Chronic Ethanol Consumption Induces Osteopenia via Activation of Osteoblast Necroptosis

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Chronic high-dose alcohol consumption impairs bone remodeling, reduces bone mass, and increases the risk of osteoporosis and bone fracture. However, the mechanisms underlying alcohol-induced osteoporosis are yet to be elucidated. In this study, we showed that excess intake of ethyl alcohol (EtOH) resulted in osteopenia and osteoblast necroptosis in mice that led to necrotic lesions and reduced osteogenic differentiation in bone marrow mesenchymal stem cells (BMMSCs). We found that EtOH treatment led to the activation of the RIPK1/RIPK3/MLKL signaling, resulting in increased osteoblast necroptosis and decreased osteogenic differentiation and bone formation both in vivo and in vitro. We further discovered that excessive EtOH treatment-induced osteoblast necroptosis might partly depend on reactive oxygen species (ROS) generation; concomitantly, ROS contributed to necroptosis of osteoblasts through a positive feedback loop involving RIPK1/RIPK3. In addition, blocking of the RIPK1/RIPK3/MLKL signaling by necrostatin-1 (Nec-1), a key inhibitor of RIPK1 kinase in the necroptosis pathway, or antioxidant N-acetylcysteine (NAC), an inhibitor of ROS, could decrease the activation of osteoblast necroptosis and ameliorate alcohol-induced osteopenia both in vivo and in vitro. Collectively, we demonstrated that chronic high-dose alcohol consumption induced osteopenia via osteoblast necroptosis and revealed that RIPK1 kinase may be a therapeutic target for alcohol-induced osteopenia.

1. Introduction

Chronic and excessive alcohol consumption has harmful effects on multiple organs, including the brain, heart, liver, and skeleton [1–4]. Chronic heavy alcohol consumption is recognized as a major cause of secondary osteoporosis and is positively associated with impaired bone remodeling, extensive bone loss, and fracture [5–7]. Multiple mechanisms are responsible for alcohol-induced osteoporosis (AOP), including a direct effect on bone formation and bone resorption [8, 9]. Ethyl alcohol (EtOH) intake inhibits bone marrow mesenchymal stem cell (BMMSC) derived osteoblast proliferation and enhances osteoclast differentiation via the RANKL/RANKL/OPG signaling pathway that promotes bone resorption in long-term alcohol users [10, 11]. The imbalanced osteo/adipogenic lineage commitment of BMMSCs is modulated via activation of the PI3K/AKT/mTOR signaling that results in downregulation of runt-
related transcription factor 2 (Runx2) and upregulation of peroxisome proliferator-activated receptor γ (PPARγ) [12–14]. The indirect action of chronic alcohol-induced bone loss may also be regulated by inducing oxidative stress [13, 15]. It has been reported that with alcohol exposure, increased levels of reactive oxygen species (ROS) may directly precede the production of TNF-α and other inflammatory cytokines in the bone marrow and contribute to bone formation decrease [16]. Furthermore, increased endothelial nitric oxide synthase (eNOS) and overexpressed inducible nitric oxide synthase (iNOS) in the bone may contribute to osteocyte apoptosis and necrosis [17].

Apoptosis was identified as the early stage in alcohol-mediated osteoporosis, specifically an increase in osteoblast apoptosis and osteocytes [18, 19], while necrosis occurs in succession following extensive apoptosis. Deficiency or inhibition of the aspartate-specific cysteine protease-8 (caspase-8) can switch cell death into a specific form of necrosis, resulting in necroptosis [20–22]. Necroptosis, having typical morphological characteristics of necrosis, can be induced by tumor necrosis factor-α (TNF-α), a factor associated with suicide (Fas, which binds to the ligand CD95L, known as FASL), TNF-related apoptosis-inducing ligand (TRAIL), and other death cytokines [23–25]. Receptor-interacting serine/threonine protein kinase 1 (RIPK1) and 3 (RIPK3) activate the mixed lineage kinase domain-like protein (MLKL), which then recruits the necosome to localize on the cell membrane and causes cell necroptosis [26, 27]. Furthermore, necroptosis can be regulated and pharmacologically inhibited by necrostatin-1 (Nec-1), the first small-molecule inhibitor of RIPK1 kinase to be developed [28–30]. Nec-1 is a cellular protector and has been widely used to investigate the role of RIPK1 in multiple disease models, including neurodegenerative and cerebrovascular diseases [31–33], atherosclerosis [34], heart attack [35, 36], and liver, retinal, and renal injury [37, 38]. Nec-1 has been reported to inhibit the death of chondrocytes in cartilage injury and promote cartilage repair [39]. Treatment with Nec-1 also reduces osteocyte necrosis and increases bone formation in glucocorticoid-induced osteoporosis (GIOP) rats and ovarioctomized rats that resulted in the kinase RIPK1-dependent necroptosis [40, 41]. In addition, the necroptosis signaling pathway is also found to be involved in the pathophysiology of osteosarcoma [42]. However, it remains unknown whether RIPK1 mediates damage via activation of necroptosis in an alcohol-induced osteopenia model and the effect of necrostatin-1 on osteopenia requires further elucidation.

In the current study, we investigated the effect of ethanol treatment on osteogenesis and osteopenia and the underlying mechanisms that mediate necroptosis in alcohol-induced osteopenia. Our investigations revealed that EtOH treatment increased osteoblast necroptosis and decreased osteogenic differentiation and bone formation in vivo; ROS have been considered, at least in part, as a critical driving force for necroptosis; necrostatin-1 and N-acetylcycteine (NAC) ameliorated osteopenia via inhibition of the RIPK1/RIPK3/MLKL signaling pathways. Therefore, RIPK1 kinase and targeted antioxidants may be therapeutic targets for alcohol-induced osteopenia.

2. Materials and Methods

2.1. Animals. All animal experimental protocols were approved by the Animal Research Center of Southwest Medical University (No.: 20210928-007) and performed in accordance with the policies of Chinese animal research committees. Male C57BL/6J mice at 8 weeks were purchased from the SPF (Beijing) Biotechnology Co., Ltd. (China, SCXK 2016-0002) and acclimatized for two weeks with temperature and humidity controlled at 21 ± 0.5°C and 55 ± 10%, respectively. Then, mice were randomly divided into a control vehicle-treated (control), necrostatin-1 drug control group (Nec-1), chronic high-dose alcohol group (EtOH), and chronic high-dose alcohol add on necrostatin-1 treatment group (EtOH+Nec-1) (n = 15/group). Mice were administered 0.2 ml/10 g/d of alcohol (gradually increase concentration to 30% v/v) via gavage [43, 44] and 1.65 mg/kg/d of necrostatin-1 (Sigma-Aldrich, St. Louis, USA) by intraperitoneal injection (ip) [40]; the control received the ddH2O and 1000 μl/kg/d of 1% DMSO solution (Sigma-Aldrich, St. Louis, USA) in the same volume by the same route for 22 weeks.

2.2. Dual-Energy X-Ray Absorptiometry (DXA). Mice were anesthetized with 1% pentobarbital sodium (50 mg/kg body weight, ip). The whole body, lumbar spine, and bilateral femur scans were analyzed using the region of interest (ROI) tools (GE Healthcare Lunar, Germany). Bone mineral density (BMD) was automatically calculated for each ROI scan. A quality assurance scan of the calibration block provided with the scanner was performed daily.

2.3. Micro-Computed Tomography Assay (Micro-CT). The distal femurs were harvested, fixed in 4% paraformaldehyde (PFA; Sigma-Aldrich, St. Louis, USA), and placed in fresh 70% ethanol. The specimens were then scanned using a high-resolution μCT scanner (SCANCO Medical AG, μCT 50, Brüttiselen, Switzerland) with a voxel size of 15 μm, an X-ray tube voltage of 70 kVp, a current of 200 μA, and an integration time of 300 ms. The scanner special software (SCANCO Medical Evaluation) was used to locate and analyze the distal femurs. The volumes-of-interest (VOIs) of bone tissue were 29 × 29 × 29 μm3, and a total of 1800–2000 layers of cancellous bone tissue were defined as the final ROI. The datasets were loaded onto the Amira software with 345 analysis thresholds.

2.4. Histological and Immunofluorescent and Immunoistochemistry Staining. The femurs were fixed in 4% PFA for 24 h, decalcified with 10% ethylenediaminetetraacetic acid (EDTA, pH 7.6–7.8) for 40 days at room temperature, and embedded in paraffin. Sections of 2 μm thickness were deparaffinized, dehydrated, and stained with hematoxylin and eosin (H&E) and TRACP staining for histological and
histomorphometric analysis (Leica, Germany). Bone remodeling activity was evaluated by the ratio of osteoid surface to bone surface (OS/BS, %), osteoblast surface to bone surface (Ob.S/BS, %), and eroded surface to bone surface (ES/BS, %), according to the recommendations of the ASBMR Histomorphometry Nomenclature Committee [45]. Sections of 4 μm thickness were subjected to immunofluorescent and immunohistochemistry staining, blocked with 10% goat serum (Gibco, USA), and incubated with Runx2- or RIPK1- or RIPK3- or cleaved caspase-3-specific antibodies (1:100; Abcam, Cambridge, UK) at 4°C overnight, followed by a fluorescent secondary antibody (1:200; Cell Signaling Technology, USA) for immunofluorescent staining; treated with secondary antibody, incubated with the streptavidin-horseradish peroxidase complex, visualized with diaminobenzidine, and counterstained with hematoxylin for immunohistochemistry staining. The trabecular bone structure parameters and area percentage and the percentage of Runx2-, RIPK1-, and RIPK3-positive cells were calculated using the Image-Pro Plus software.

2.5. Isolation and Culturing of Mouse BMMSCs. Mouse BMMSCs were isolated from the bilateral femoral marrow of six-week-old male C57BL/6J mice. Single-cell suspensions were cultured in alpha-minimum essential medium (α-MEM; Hyclone, Logan, UT, USA) containing 15% fetal bovine serum (FBS; Gibco, Grand Island, NY, USA), 1% penicillin/streptomycin (Invitrogen, Grand Island, NY, USA), 2 mM L-glutamine (Sigma-Aldrich, St. Louis, USA), and 55 μM 2-mercaptoethanol (Sigma-Aldrich, St. Louis, USA) in a 5% CO₂ humidified atmosphere at 37°C. Nonadherent cells were discarded after 48 h, and the adherent BMMSCs were cultured for 16 days until colonies are formed and then passaged once for further experimental use. The essential medium was changed every 2 days.

2.6. Flow Cytometry. Flow cytometry was used to test the surface markers of the isolated BMMSCs at passage three. Adherent cells were collected and incubated with a positive surface epitope profile with monoclonal antibodies for mouse CD105 and SC1 and negative cell surface markers with CD34 and CD45 (FITC anti-mouse CD34, CD45, CD105, and SC1 antibodies; BD Biosciences, USA) and then identified by flow cytometry analysis (BD Biosciences, San Jose, CA, USA), and the data were analyzed with EPICS XL software.

2.7. Osteogenic Differentiation and Mineralization Assays. The flow cytometry was used to identify BMMSCs in all experiments between three and seven passages. After the BMMSCs cultured in essential medium reached 70% confluence, the medium was replaced with an osteogenic medium containing essential medium and supplemented with 2 mM β-glycerophosphate (Sigma-Aldrich, St. Louis, USA), 100 μM L-ascorbic acid 2-phosphate (Sigma-Aldrich, St. Louis, USA), and 10 nM dexamethasone (Sigma-Aldrich, St. Louis, USA).

Identified BMMSCs (1 × 10^5) and mouse preosteoblastic MC3T3-E1 cells purchased from American Type Culture Collection (ATCC, USA) were seeded in 6-well plates with the osteogenic medium for 7 days. Alkaline phosphatase (ALP) staining (Beyotime Institute of Biotechnology, Shanghai, China) was performed using a standard protocol after 7 days of induction in 24-well plates. Further, alizarin red staining (Beyotime Institute of Biotechnology, Shanghai, China) was performed after culturing in osteogenic medium for 28 days to detect mineralized nodule formation. Images were captured with a converted microscope (Olympus, Tokyo, Japan), and quantitative analysis was performed with the Image-Pro Plus software.

2.8. Cell Proliferation Assay. The effects of ethanol or necrostatin-1 on cell proliferation were evaluated using the Cell Counting Kit-8 assay (Beyotime Institute of Biotechnology, Shanghai, China). Briefly, BMMSCs and MC3T3-E1 (1 × 10^5 cells/well) were seeded in 96-well plates and cultured in 100 μl growth medium until the cells proliferated to about 30%, respectively. The cells were then starved for 6 h, and the medium was replaced with osteogenic differentiation induction medium, and different concentrations of alcohol were added at 24, 48, and 72 h, respectively. After alcohol stimulation, 10 μl CCK-8 solution and 90 μl fresh medium were added to each well, with 5 repeat wells, and incubated at 37°C for 2 h, according to the manufacturer's protocol. Cell proliferation was assessed by measuring the absorbance at a wavelength of 450 nm (Thermo Fisher Scientific, MA, USA), and the appropriate time and concentration of alcohol intervention were determined. The same method was followed for necrostatin-1 (50 μmol/l).

2.9. Real-Time PCR Analysis. Cells were cultured in 6-well plates with osteogenic medium and treated with alcohol (150 mM) and necrostatin-1 (50 μM) for 48 h to detect the gene expression of necroptosis markers (RIPK1, RIPK3, and MLKL), treated for 7 days to test the expression of Runx2, then extracted total RNA using TRIzol reagent (Qiagen, Valencia, CA, USA), and assessed the concentration and purity by a spectrophotometer. Reverse transcriptase reactions were completed by ReverTra Ace® qPCR RT Kit (Toyobo, Japan), and real-time PCR was performed using SYBR® Premix Ex Taq (2x) (Toyobo, Japan). Finally, the CT values were calculated in relation to β-actin CT values (RQ = −ΔΔCt). Primers were synthesized by Primer Premier 5.0 software, using Primer-BLAST to confirm the definition of primers in NCBI website, and synthesized by Shanghai Biotechnology Co., Ltd. (Supplementary Table S1). The same method also applied to examine the gene expression of mouse bone tissue.

2.10. Western Blot Analysis. BMMSCs were cultured and treated as described previously, and proteins were harvested with RIPA cell lysis buffer system (Cell Signaling Technology, USA) supplied with phosphatase inhibitors. Total protein concentrations were quantified by BCA protein assay kit (Beyotime Institute of Biotechnology, Shanghai, China). Equal amounts of proteins (20 μg) were separated on 10% SDS-PAGE for protein electrophoresis and transferred to 0.2 μm polyvinylidene fluoride (PVDF) membranes, which
were blocked with 5% nonfat dry milk for 1 h at room temperature and incubated at 4°C overnight with primary antibodies to mouse Runx2, RIPK1, RIPK3, MLKL, phosphor-RIPK3 (p-RIPK3), phosphor-MLKL (p-MLKL), cleaved caspase-3, and β-actin (1:1000; Abcam, Cambridge, UK) and phosphor-RIPK1 (p-RIPK1; 1:1000; Cell Signaling Technology, USA). After washing three times in PBST and incubated for 1 h with HRP-conjugated second antibody at room temperature (1:2000, Abcam, Cambridge, UK), immunoreactive protein was detected using enhanced chemiluminescence reagents (Millipore, USA) and immunoblots were quantified with ImageJ software. MC3T3-E1 were cultured with osteogenic differentiation induction medium and treated as described previously to verify EtOH-induced osteoblast necroptosis.

2.11. Double-Label Immunofluorescent Staining. After 48 h of intervention on confocal dishes, cells were fixed with 4% PFA for 30 min, permeabilized in 0.5% Triton X-100 for 10 min, and incubated with primary antibodies of anti-mouse RIPK1 (1:100; Abcam, Cambridge, UK) and anti-rabbit RIPK3 (1:100; Abcam, Cambridge, UK) at 4°C for overnight and then incubated with fluorescent goat anti-mouse and goat anti-rabbit secondary antibodies (1:200; Abcam, Cambridge, UK) for 4 h at room temperature. After stained with DAPI, acquired confocal images by OLYMPUS inverted laser scanning confocal microscope (Olympus, Tokyo, Japan) and mounted the positive coexpression of RIPK1 and RIPK3 rates.

2.12. Transmission Electron Microscopy (TEM). After EtOH treatment for 48 h, the specimens were fixed in 2.5% glutaraldehyde for 2 h and postfixed in 1% osmium tetroxide for 90 min at 4°C, dehydrated in ethanol, and embedded in epoxy resin. The cells were examined under a transmission electron microscope (H-7700, Hitachi, Japan).

2.13. Intracellular Reactive Oxygen Species Assay. After EtOH treatment for 48 h, the ROS generation of MC3T3-E1 was detected by a ROS detection kit (Beyotime Institute of Biotechnology, Shanghai, China). The fluorescence of 2′,7′-dihydrodichlorofluorescein diacetate (DCFH-DA) was determined using a spectrofluorophotometer, and the mean fluorescence intensity (MFI) was analyzed by flow cytometry.

2.14. ELISA Assay. Peripheral blood serum was collected; serum alcohol concentration was measured with mouse EnzyChrom™ Ethanol Assay Kit (BioAssay Systems, USA). TNF-α level was analyzed using mouse tumor necrosis factor-α ELISA kit (Andy Gene, China) with mouse peripheral blood serum and cell culture medium, according to the manufacturer’s instructions. BALP; C-terminal telopeptide of type I collagen (CTX-I), tartrate-resistant acid phosphatase 5b (TRACP 5b) levels, and proinflammatory cytokines (IL-1β and IL-6) were analyzed using mouse ELISA kit (Andy Gene, China) with mouse peripheral blood serum.

2.15. Statistical Analysis. All experimental data were expressed as the mean ± standard deviation (SD) from at least three independent experiments. Comparisons between the two groups were analyzed using independent two-tailed Student’s t-test, and comparisons between multiple groups were accomplished using one-way ANOVA with the Bonferroni adjustment with SPSS 22.0 software (SPSS Inc., Chicago, IL, USA). p values < 0.05 were considered statistically significant.

3. Results

3.1. EtOH Treatment Leads to Osteogenic Differentiation Reduction and Osteopenia. In order to verify the effects of alcohol treatment on bone tissue, C57BL/6J male mice were administered via gavage with 0.2 ml/10 g/d (30% v/v) of alcohol for 22 weeks. All the animals were alive with no complications, and there was a significant difference in the body weight between the control group and EtOH-treated group after administration of alcohol for 18 weeks (p < 0.05, Figure 1(a)). The serum alcohol concentration of the mice in the EtOH-treated group was twice as much as the control group when sacrificed (p < 0.05), indicating the success of alcohol gavage (Figure 1(b)). BALP, an early marker of osteoblastic differentiation, in the serum was significantly decreased in the EtOH-treated group compared to that in the control group (p < 0.05, Figure 1(c)). Moreover, chronic alcohol consumption significantly reduced the bone mineral density (BMD) and mineral content of bilateral femurs, as measured by DXA (p < 0.05, Figures 1(d) and 1(e)), and markedly decreased the BMD, BV/TV, Tb.Th, Tb.N and Ct.Th, concomitantly, increased the Tb.Sp by micro-CT analysis (p < 0.05, Figures 1(f) and 1(g)). Histomorphometric analysis performed on the H&E stained sections revealed that the distal trabecular bone structure was disordered and loose, and the numbers of trabecular bone (yellow rectangle area, p < 0.01) and the ratio of OS/BS (two yellow dotted lines) and Ob.S/BS were significantly decreased (p < 0.05); the ratio of ES/BS (two green dotted lines) was increased (p < 0.01) compared to untreated mice (Figures 1(h) and 1(i)). Immunofluorescent staining, western blotting, and RT-PCR detected the expression of Runx2, a specific marker of osteogenic differentiation, and the results revealed that mRNA and protein levels of Runx2 were significantly decreased (Figures 1(j)–1(n)). Thus, this indicated that chronic alcohol consumption can lead to osteogenesis reduction and osteopenia in mice.

3.2. EtOH Treatment Leads to Osteoblast Necroptosis. To investigate the cause of EtOH-induced osteogenesis reduction and osteopenia, we detected necroptosis signaling. Interestingly, we found that high doses of EtOH markedly elevated the osteoblast necroptosis of bone tissue, as located and evidenced by the increased number of RIPK1-positive cells (Figures 2(a) and 2(b)) and RIPK3-positive cells (Figures 2(c) and 2(d)). The mRNA levels of RIPK1 and RIPK3 were correspondingly increased (Figures 2(e) and 2(f)). The levels of bone resorption markers, CTX-I and TRACP 5b, in serum (p < 0.05, Supplementary Figure S1a, b) and the number of osteoclasts in the femurs identified by TRACP staining increased in the EtOH-treated mice.
Figure 1: Continued.
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Surprisingly, the expression of RIPK1 in osteoclasts had no significant difference between the alcohol intervention group and the control group with colocalization (Supplementary Figure S1c). Overall, necroptosis of osteoblasts was more obvious than osteoclasts with alcohol intervention, so our experiment mainly focused on bone formation rather than bone resorption.

To determine the type of cell death that a cell undergoes, antiactive caspase-3 was detected. The results showed that the protein levels of cleaved caspase-3 in mouse bone tissue and MC3T3-E1 cells were increased in the EtOH-treated group evaluated by immunofluorescent staining and western blot analysis, but the difference was not statistically significant (p > 0.05, Figures 2(g) and 2(h)). Consistently, TEM analysis revealed that cells displayed typical necrotic ultrastructural changes including a disrupted plasma membrane, fragmented and vacuolated mitochondrial membranes, and overflown cytoplasmic content after exposure to 150 mM EtOH for 48 h (Figures 3(i) and 3(j)) that exhibited the typical characteristic of necroptosis rather than apoptosis. The above results suggested that apoptosis accounts for a small proportion in the occurrence of alcohol-induced osteopenia, but mainly necroptosis.

Further, to confirm the activation of necroptosis in the EtOH-treated group, we performed western blotting to semiquantitatively analyze the changes in the protein...
Figure 2: Continued.
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expression levels. The results showed upregulated expression of RIPK1 and p-RIPK1 (Figures 2(k) and 2(l)) and RIPK3 and p-RIPK3 (Figures 2(m) and 2(n)) with EtOH treatment in vivo, indicating that chronic heavy alcohol-induced bone loss in mice was associated with osteoblast necroptosis activation.

3.3. EtOH Induces Osteoblast Necroptosis by Activating the RIPK1/RIPK3/MLKL Signaling. To investigate the mechanism of EtOH-induced osteoblast necroptosis, we added 150 mM EtOH to BMMSCs, which were identified by flow cytometry analysis (Supplementary Figure S2a) and then induced to differentiate into osteoblasts that were evaluated by alkaline phosphatase (ALP) staining (Supplementary Figure S2b) and alizarin red staining (Supplementary Figure S2c), for 48h using the CCK-8 assay (Supplementary Figure S3a, b). Quantitative western blotting showed that EtOH treatment elevated the expression of RIPK1 and p-RIPK1 (Figures 2(k) and 2(l)) and RIPK3 and p-RIPK3 (Figures 2(m) and 2(n)) with EtOH treatment in vivo, indicating that chronic heavy alcohol-induced bone loss in mice was associated with osteoblast necroptosis activation.

3.4. Necrostatin-1 Treatment Ameliorates Osteopenia and Osteogenic Differentiation Reduction in EtOH-Treated Mice. Next, to examine whether necrostatin-1 treatment ameliorated osteopenia in EtOH-treated mice, we treated the mice with 1.65 mg/kg/d of Nec-1 for 22 weeks and BMMSCs with 50 μM of Nec-1 for 48h, respectively (Supplementary Figure S3c, d). We found that Nec-1 treatment ameliorated EtOH-induced low BMD, trabecular BV/TV, Tb.Th, Tb.N, and Cl.Th in the distal femurs, as assessed by DXA, micro-CT cross sections, and 3D reconstructions, respectively (Figures 4(a)–4(d)). Further, H&E staining revealed that Nec-1 treatment inhibited the reduction in trabecular bone volume and number, facilitated partial recovery of the loose trabecular structure, and increased the ratio of OS/BS and Ob.S/BS compared to EtOH-treated mice (Figures 4(e) and 4(f)). Moreover, Nec-1 treatment significantly increased bone formation, as indicated by the increase in alkaline phosphatase activity and the formation of mineralized nodules in osteoblasts (Figure 4(g)) and serum ALP activity in mice (Figure 4(h)). Further, immunofluorescence staining (Figures 5(a) and 5(b)), qPCR (Figure 5(c)), and western blot analysis both in vivo (Figures 5(d) and 5(e)) and in vitro (Figures 5(f) and 5(g)) showed that the expression level of Runx2 was downregulated after EtOH treatment; notably, necrostatin-1 treatment upregulated these expression levels.

3.5. Necrostatin-1 and NAC Treatment Ameliorates Osteopenia by Inhibiting the RIPK1/RIPK3/MLKL Signaling. We further explored whether necrostatin-1 treatment could inhibit RIPK1 kinase and investigated its effects
on the downstream signaling pathway of RIPK1 with EtOH intervention. Western blot analysis showed that the expression of RIPK1 and p-RIPK1 (a, b), RIPK3 and p-RIPK3 (c, d), and p-MLKL (e, f) on MC3T3-E1 cells were elevated in the EtOH treatment group, while the expression of caspase-8 was not significantly influenced (g, h). Double-labeled immunofluorescence staining confirmed that necroptosis makers RIPK1 and RIPK3 coexpressed in the EtOH-treated MC3T3-E1 cells (i). Bar: 15 μm. And p-MLKL aggregated on the cell membrane, leading to perforation and destruction of the integrity of proliferating MC3T3-E1 osteoblasts (j). Bar: 20 μm. Error bars represent the SD from the mean values. *p < 0.05. p-MLKL: phosphorylated mixed lineage kinase domain-like protein.

Oxidative stress is known as one trigger for alcohol-induced osteopenia, and increased levels of ROS may directly precede the production of inflammatory cytokines in the bone marrow and contribute to decreased bone formation [16]. However, whether ROS could increase osteoblast necroptosis in alcohol-induced osteopenia remains unclear. Therefore, we measured the effect of alcohol intervention on the intracellular ROS levels by flow cytometry, and our results supported that high dose of EtOH consumption increased the intracellular ROS levels compared with the control group, and Nec-1 directly inhibited the elevated ROS (Figure 6(f)). Several necroptotic stimuli have been proposed to induce necroptosis, and ROS are essential factors [46]. Next, we explored whether ROS are important for EtOH-induced necroptosis in osteoblastic cells, using the
Figure 4: Continued.
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Figure 4: Necrostatin-1 treatment ameliorates osteopenia in EtOH-treated mice. Nec-1 treatment ameliorated EtOH-induced low BMD in femurs (a-d) and BV/TV, Tb.Th, Tb.N, and Ct.Th in the distal femurs, as assessed by DXA (a, b) and axial micro-CT cross sections and 3D reconstructions (c, d), respectively. H&E stained sections revealed that Nec-1 treatment inhibited the reduction in trabecular bone volume and number and facilitated partial recovery of the destroyed trabecular structure (e, yellow rectangle area), increased the ratio of OS/BS (e, two yellow dotted lines) and Ob.S/BS, and decreased the ratio of ES/BS (e, two green dotted lines) compared to EtOH-treated mice (f) Bar: 100 μm. Nec-1 treatment significantly increased bone formation, as indicated by the increase in alkaline phosphatase activity (ALP staining) and the formation of mineralized nodules (alizarin red staining) in osteoblasts (g) and serum alkaline phosphatase activity in mice (h). Error bars represent the SD from the mean values. *p < 0.05. Nec-1: necrostatin-1.
Figure 5: Necrostatin-1 treatment ameliorates osteogenic differentiation reduction. Immunofluorescence staining (a, b; bar: 100 μm), RT-PCR (c), and western blot analysis both in vivo (d, e) and in vitro (f, g) showed that the expression level of Runx2 was downregulated after EtOH treatment; however, necrostatin-1 treatment upregulated the expression level. (a–e) Data from mice. (f, g) Osteoblasts. Error bars represent the SD from the mean values. *p < 0.05 compared with the control group. #p < 0.05 compared with the EtOH-treated group.
Figure 6: Continued.
Figure 6: Continued.
antioxidant NAC to inhibit the production of ROS. In this study, the addition of NAC effectively reduced EtOH treatment necroptosis of the osteoblastic cells as NAC attenuated the upregulation of RIPK1 and RIPK3 (Figure 6(g)). Meanwhile, NAC treatment significantly increased bone formation, as indicated by the increase in ALP staining and the formation of mineralized nodules (alizarin red staining) in osteoblasts (Figure 6(h)). Together, these results highlight that osteoblast necroptosis, at least in part, promotes the generation of ROS, and on the other hand, ROS play a critical role in the regulation of excessive EtOH treatment-induced osteoblast necroptosis via a positive feedback loop.

4. Discussion

Chronic and excessive alcohol consumption causes a reduction in bone mass and destruction of the bone microstructure due to an impairment and/or imbalance in bone remodeling. However, the underlying mechanisms have not yet been elucidated. In this study, we successfully established an alcohol-induced osteopenia mouse model and found that osteopenia is accompanied by decreased bone formation and increased osteoblast necrosis, which was reported in studies on GIOP and ovariectomized rats [40, 41, 47]. Long-term detrimental effects of alcoholism have toxic effects on BMMSCs and affect their differentiation into...
osteoblasts by downregulating the expression of Runx2, which in turn leads to low bone mass [48, 49]. In addition, previous studies have confirmed that apoptosis was involved in alcohol-induced osteoporosis [18, 50, 51]. Herein, we found that loose trabeculae were more obvious, and the lacunae volume was larger in the EtOH-treated mice than in the control mice. Moreover, some of the MC3T3-E1 treated with alcohol suffered a necrotic change as observed from certain morphological changes and specific markers. These conclusions indicate the involvement of necrosis-associated mechanisms in the osteoblasts treated with EtOH.

Necroptosis is a caspase-independent form of programmed cell death that is triggered by ligation of death receptors or initiators that bind deubiquitylated RIPK1 to autoactivated RIPK3 and initiate the phosphorylation of a pseudokinase substrate MLKL, which then combines with autoactivated RIPK3 and initiate the phosphorylation of programmed cell death that is triggered by ligation of death mechanisms in the osteoblasts treated with EtOH.

Studies revealed that necroptosis was involved in the regulated cell death induced by TNF-α in nonalcoholic fatty liver disease [66], osteoblasts [67], glycerol-induced acute kidney injury [68], etc. However, the specific regulatory mechanism of TNF-α on bone cells and the molecular mechanism in the pathogenesis of osteoporosis remain unclear. It has been reported that TNF-α plays a role in promoting alcohol-induced osteoporosis [69, 70]. Given that necroptosis is a special type of necrosis mainly mediated by the stimulating death receptor of TNF-α [27, 71], we examined the level of TNF-α in mouse serum and osteoblastic culture medium after EtOH intervention and Nec-1 treatment by ELISA. Both serum and medium contained elevated levels of TNF-α in the EtOH-treated mice and cells but had no significant difference after Nec-1 treatment (Figures 6(k) and 6(l)). Therefore, we conjectured that alcohol could induce necroptosis by activating the expression of TNF-α and downstream molecules. Unfortunately, we could not prove the role of TNF-α in the process of EtOH-induced necroptosis. Moreover, detecting the RIPK1/RIPK3/MLKL axis by treating osteoblasts with TNF-α plus the specific inhibitor of necroptosis with or without Nec-1 is necessary to validate that chronic alcohol conditions promote RIPK1 activation with TNF signaling.

ROS have been considered as a driving force for necroptosis, the mitochondrial ROS derived in response to the TNF/TNFR1 pathway [72], and nonmitochondrial ROS generated via TNF-induced necroptosis [73]. Studies showed that treatment with TNF-α could stimulate ROS production, which enhanced necrosome formation [74]. ROS promote RIPK1 autophosphorylation on serine residue 161 (S161), and three crucial cysteines (cysteines 257, 268, and 586) in RIPK1 are required for sensing ROS to regulate RIPK1. The major function of RIPK1 kinase activity in TNF-induced necroptosis is to autophosphorylate S161 which then forms a functional necroosome [75]. TNF-induced ROS function in a positive feedback to enhance necroosome formation and induce necroptosis, when the necrosome complex was impaired or knockeddown the RIPK1 or RIPK3,
led to a decrease in ROS production [74]. In our study, alcohol treatment increased the production of intracellular ROS levels and necroptosome complex. Conversely, treatment with Nec-1 markedly suppressed necroptosis and ROS production in EtOH intervention osteoblasts. Altogether, osteoblast necroptosis partly promotes the generation of ROS; on the other hand, ROS contributed to necroptosis of osteoblasts through a positive feedback loop involving RIPK1/RIPK3. Nec-1 and NAC inhibit the expression of RIPK1, affect the activation of necroptosis, and promote the survival and proliferation of osteoblast in the presence of death receptor ligands, eventually leading to a reduction in osteoblast necrosis and increased bone formation. TNFR: tumor necrosis factor receptor; FADD: Fas-associated protein with death domain; TRADD: TNFR1-associated death domain.

**Figure 7**: EtOH treatment induced osteopenia by increasing osteoblast necroptosis and decreasing BMMSC osteogenic differentiation. Chronic EtOH consumption results in osteopenia and osteoblast necroptosis in mouse model and leads to a necrotic morphology lesion and osteogenic differentiation reduction in bone marrow mesenchymal stem cells (BMMSCs). The ligation is stimulated by EtOH treatment that binds deubiquitylated RIPK1 to autoactivated RIPK3 and initiates the phosphorylation of a pseudokinase substrate MLKL, which then combines with p-RIPK3 to form a necrosome and translocates to the membrane to promote cell lysis and inflammation release (IL-1β and IL-6). Excessive EtOH treatment-induced osteoblast necroptosis might partly depend on ROS generation; concomitantly, ROS contributed to necroptosis of osteoblasts through a positive feedback loop involving RIPK1/RIPK3. Nec-1 and NAC inhibit the expression of RIPK1, affect the activation of necroptosis, and promote the survival and proliferation of osteoblast in the presence of death receptor ligands, eventually leading to a reduction in osteoblast necrosis and increased bone formation. TNFR: tumor necrosis factor receptor; FADD: Fas-associated protein with death domain; TRADD: TNFR1-associated death domain.

5. **Conclusions**

In summary, this study provides evidence that excessive EtOH treatment leads to activation of the RIPK1/-RIPK3/MLKL signaling, promotes the generation of ROS, resulting in an increase in osteoblast necroptosis and a decrease in BMMSC osteogenic differentiation both in vivo and in vitro, and causes bone mass reduction. Necrostatin-1-mediated treatment effectively inhibits the RIPK1/-RIPK3/MLKL-dependent signaling, and NAC partly prevents the production of ROS and decreases the activation of necroptosis through a positive feedback loop involving RIPK1/RIPK3 and finally ameliorates alcohol-induced osteopenia by reducing osteoblast necroptosis. Collectively,
we demonstrated that chronic high-dose alcohol consumption induces osteopenia via necroptosis and thereby discovered that RIPK1 kinase may be a therapeutic target for alcohol-induced osteopenia. However, the detailed mechanism and other molecules involved need to be further investigated.

Data Availability
The original data used to support the findings of this study are available from the corresponding authors upon request.

Conflicts of Interest
The authors declare that they have no conflict of interest.

Authors’ Contributions
M. G. and Y. X. participated in the study conception and design. M. G., Y. H., Z. J., and F. T. performed the cell culture, qPCR, and signaling studies. M. G., Q. W., L. C., and X. T. designed and performed the animal experiments. M. G. and S. W. scanned the specimens with the micro-computed tomography (micro-CT) scanner. Q. W., Y. L., Y. Z., and J. G. prepared the figures and performed data analysis. M. G., F. T., and Y. X. prepared and revised the manuscript. Man Guo, Yong-Li Huang, and Qi Wu contributed equally to this work.

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Supplementary Materials
Table S1: DNA primers used in quantitative real-time PCR. Figure S1: bone resorption in alcohol-induced osteopenia. Figure S2: identified the isolated mouse BMMSCs and osteoblast. Figure S3: CCK-8 assay was performed to determine the cell proliferation ability. Figure S4: necrostatin-1 treatment inhibited the RIPK1/RIPK3/MLKL signaling. (Supplementary Materials)

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