figure 1(a)). The formation of TRAP-positive cells slowly diminished in BMMs incubated with 1 or 3 μM auranoïn (Figure 1(a)). Similar decreases in the number of multinucleated cells were also observed in response to auranoïn in a concentration-dependent manner (Figure 1(b)). To assess whether the antosteoclastogenesis leads to inhibition of bone resorption, BMMs were plated on a fluoresceinated calcium phosphate-coated plate and stimulated with M-CSF and RANKL in the presence or absence of auranoïn. BMMs stimulated with M-CSF and RANKL formed a number of resorption pits, while the area of pits was significantly reduced by 1 and 3 μM auranoïn (Figures 1(c) and 1(d)). Moreover, the fluorescence intensity
of medium resulting from RANKL-induced bone resorption activity was decreased by aurano
tin treatment (Figure 1(e)).

Those observations were further supported by real-time qPCR. Aurano
tin (1 and 3 μM) almost completely inhibited RANKL-induced mRNA expression of CSTK and TRAP, two osteoclast-selective markers (Figures 2(a) and 2(b)). RANKL-induced activation of nuclear factor of activated T-cells1 (NFATc1) is essential for achieving terminal differ-
entiation to osteoclasts [12]. To test the effects of RANKL on osteoclast differentiation, BMMs were exposed to RANKL with and without aurano
tin and screened for NFATc1 and CSTK protein expression. Treatment with 1 and 3 μM aurano
tin potently suppressed RANKL-mediated expression of both NFATc1 and CSTK (Figure 2(c)). Next, an MTT assay was carried out to exclude the possibility that the inhibition of osteoclastogenesis by aurano
tin is due to nonspecific cytotoxicity. No cytotoxicity was evident in BMDMs treated with 0.3–3 μM aurano
tin (Figure 2(d)). We further confirmed that aurano
tin (0.3–3 μM) inhibited NFATc1 expression in a concentration-dependent manner using RANKL-exposed Raw264.7 macrophages, a surrogate cell line of BMMs (Figure 2(e)). Treatment of Raw264.7 cells with 0.3–3 μM aurano
tin for 48 h marginally affected cell viability (data not shown).

3.2. Effect of Aurano
tin on RANKL-Activated IKK/IKBα.
Transcription of NFATc1 is mainly controlled by NF-κB [12]; however, the AP-1 family, including c-Fos, has been shown to trigger a transcriptional regulatory cascade by producing and cooperating with NFATc1. Furthermore, mice lacking c-Fos are osteoporotic because of a lack of osteoclast differentiation [13]. To investigate whether the inhibition of NF-κB activity by aurano
tin leads to blockade of osteoclast differentiation, we performed reporter gene analyses using a pNF-κB-luciferase reporter. In Raw264.7 cells transiently transfected with a luciferase reporter
construct containing NF-κB-binding regions, auranofin significantly inhibited RANKL-induced NF-κB luciferase reporter activity (Figure 3(a)). Although c-Fos plays a pivotal role in osteoclast differentiation, enhanced c-Fos expression by RANKL was only moderately affected by auranofin treatment in Raw264.7 cells, although significant decreases were observed in cells treated with the highest dose (3 μM) of auranofin (Figure 3(b)). As low concentrations of auranofin were found to effectively inhibit NF-κB activity and NFATc1 expression in osteoclast precursor cells, we next focused on how auranofin suppresses RANKL-induced NF-κB activation. Since p65 is a major transcription factor for NF-κB activation by RANKL, we examined nuclear translocation of p65 by subcellular fractionation with immunoblotting. Nuclear p65 protein levels increased from 5 to 15 min after treatment with RANKL (100 ng/mL), with 3 μM auranofin substantially inhibiting nuclear translocation of p65 at 15 min after RANKL treatment (Figure 3(c)). Immunocytochemistry using a p65 antibody confirmed that the majority of p65

**Figure 2:** Inhibition of RANKL-induced osteoclast differentiation by auranofin. (a, b) mRNA levels of CSTK and TRAP (n = 4). mRNA levels were determined by real-time qPCR. (c) The protein expression level of NFATc1 and CSTK (n = 3). (d) Cytotoxicity of auranofin in BMMs (n = 6). MTT assays were performed to evaluate cytotoxicity. (e) Effect of auranofin on the protein expression of NFATc1 in RANKL-treated Raw264.7 cells (n = 3). Data represents mean ± SD; *P < 0.05 and **P < 0.005 compared with the control group; #P < 0.05 and ###P < 0.005 compared with the RANKL-treated group.
Fig. 3: Continued.
was localized in the cytoplasm of quiescent Raw264.7 cells (Figure 3(d)). After stimulus with RANKL for 15 min, nuclear p65 was detected in some populations of Raw264.7 cells (Figure 3(d)); however, the nuclear translocation of p65 was completely blocked by 3 μM auranoﬁn (Figure 3(d)).

Next, we assessed the effects of auranoﬁn on upstream signaling pathways of NF-κB in RANKL-treated Raw264.7 cells. Like most members of the TNF receptor superfamily, RANK engages TNF receptor-associated factors (TRAFs) to activate several kinase cascades including IKK-dependent IxBα phosphorylation [14]. Under the control of TRAF6, transforming growth factor-β- (TGF-β-) activated kinase 1 (TAK1-) mediated phosphorylation of IKK plays a central role in activating the NF-κB pathway in response to inﬂammatory cytokines and RANKL [15]. Compared to Raw264.7 cells treated with RANKL only, coinubcation with 3 μM auranoﬁn led to signiﬁcant reductions in IKKβ phosphorylation (Figure 3(e)). TAK1-dependent IKK complex activation is followed by the phosphorylation and subsequent degradation of the IxBα subunit [16]. Immunoblot analyses using speciﬁc antibodies conﬁrmed that the phosphorylation and degradation of IxBα by RANKL were also prevented by 3 μM auranoﬁn (Figure 3(f)). These data strongly support the notion that NFATc1 downregulation by auranoﬁn may result from the inhibition of TAK1/IKK/IxBα axis-dependent NF-κB activation.

3.3. Inhibition of Inflammamosome-Mediated IL-1β Secretion by Auranofin. IL-1β, one of the major proinﬂammatory cytokines, is known to promote RANKL expression in marrow stromal cells and directly induces osteoclastogenesis [17]. IL-1β is expressed by hematopoietic cells as a 31 kDa precursor, primarily in response to inﬂammatory stimuli. However, the full-length pro-IL-1β is inactive and must be cleaved into its mature 17 kDa form before binding to the IL-1 receptor [18]. In BMDMs, IL-1β secretion is thought to occur via a two-step process. The ﬁrst signal can be triggered by various pathogen-associated molecular patterns (PAMPs) after Toll-like receptor (TLR) activation, resulting in NF-κB-dependent synthesis of NLRP3 and pro-IL-1β. The second signal is mediated by activation of the inﬂammamosome, resulting in cleavage of IL-1β by caspase-1 [19].

To test whether inﬂammamosome-dependent IL-1β secretion is involved in the differentiation of BMMs to osteoclasts, we examined the effect of nivocasan (GS-9450), a pan-caspase inhibitor, on osteoclastogenesis. Nivocasan irreversibly arrests caspase-1 activity and inhibits that of caspase-8, both of which play a key role in IL-1β secretion [20]. BMDMs were exposed to lipopolysaccharide (LPS) and ATP to induce NLRP3 inﬂammamosome-dependent IL-1β secretion. As shown in Figure 4(a), 10 μM nivocasan treatment inhibited cleavage of pro-IL-1β into the active p17 fragment. The inhibition of IL-1β cleavage by nivocasan resulted in blockade of osteoclast differentiation, as evidenced by a decrease in the number of TRAP-positive and multinucleated cells (Figure 4(b)). Furthermore, protein expression of NFATc1 and CSTK was suppressed by nivocasan (Figure 4(c)). These results indicate that inﬂammamosome signaling is actively involved in the differentiation of BMMs to osteoclasts.

As auranoﬁn inhibits IL-1β secretion in the differentiated cells from BMMs [21], we hypothesized that auranoﬁn may block osteoclast differentiation by interfering with inﬂammamosome-mediated IL-1β secretion in BMDMs. In a manner similar to that of nivocasan, auranoﬁn impairs inﬂammamosome mediated IL-1β secretion in BMDM and J774.A1 cells (ASC-positive mouse monocyte/macrophage cell line) (Figure 4(d)). Western blot analyses conﬁrmed that 0.3–3 μM auranoﬁn potently reduced the amount of cleaved IL-1β in both BMDM and J774.A1 cells exposed to LPS and ATP (Figure 4(e)). Interestingly, the inhibitory effects of auranoﬁn on inﬂammamosome activation were clearly observed at very low concentrations (0.3 and 1 μM), suggesting that the antiosteoclastogenesis effect of auranoﬁn is partly dependent on its ability to inhibit inﬂammamosome activation.

3.4. Inhibition of Osteoporosis by Auranofin in Ovariectomized Mice. Finally, we investigated the effects of
Figure 4: Inhibition of inflammasome-mediated IL-1β secretion by auranoﬁn. (a) Effects of auranoﬁn on the protein expression of pro-IL-1β and cleaved IL-1β. BMMs were primed with LPS (100 ng/mL) for 4 h and further incubated with 1 mM ATP and nivocasan (3-30 μM) for 1 h. (b) Effect of the inflammasome inhibitor (nivocasan) on osteoclast differentiation. The left panel shows the representative images of TRAP stainings, and the number of TRAP-positive multinuclear cells was illustrated in the right panel (n = 3). (c) Effects of nivocasan on NFATc1 and c-Fos expression in RANKL-treated BMMs. (d) IL-1β ELISA results (n = 3). The secreted IL-1β levels were determined in culture media from J774.1A cells and BMDMs incubated with 1 mM ATP and auranoﬁn (0.3-3 μM) for 1 h followed by preincubation with 100 ng/mL LPS for 4 h. (e) Cell lysates from J774.1A cells (left) and BMDMs (right) were analyzed for protein expression of pro-IL-1β and cleaved IL-1β by immunoblottings. ***P < 0.005 compared with the control group; ###P < 0.005 compared with the RANKL-treated group.
auranofin on bone loss in vivo using an ovariectomized (OVX) mouse model of osteoporosis. OVX mice were orally administered with vehicle or 10 mg/kg auranofin for 43 days. The body weight of mice was measured daily for 43 days, with a final reading performed just before sacrifice. Body weights of the OVX and control groups treated with 10 mg/kg auranofin were virtually identical prior to initiation of the study. However, the total body weight of the auranofin-treated mice gradually decreased over the course of the study, supporting a previous finding that the major gastrointestinal side effect of auranofin is diarrhea [22] (Figure 5(a)).

As evidenced by representative H&E stains, auranofin administration inhibited trabecular bone loss in OVX mice (Figure 5(b)). Auranofin treatment almost completely restored weight loss in the left or right femur of OVX mice after 43 days (Figure 5(c)), whereas serum levels of bone alkaline phosphatase (BALP) in OVX mice were significantly higher than those of the sham-operated animals. BALP is released in the circulation during the bone-forming phase of the remodeling process [23]. Compared to OVX mice, auranofin-administered mice showed decreased levels of serum BALP (Figure 5(d)).

Results from these experiments were further analyzed by microcomputed tomography (micro-CT). Although calibrated BMD marked no significant decrease in OVX mice (data not shown), representative pictures of micro-CT sections showed improvement in bone density in auranofin-administered mice (Figure 6(a)). Auranofin suppressed the two- and three-dimensional structural bone loss in the OVX mice, as evidenced by changes in BMD, percent bone volume, trabecular thickness, and number (Figures 6(b) and 6(e)–6(g)). Although trabecular pattern factors and structural model indices showed no changes, the trabecular separation number was much lower in auranofin-administered mice, demonstrating the utility of auranofin for prevention of bone loss.

4. Discussion

Drug repositioning is an increasingly popular strategy in drug development, as it has the potential to both speed up the discovery of novel treatments and reduce the risk of drug failure. Many pharmaceutical companies are scanning the existing pharmacopoeia for repositioning candidates and the number of repositioning success stories is increasing [24]. Auranofin, which is clinically approved for use in RA therapy, is one such candidate, with this agent having been investigated for potential therapeutic application in a wide range of diseases, including cancer [25, 26].

Figure 5: Inhibitory effect of auranofin on bone loss in ovariectomized mice. (a) Body weight change (n = 6 for each group). (b) H&E staining of the vertical and horizontal sections of femurs. (c) Weight of left, right, and total femur relative to body weight (n = 6) (d) Serum levels of BALP (n = 4). *P < 0.05 compared with the control group; #P < 0.05 and ##P < 0.01 compared with the RANKL-treated group.
Gold-based compounds, such as aurano
din, are known
to exhibit anti-in
flammatory activity via the downregula-
tion of NF-κB [27] and may also exert e
ffects on upstream
targets of IKK-dependent IκBα
phosphorylation [28]. Based
on previous studies demonstrating an essential role of NF-
κB activation in osteoclastogenesis [3, 29], we hypothesized that
aurano
din may block osteoclast di
fferentiation. Here, we
showed that aurano
din treatment led to reduced di
fferentiation of bone marrow cells to osteoclasts by inhibiting
RANKL-induced osteoclast di
fferentiation in Raw264.7 cells
and BMMs. Under normal circumstances, osteoclast di
fferentiation is mediated by the binding of RANKL to its target
receptor, activating multiple intracellular signaling pathways
including IκB kinase and NF-κB [14]. The RANKL-induced
phosphorylation of IKK/IκBα was suppressed by aurano
din, followed by reduced NF-κB activation.

Aurano
din also inhibited secretion of IL-1β from
BMDMs. In an in vitro osteoclastogenesis assay using a
pan-caspase inhibitor, we observed a close correlation
between IL-1β secretion and osteoclast differentiation. Mature IL-1β is secreted via in
flammasome-mediated activation and is responsible for osteoclast-induced bone resorp-
tion [17]. IL-1 was shown to directly induce osteoclast di
fferentiation in BMMs expressing the IL-1 receptor via a
RANKL-independent pathway [21]. In this study, we dem-
onstrated that aurano
din dramatically blocks IL-1β secretion
from BMDMs. The antosteoclastogenic activity of aurano
din was further confirmed in vivo using an OVX mouse model
of osteoporosis. Oral administration of 10 mg/kg aurano
din led to significant recovery of weight loss in the femur of
OVX mice. That inhibitory effect is thought to be closely
correlated with the antiosteoclastogenic activity of aurano
din.
as it showed no effect on bone morphogenetic protein-(BMP-) induced osteoblastic differentiation of C2C12 cells (Supplementary Figure 1). Serum BALP levels, which are indicative of bone turnover activity, were also reduced in OVX mice treated with auranoﬁn. Furthermore, data from our micro-CT analyses strongly supported the results of in vitro assays examining anti-bone resorptive activity.

Taken together, the in vitro and in vivo results presented here clearly support the use of auranoﬁn as a potent inhibitor of BMM diﬀerentiation to osteoclasts. Denosumab is the first drug targeting osteoclast differentiation and has been approved for the treatment of malignant osteoporosis in both the United States and Europe. Despite its high eﬃcacy and low rate of adverse events in clinical trials [30], its high cost and inconvenient parenteral injection led to continued interest in alternative therapeutic options. Here, we propose auranoﬁn as a potential substitute for denosumab, as it is cost-eﬀective and orally available. Moreover, auranoﬁn has undergone extensive toxicological analyses and has proven safe for human use.

RANKL, through its ability to stimulate osteoclast formation and activity, is a critical mediator of bone resorption and overall bone density. Overproduction of RANKL is implicated in a variety of degenerative bone diseases, including RA and psoriatic arthritis. In RA, RANKL is expressed by both T cells and synoviocytes, leading to persistent osteoclast activation and bone erosion [31]. The data presented here further suggest that auranoﬁn exerts anti-RA activity via targeting of RANKL-induced osteoclast maturation.

5. Conclusion

The current results suggested that auranoﬁn could serve as a novel inhibitor of osteoclastogenesis by suppressing IKK-IxBα phosphorylation and inﬂammasome-mediated IL-1β secretion. Based on these ﬁndings, auranoﬁn represents an excellent candidate for drug repurposing, offering potential therapeutic eﬃcacy for osteoclast-related disorders outside of its approved use in RA.

Data Availability

The data used to support the ﬁndings of this study are available from the corresponding author upon request.

Conflicts 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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Supplementary Materials

Supplementary Figure 1: alkaline phosphatase (ALP) staining. Osteoblast differentiation of C2C12 cells was induced by bone morphogenetic protein (BMP). ALP staining was performed to evaluate the eﬀect of auranoﬁn on the differentiation capacity. (Supplementary Materials)

References

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