Valproic acid prevents glucocorticoid-induced osteonecrosis of the femoral head of rats

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Abstract. Glucocorticoids (GCs) are the most common cause of atraumatic osteonecrosis of the femoral head (ONFH) because their effect compromises the osteogenic capability of bone marrow-derived mesenchymal stem cells (BMSCs). Valproic acid (VPA) is a widely used anti-epileptic and anti-convulsant drug. Previous studies have reported that VPA promotes osteogenic differentiation of BMSCs in vitro and osteogenesis in vivo as a histone deacetylase (HDAC) inhibitor. The purpose of the present study was to investigate the efficacy of VPA as a precautionary treatment of ONFH after GC treatment in rats. In vitro, the effect of VPA, dexamethasone or a combination treatment of the two on the proliferation and osteogenic differentiation of human BMSCs was assessed using a cell counting Kit-8 and apoptosis assays, and by measuring the expression of proteins associated with osteogenesis. In vivo, a GC-induced ONFH model was established in rats and VPA was added during GC treatment to investigate the preventive effect of VPA against ONFH. Rat BMSCs were also extracted to investigate the osteogenic capacity. The results of micro-computed tomography scanning, angiography of the femoral head and histological and immunohistochemical analyses indicated that 11 of 15 rats induced with methylprednisolone (MP) presented with ONFH, while only 2 of 15 rats treated with a combination of MP and VPA developed ONFH. VPA produced beneficial effects on subchondral bone trabeculae in the femoral head with significant preservation of bone volume and blood supply, as well as improved osteogenic capability of BMSCs compared with those in rats treated with GC alone. In conclusion, VPA attenuated the inhibitory effect of GC on BMSC proliferation and osteogenesis by inhibiting apoptosis and elevating the expression of proteins associated with osteogenesis, which may contribute to the prevention of GC-induced ONFH in rats.

Introduction

Osteonecrosis of the femoral head (ONFH) is a refractory disease affecting young individuals, which is characterised by hip pain and dysfunction or even lameness (1,2). While several risk factors are known, glucocorticoid (GC) medication is thought to be the principle one, causing 10-30% of non-traumatic ONFH (3). GC has been widely used in the treatment of rheumatic and auto-immune diseases, and for chemotherapy and acute medical conditions, including traumatic spinal cord injury (4-7). Hence, ONFH is a potential complication for numerous patients receiving GC treatment. The pathogenesis of ONFH includes ischaemia, necrosis, repair and deformity in the femoral head (8-10). All of these changes occur sequentially, resulting in collapse of the femoral head. GC may cause ONFH through several mechanisms. First, a GC overdose often exerts adverse effects on the vascular system, including hypertension, atherosclerosis and coagulation abnormalities, leading to diminished blood supply in the femoral head (11,12). Secondary to GC administration, patients present with a decreased pool of mesenchymal stem cells (MSCs) in the proximal femur (13) and a reduced activity of MSCs (14). MSCs have a vital role in the osteonecrosis repair process. When osteonecrosis appears, fibrous tissue carrying undifferentiated mesenchymal cells invades the necrotic area and then initiates new bone formation; however, this is slower than the spread of fibrous tissue, therefore leading to incomplete reconstruction of the necrotic area (15-18). GC has been reported to inhibit the proliferation and osteogenic differentiation of MSCs. The inhibitive role of GC may be associated with the induction of apoptosis and downregulation of runt-related transcription factor 2 (Runx2)/core-binding factor α1 (Cbfα1), which is considered to be important in osteogenesis, particularly for...
the development of osteoblasts (19,20). Furthermore, MSCs extracted from steroid-induced ONFH patients have a compromised osteogenic ability, an elevated adipogenic ability and a lower proliferation rate (21-25). Therefore, the enhancement of proliferation and osteogenesis in bone marrow-derived MSCs (BMSCs) in patients receiving GC by other drugs may improve the repair process or even prevent ONFH.

Valproic acid (VPA) has been approved by the Food and Drug Administration of the US to treat epilepsy for >20 years (26). It has been reported that VPA promotes the osteoblast differentiation and maturation processes through the regulation of cell histone acetylation (27,28). Histone acetylation is a reversible epigenetic process delicately modulated by histone acetyltransferases (HATs) and histone deacetylases (HDACs). HATs are responsible for histone hyperacetylation, resulting in relaxation of the chromatin structure, and allow the binding of DNA sequences with transcription factors, thereby activating transcription and downstream expression. Conversely, HDACs deacetylate histones and silence transcription, thereby counteracting the effects of HATs (29). This epigenetic regulation of gene expression has a dominant role in cell stemness, determination, commitment and differentiation (30-33). HDACs are mainly classified into 4 groups: class I HDAc (1, 2, 3 and 8), class IIa HDacs (4, 5, 7 and 9), class IIb HDacs (6 and 10), class III HDacs (sirtuin-1 to -7) and a class IV H dAc (H dAc11) (34). VPA, a potent HATi, has been reported to be a dominant H dAci, which has been reported to inhibit cell migration and invasion (35-38). To the best of our knowledge, the effect of VPA on BMSc proliferation and osteogenesis under the prevalence of GC has not been explored. Previous studies (46,47) have reported that VPA has a dominant role in cell proliferation and osteogenesis reported in vivo. VPA promotes the osteogenic differentiation of BMSCs and thus prevent the occurrence of ONFH after GC administration. The results of the present study indicated that VPA reduces the inhibitory effect of GC on proliferation, apoptosis and osteogenic differentiation of BMSCs in vitro and prevents the occurrence of ONFH in rats.

Materials and methods

Cell culture. The present study was performed according to the principles of the Helsinki Declaration, and written informed consent was obtained from each patient. The experimental procedures were approved by the Ethical Review Board of Shanghai Jiaotong University Affiliated Sixth People’s Hospital (Shanghai, China). BMSCs were isolated from human bone marrow harvested from patients suffering femoral neck fracture during hip arthroplasty surgery. The BMSCs were purified and cultured according to a previously published protocol by our group (45). BMSCs of passages 3-6 were used in all experiments. Patient consent was provided before any procedures were performed. BMSCs from proximal femurs of rats were extracted and cultured using the same methods.

Cell proliferation and apoptosis assay. First, the effect of VPA, dexamethasone (DEX) and a combination of the two drugs on BMSC proliferation and apoptosis was assessed. BMSCs were seeded in 96-well plates (3-wells per group) with the same number of cells and divided into 5 groups: i) Control; ii) VPA 0.5 mM; iii) VPA 1 mM; iv) VPA 2 mM; and v) VPA 5 mM. DEX at 10⁻⁷ M was selected due to its significant inhibitory effect on cell proliferation and osteogenesis reported in previous studies (46,47). In addition, the following DEX groups with or without VPA treatment were established: i) DEX; ii) DEX+VPA 0.5 mM; iii) DEX+VPA 1 mM; iv) DEX+VPA 2 mM; and v) DEX+VPA 5 mM. The Cell Counting Kit-8 (CCK-8) assay was used according to the manufacturer’s instructions at 24, 72, 120 and 168 h after BMSC adherence and the above treatments. At these time-points, 10 µl CCK-8 reagent was added into 100 µl of culture medium in each well, followed by incubation for another 3 h. The absorbance value was measured using a microplate reader (Bio-Rad Laboratories, Inc., Hercules, CA, USA) at 450 nm.

To evaluate the effect on apoptosis, BMSCs cultured with 5 mM VPA, DEX and DEX plus 5 mM VPA for 5 days were subjected to Annexin V-fluorescein isothiocyanate and propidium iodine double staining (Dojindo, Kumamoto, Japan), and the apoptotic rate was measured by flow cytometry according to the manufacturer’s protocols.

Osteogenic induction assay. BMSC differentiation was initiated 48 h after the cells were plated, with the basic medium in each group supplemented with 10⁻² M β-sodium glycerophosphate, 50 µg/ml L-aspartic acid and 10⁻⁷ M DEX. The medium was changed every 3 days. First, the osteogenesis-associated mRNA levels of BMSCs treated with VPA alone for 7 and 14 days were measured via a polymerase chain reaction (PCR) assay. Next, BMSCs were osteo-induced with VPA at gradient concentrations for 7 and 21 days in a mineralisation assay. Subsequently, the effect of DEX in the presence or absence of VPA on BMSC osteogenesis and mineralisation was tested. For PCR and western blot assays, BMSCs were cultured for 7 days. For alizarin red S staining, BMSCs in each group were cultured for 21 days. Alkaline phosphatase (ALP) activity was performed using EasyScript one-step gDNA Removal and cDNA Synthesis Supermix (TransGen Biotech, Beijing, China) from 1 µg of total RNA according to the manufacturer’s protocols. qPCR of type I collagen (COL I, ALP, osteocalcin (OCN) and Runx2 were performed with the TransStart Tip Green qPCR SuperMix (TransGen Biotech). The relative amount of mRNA was normalised to β-actin (48). The forward and reverse primers of each complementary (c)DNA were designed as follows: β-actin, 5’-GTCATCCATGGCGAACTCGT-3’ and 5’-CGTCATTCAGCGCCAATCTG-3'; Runx2, 5’-CCGAGACCAACCCAGTCTTTTA-3’ and 5’-AAGAGCTGTGTACCCCAT-3'; ALP, 5’-CAAGGATGCTGGGAAGTCG-3’ and 5’-CTCTGGCCGCATCTCATGTT-3’;

Reverse transcription-quantitative PCR (RT-qPCR) analysis. Total RNA was extracted from osteogenic-induced BMSCs in each group with TRIzol reagent (Invitrogen; Thermo Fisher Scientific, Inc., Waltham, MA, USA). The RT reaction was performed using EasyScript one-step gDNA Removal and cDNA Synthesis Supermix (TransGen Biotech, Beijing, China) from 1 µg of total RNA according to the manufacturer’s protocols. qPCR of type I collagen (COL I, ALP, osteocalcin (OCN) and Runx2 were performed with the TransStart Tip Green qPCR SuperMix (TransGen Biotech). The relative amount of mRNA was normalised to β-actin (48). The forward and reverse primers of each complementary (c)DNA were designed as follows: β-actin, 5’-GTCATCCATGGCGAACTCGT-3’ and 5’-CGTCATTCAGCGCCAATCTG-3'; Runx2, 5’-CCGAGACCAACCCAGTCTTTTA-3’ and 5’-AAGAGCTGTGTACCCCAT-3'; ALP, 5’-CAAGGATGCTGGGAAGTCG-3’ and 5’-CTCTGGCCGCATCTCATGTT-3’;
OCN, 5'-CCCCCTCTAGCTAGGACCC-3' and 5'-ACCGG
TAATGCGAGTGTTCG-3'; COL I, 5'-CAGCCGCTTCCACCTA
CACG-3' and 5'-TTTTGATATCCATCTA.GTTGCCC-3'.

The PCR system was as follows: cDNA, 1 µl; double-distilled
water, 3.4 µl; Tip Green qPCr SuperMix, 5 µl; passive reference
dye (50X), 0.2 µl; forward primer (10 µmol/l), 0.2 µl;
reverse primer (10 µmol/l), 0.2 µl. The total volume of the
system was 10 µl. The reaction conditions were 95° C for 30 sec
initial denaturation, followed by 40 cycles of 95° C for 5 sec
and finally 60° C for 30 sec. Furthermore, a 65-95° C solubility
curve was constructed.

Western blot analysis. Proteins were extracted with a cell lysis
buffer (radioimmunoprecipitation assay lysis and extraction Buffer, Thermo Fisher Scientific, Inc.) supplemented with
protease inhibitor (#78430; Thermo Fisher Scientific, Inc.), and the total protein concentration was detected with a bichinchoninic acid assay. Following denaturation at
95° C for 5 min, 30-µg aliquots of protein were subjected to
10% SDS-PAGE and transferred to a polyvinylidene fluoro
deride membrane (Merck KGaA, Darmstadt, Germany). After
being blocked with 5% dried skimmed milk, the membranes
were labelled with a primary antibody to COL I (ab138492),
Runx2 (ab23981), ALP (ab186422) and β-actin (ab15777)
(Abcam, Cambridge, UK) at a concentration of 1:1,000 at
4°C overnight and then immersed in the secondary antibody
working solution containing anti-rabbit immunoglobulin G
(#14708; Cell Signaling Technology, Inc., Danvers, MA, USA)
(1:1,000) at 37°C for 1 h. After chemiluminescence staining
(Pierce™ ECL, #32106; Thermo Fisher Scientific, Inc.) with
a commercial assay, the target bands were detected with a gel
image-processing system. The protein levels were normalised
against β-actin. Histone extraction was performed according
to the protocol of a previous study (35), and the membranes
were incubated with primary antibodies to HDAC1 (AH379;
Beyotime Institute of Biotechnology, Shanghai, China),
HDAC2 (AH382; Beyotime Institute of Biotechnology), acety-
lated histone H3 (ac-H3) (#06-599; EMD Millipore, Billerica,
MA, USA), ac-H4 (#06-598; EMD Millipore) and histone H4
(ab31830; Abcam), which was used as the loading control, at a
concentration of 1:500 at 4°C overnight. Proteins were detected
after incubation with goat anti-rabbit IgG secondary antibody
(1:1,000; #14708; Cell Signaling Technology, Inc.) at 37°C for
1 h and visualized using chemiluminescence in the same way
according to manufacturer's protocols mentioned above.

ALP activity assay. The osteogenic differentiation of human
(h)BMSCs was evaluated by the ALP activity assay. After
culture for 7 days, the cell layers were gently washed with
cold PBS and lysed in 200 µl of 0.2% Triton X-100 for 30 min.
The lysates were centrifuged at 20,000 g for 15 min at 4°C
and sonicated. Next, 30 µl of the supernatant was mixed with
150 µl of the working solution (Nanjing Jiancheng
Bioengineering Institute, Nanjing, China) according to the
manufacturer's protocol. The formation of p-nitrophenol from
p-nitrophenylphosphate, the substrate of ALP, was evaluated
by measuring the absorbance at 405 nm with a microplate
reader (Bio-Rad 680; Bio-Rad Laboratories, Inc.). The ALP
activity was calculated by determining the ratio of the absorb-
bance of the experimental samples to that of the standard and
was expressed as mM of p-nitrophenol produced per minute
per mg of protein.

Alizarin red S staining. After osteogenic induction for 21 days,
cells in 48-well plates (3-wells in each group) were fixed with
4% paraformaldehyde for 20 min, then rinsed twice with
PBS (pH 7.4) and stained with 40 mM alizarin red working
solution for 10 min at room temperature. After being rinsed
twice with PBS again, these cells were visualised under a
light microscope. Subsequently, 100 mmol/l cetylpyridinium
chloride was added to each well and semi-quantitative analyses
were performed by measurement of optical density values at
560 nm.

Animal model and grouping. All animal experiments in the
present study were approved by the Animal Care and Use
committee of Shanghai Sixth People's Hospital, Shanghai
Jiaotong University School of Medicine (Shanghai, China).
A total of 36 Sprague Dawley (SD) rats (weight, 250-300 g,
7 weeks old) purchased from Shanghai Animal Experimental
Centre, (Shanghai, China) were divided into 3 groups. Six rats
in the control group received no treatment; 15 rats in the
methylnprednisolone (MP) group were intramuscularly injected with
MP 20 mg/kg/day (49,50) for 3 continuous days per week over
a period of 3 weeks (total MP, 180 mg/kg) to induce ONFH,
and 15 rats in the MP+VPA group were intraperitoneally injected with
300 mg/kg VPA (51,52) once a day for 3 weeks in a row until MP injection was ceased. MP rather than DEX
was selected, as it is more commonly used in the clinic.

Serum ALP activity. Blood samples of 1 ml were harvested
randomly from 3 SD rats in each group on the last day of
weeks 1, 2 and 3 after injection. Serum separation was
performed by centrifugation at 300 x g for 15 min at 4°C. The
serum was then mixed with the working solution for the ALP
activity assay (#A059-2; Nanjing Jiancheng Bioengineering
Institute, Nanjing, China) and processed according to the
manufacturer's protocol. The optical density values were
measured at 405 nm with a microplate reader (Bio-Rad 680;
Bio-Rad Laboratories, Inc.).

Micro-computed tomography (CT) scanning. To evaluate bone
morphologic changes in the rats, the right femoral head of
each rat was scanned with a micro-CT scanner (Skyscan 1176;
Bruker MicroCT, Kontich, Belgium) at a resolution of
9 microns. Two-dimensional (2D) images were analysed using
CTan software (v.1.13; Bruker MicroCT). Primarily, the
trabecular bone parameters of whole subchondral bone in the
femoral head were measured. Specifically, the bone mineral
density (BMD), bone volume per tissue volume (BV/TV) and
trabecular thickness (Tb.Th) were quantified.

Angiography. After cardiac perfusion with heparinised saline,
Microfil (MV-112; Flow Tech, Inc., Carver, MA, USA) was
injected into the abdominal aorta until a constant outflow of
the compound was observed from the abdominal vein. All
the surgical steps were under anaesthesia after intraperitoneal
injection of 4% chloral hydrate (360 mg/kg). Subsequently, the
rats were stored at 4°C overnight to ensure the polymerisa-
tion of the contrast agent. Femoral heads were fixed with 10%
formalin and decalcified with a 10% EDTA solution. Finally, the samples were scanned via micro-cT as described above.

**Histological and immunohistochemical (IHC) analyses.**

After decalcification and paraffin embedding, femoral heads were sectioned at a thickness of 5 µm in the coronal plane. Certain sections were stained with haematoxylin and eosin (H&E) at room temperature for 5 min to evaluate the trabecular structure, while others were deparaffinised, subjected to antigen retrieval and incubated with anti-OcN (1:50; ab13420; Abcam), anti-Runx2 (1:200; ab23981; Abcam) and anti-vascular endothelial growth factor (VEGF; 1:50; BA0407; Boshide, Wuhan, China) primary antibodies at 4˚C overnight and then incubated with the Super Vison Polymer anti‑rabbit lgG‑HRP kit at 37˚C for 30 min according to the manufacturer's protocols (SV0002; Boshide). Staining was visualized with 3,3-diaminobenzidine and samples were counterstained with haematoxylin for 1 min at room temperature. Photomicrographs were acquired using a Leica DM 4000 (Leica Microsystems, Wetzlar, Germany).

**Statistical analysis.** SPSS 20.0 (IBM Corp., Armonk, NY, USA) was used to analyse the values in each group. Values are expressed as the mean ± standard deviation. Comparisons of data among the groups were performed using one-way analysis of variance with Student-Newman-Keuls post hoc analysis. P<0.05 was considered to indicate a statistically significant difference.

**Results**

**VPA at a high concentration inhibits BMSC proliferation,** while **VPA at lower concentrations inhibits DEX-induced decreases in BMSC proliferation.** The CCK-8 assay indicated that the amount of BMSCs in all groups gradually increased in a time-dependent manner, but proliferation was significantly repressed after incubation with 5 mM VPA and with 10⁻⁵ M DEX in the presence or absence of VPA. The BMSC proliferation rate was not significantly influenced by 0.5-2 mM VPA, but these concentrations of VPA attenuated the inhibitory effect on BMSC proliferation exerted by DEX in a dose-dependent manner, except for the 5-mM concentration of VPA, which decreased the proliferation compared with that in the groups treated with DEX alone (Fig. 1).

**VPA decreases the apoptotic rate of BMSCs induced by DEX.** The apoptosis assay indicated that DEX promoted cell apoptosis. The apoptotic cells in the DEX group was >40%, which was significantly higher than that in the control group (24.83%). Although 5 mM VPA inhibited BMSC proliferation, it exerted no significant effect on cell apoptosis. However, when 5 mM VPA was added to the DEX culture, the amount of apoptotic BMSCs was significantly reduced (Fig. 2).

**VPA increases osteogenic differentiation and improves the osteogenic capacity of BMSCs compromised by DEX treatment by inhibition of HDAC activity.** First, BMSCs were treated with different concentrations of VPA, and RT-qPCR analysis revealed that 1 mM VPA significantly increased the mRNA levels of Runx2, ALP, COL I and OcN after 7 days of culture. After osteogenic induction for 14 days, VPA at all concentrations promoted the expression of these osteogenesis-associated genes to varying degrees, while decreasing that of Runx2. The most efficacious concentration of VPA at 7 and 14 days was 1 mM (Fig. 3). Next, the protein expression of Runx2, COL I and ALP, as well as ALP activity, was investigated in the same treatment groups. The results indicated that VPA elevated ALP activity and the protein expression of ALP, COL I and Runx2 (Fig. 4A-C). With VPA treatment at 1 mM, a similar trend to that in ALP activity was observed regarding ALP expression (Fig. 4D). Given that 5 mM VPA caused a downward trend in proliferation, only the lower concentrations (0.5-2 mM) were added into osteogenic medium containing 10⁻⁵ M DEX, in which BMSCs were cultured for 7 days. The results indicated that DEX decreased the protein levels of Runx2, COL I and ALP in comparison with those in the control group, while the addition of 0.5 and 1 mM VPA enhanced the levels of all of these proteins to varying degrees. Specifically, 0.5 and 1 mM VPA recovered Runx2 expression to the same level as that in the control group, and rescued ALP expression. Furthermore, supplementation with 0.5 mM VPA gave rise to
significantly higher expression levels of COL I than those in the control and DEX groups. However, the addition of 2 mM VPA repressed the expression levels of Runx2, COL I and ALP (Fig. 5A). Considering that VPA is an HDACi, the protein expression of HDAC1 and HDAC2 as class I HDACs as well as the levels of ac-H3 and ac-H4 were measured. The results indicated that the two HDACs were lowered in all experimental groups, including the DEX and DEX+VPA groups, while ac-H3 and ac-H4 were lowered in the DEX group and upregulated with the addition of 0.5 and 1 mM, but not 2 mM VPA (Fig. 5B and C). Similarly, the mRNA levels of Runx2 and COL I were significantly lowered by DEX and elevated by the addition of VPA. Although DEX increased ALP, even higher ALP levels were observed after the addition of VPA. Likewise, 1 mM VPA exerted the greatest effect on COL I, Runx2 and ALP gene expression. However, VPA did not rescue the lowered OCN expression under DEX treatment (Fig. 6).

VPA promotes bone mineralisation and improves mineralisation compromised by DEX treatment. Alizarin red
S staining was used to assess calcium deposition, a characteristic of late-stage osteogenic differentiation of BMSCs. Fewer calcium deposits in the dEX group compared with those in the control group were observed, while more deposits were visible after VPA treatment. In addition, supplementation of VPA in addition to DEX significantly improved the mineralisation of BMScs in comparison with dEX treatment alone (Fig. 7A).

The staining eluate was then harvested with 1 ml cetylpyridinium chloride and the absorbance values were measured at 560 nm. The quantified results indicated the same trend as that of the microscopic views presented (Fig. 7B).

**VPA increases serum ALP in rats treated with MP.** Within the 3 weeks of the animal experiment, serum ALP activity in the MP group was highest in the first week. At 1 week, it was also significantly higher than that in the MP+VPA group. However, ALP activity in the MP group waned in the following 2 weeks and became lower than that in the control. Of note, with the addition of VPA, ALP activity was rescued to the level of the control in the second week, while this effect was attenuated but still significant in the third week (Fig. 8).

**VPA improves subchondral bone formation under MP treatment.** The trabecular changes in the subchondral area of the femoral heads were detected by micro-CT scan at 6 weeks after the first injection of MP. A total of 11 rats in the MP group exhibited visible osteonecrosis of the femoral head in the micro-CT images, while only 2 rats in the MP+VPA group had obvious osteonecrosis (Fig. 9A). The BMD of the rats in the MP group was 0.27±0.01 g/cm³, which was significantly lower than that in the control group (0.61±0.01 g/cm³), while supplementation with VPA significantly increased the BMD of the area to 0.43±0.01 g/cm³. In addition, a similar effect was identified regarding the BV/TV and Tb.Th among these 3 groups of rats (Fig. 9B).

Similar to the micro-CT results, H&E staining of subchondral bone tissue revealed typical signs of osteonecrosis. Specifically, at low magnification, the MP group exhibited amorphous substance in the inter-trabecular spaces with few normal haematopoietic and fat cells on H&E staining. The trabeculae decreased in volume and became thinner in disarray and pale in staining in comparison with the control. At a magnification of x100, diffuse karyolytic and karyorrhexic nuclei of osteocytes were identified, and the marrow of amorphous matter was lacking normal cell nuclei, where there were mainly karyorrhectic and pyknotic speckles, revealing coagulation necrosis of the fatty hematopoietic tissue of the marrow. Furthermore, the ingrowth of fibrous tissue, a characteristic property of the repair process, was observed in this stage. However, with supplementation of VPA, no obvious necrotic area was present (Fig. 10A). The present study further assessed OCN expression as a marker of mature bone by IHC.
staining. The results indicated almost no positive area in the necrotic part of the subchondral bone after MP treatment, while the control and MP+VPA groups featured widespread OcN-expressing cells in the same area (Fig. 10B).

**VPA preserves the blood supply to the femoral head.** To detect the blood supply to the femoral head in the present study, micro-angiography of the femoral head was performed by perfusing the vessels with Microfil and observing them by micro-CT. The reconstructed 3D micro-CT images exhibited disrupted blood supply in the femoral head in the MP group, with vascularisation blocked around the femoral neck. However, the main vessel distributing to the femoral head in rats treated with MP+VPA was preserved (Fig. 11A). Correlation of micro-CT images and H&E staining revealed structural trabeculae in the vascularised area of the femoral heads. By contrast, the devascularised area displayed a lack of trabeculae and was referred to as the necrotic area. The vascularisation of the subchondral area of the femoral head was further detected by IHC analysis of VEGF. Nearly no staining for VEGF was observed in the subchondral bone area in the MP group, while a stained area was observed in the marrow cavity around the trabeculae in the MP+VPA and control groups (Fig. 11B).

**VPA promotes the osteogenic capacity of BMSCs in rats.** The BMSCs were subjected to osteogenic induction for...
3 days after being extracted from the proximal femurs of rats from each group and cultured for 3-5 passages. Western blot analysis was performed to measure the expression of Runx2 directly influenced by VPA as reported in previous studies (53), as well as ALP, its downstream protein. The results indicated that Runx2 and ALP expression were lowered in the MP group and improved with the addition of VPA (Fig. 12).

**Discussion**

ONFH is a common side effect of exogenous GC usage. One of the major underlying mechanisms is the inhibitory role of GCs in the osteogenesis of BMSCs, which influences the repair process of osteonecrosis. High doses of GCs may affect numerous processes involved in the differentiation of BMSCs. GCs may downregulate the expression of Runx2/Cbfa1, the key factor associated with osteogenic differentiation and maturation of osteoblasts (54). Several studies have indicated that GC antagonises Runx2 and subsequently compromises the expression of ALP, osteopontin and OCN in MSCs, resulting in inhibition of bone remodelling and fracture healing (19,55,56). The repair process is initiated when osteonecrosis takes place and is gradually outpaced by the proliferation of fibrous tissue, leading to incomplete reconstruction of the necrotic area devastated by the ceased blood supply. The ceased blood supply due to injury of vessels distributing to the femoral head is also attributed to failure of tissue reconstruction in the necrotic area. Consequently, the approach to prevent ONFH is more promising than to treat it, avoiding the loss of vessels that contributes to necrosis, which was applied in the present study.

VPA is an effective and widely used antiepileptic drug, which has been reported to influence osteogenesis as an
HdAci (26). HdAci have been demonstrated to enhance osteoblast differentiation in vitro (34, 53, 57) and new bone formation in vivo (37, 39). To the best of our knowledge, the present study was the first to indicate that VPA attenuates the inhibitive effect of GCs on the osteogenic differentiation of BMSCs, and that systemic injection of VPA prevents GC-induced ONFH in rats.

First, the effect of VPA alone on BMSc viability and apoptosis was observed. The results indicated that concentrations of VPA as high as 5 mM inhibited BMSc proliferation after 7 days of incubation, while VPA concentrations of 0.5 - 2 mM did not significantly influence BMSc proliferation. Hatakeyama et al (58) reported that (0.625 - 6.25 mM) VPA inhibited the proliferation of mouse embryonic stem cells. Lee et al (35, 38) also reported that the proliferation of adipose-derived stem cells decreased in a dose-dependent manner after VPA treatment at 0.4 - 10 mM, but the extent of inhibition was not as great in umbilical cord blood-derived stem cells, displaying a significant decrease only with 10 mM VPA. The decrease in cell proliferation in the presence of high concentrations of VPA may be associated with cell apoptosis or cell cycle arrest. Lee et al (35, 38) reported that the G2/M phase population of the cells was increased by VPA in a dose-dependent manner, while no increase in the sub-G1 phase was observed, demonstrating that the decrease in proliferation by VPA was not due to promoting apoptosis. In line with this, the present results also indicated that VPA concentrations as high as 5 mM inhibited the proliferation of BMSCs, but did not significantly influence BMSc proliferation.

Figure 7. Effects of VPA, DEX and DEX+VPA on osteogenic differentiation of BMSCs. (A) Alizarin red S staining of BMSCs exposed to VPA, DEX and DEX plus 1 mM VPA during osteogenic differentiation (magnification, x200). (B) Quantification of alizarin red S staining was performed by measurement of the cetylpyridinium chloride eluate and presented as absorbance values at 560 nm. Values are expressed as the mean absorbance value ± standard deviation. *P<0.05 vs. control group; #P<0.05 vs. DEX group. DEX, dexamethasone; BMSCs, bone marrow mesenchymal stem cells; VPA, valproic acid.

Figure 8. Analysis of serum ALP in Sprague Dawley rats. ALP activity was measured at 1, 2 and 3 weeks after injection of MP with or without VPA. Values are expressed as the mean absorbance value ± standard deviation. *P<0.05 vs. control group; #P<0.05 vs. MP group. VPA, valproic acid; ALP, alkaline phosphatase; MP, methylprednisolone.
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... not significantly affect cell apoptosis. Accordingly, VPA may cause cell cycle arrest rather than cell apoptosis. Furthermore, cycle arrest is a prerequisite for cell differentiation, and thus, it is possible that VPA influences the osteogenic differentiation of BMSCs (58,59). However, in the present study, monotherapy with DEX significantly inhibited BMSC proliferation, while this effect was attenuated by co-treatment with 0.5-2 mM VPA. In a cell apoptosis assay, the percentage of apoptotic cells increased under DEX treatment, but improved after addition of 5 mM VPA. DEX has been reported to promote apoptosis of several types of stem cell (60-63). However, VPA may be able to hamper this negative effect of DEX by inducing cell cycle arrest, which requires further study. Taken together, the results of the present study suggested that low concentrations of VPA (0.5-2 mM) do not significantly influence BMSC proliferation and attenuate the negative effect on proliferation exerted by DEX. The underlying mechanisms may include a reduction of DEX-induced cell apoptosis by VPA.

Next, the effect of VPA on osteogenic differentiation of BMSCs was assessed. The results indicated that VPA alone promoted ALP, COL I, Runx2 and OCN mRNA expression. Specifically, BMSCs treated with 1 mM VPA for 7 days exhibited the highest osteogenesis-associated gene expression, while all concentrations of VPA significantly elevated these genes after 14 days of incubation, with the exception of Runx2. Runx2 expression was gradually reduced along with osteoblast maturation, and mature osteoblasts do not contain significant amounts of Runx2 protein (64). Hence, it is possible that VPA accelerated the osteogenic differentiation process of BMSCs, leading to a reduction of Runx2, but an elevation of COL I, ALP and OCN, all of which are expressed in mature osteoblasts. Similar results were obtained for the protein expression of ALP, COL I and Runx2, as well as for ALP activity. VPA at a concentration of 1 mM was selected due to its efficacy in the mineralisation assay, resulting in the promotion of calcium nodule deposition. All of the present results indicated that VPA promotes osteogenic differentiation of BMSCs.

DEX has been reported to inhibit osteogenic differentiation and bone formation in vitro and in vivo (19,55). The present study indicated that DEX inhibited COL I, Runx2 and ALP protein expression after osteogenic induction of BMSCs for 7 days. This inhibition was abrogated or attenuated by co-treatment with 0.5 and 1 mM of VPA for Runx2 or ALP, respectively. In addition, 0.5 mM VPA reversed the inhibitory effect of DEX on COL I, resulting in expression levels above those in the control group. Similar results were obtained for the mineralization of hBMSCs, manifesting in a protective effect of VPA on calcium deposition on BMSCs in the presence of DEX. Likewise, the gene expression of Runx2 and COL I was downregulated by DEX and reversed by the addition of VPA, and ALP expression exhibited the highest level with the addition of 1 mM of VPA. However, the lowered gene expression of OCN was not improved with the addition of VPA. Rimando et al (65) used high concentrations of DEX (10^{-5} M) for long-term treatment (7 days) of BMSCs and identified glucocorticoid receptor (GR)-HDAC6 complex as the repressor of the promoter region of OCN, leading to the downregulation of OCN mRNA. However, VPA inhibits class 1 HDACs more efficiently than class II HDACs, e.g., HDAC5 and -6 (27). It is possible that VPA at concentrations of 0.5-2 mM is not sufficient to inhibit HDAC6 to improve OCN mRNA expression. Consistent with the results of previous studies (66-71), the present study also indicated that VPA and DEX significantly inhibit HDAC1 and -2 expressions in BMSCs. Previous studies have demonstrated that GR elements are located in the promoter of the HDAC2 gene and that DEX treatment for 7 days reduced GR expression (66,67). Accordingly, HDAC2 expression may be lowered by DEX. DEX also affects histone acetylation. DEX acts by binding...
to GR, which, upon activation, translocates to the nucleus and either increases or decreases gene expression (68). DEX has been reported to inhibit Runx2 expression by recruiting HDAC1 to the Runx2 promoter, which then mediates the

Figure 10. Histological analysis of paraffin sections of the femoral heads. (A) Representative images of H&E-stained coronal sections of femoral heads in each group. The rectangle refers to a part of the subchondral bone area (magnification, x50 and x100). At x100 magnification, the ellipses indicate the necrotic area, including karyolysis, karyorrhectic osteocytes, marrow necrosis and fibrous invasion. (B) Representative immunohistochemical staining images for OCN in the coronal sections of femoral heads in each group (magnification, x100). OCN, osteocalcin; VPA, valproic acid; MP, methylprednisolone.

Figure 11. Evaluation of blood supply and vascularisation preservation in femoral heads. (A) Representative 3-dimensional micro-angiography images of femoral heads in each group processed by micro-computed tomography (magnification, x25). (B) Representative images of immunohistochemical staining for VEGF in the femoral heads of each group (magnification, x100). VEGF, vascular endothelial growth factor; VPA, valproic acid; MP, methylprednisolone.
deacetylation of histone H4 and downregulates Runx2 expression (69). In addition, the complex of DEX and GR recruits HDAC2 to the complex of nuclear factor-κB and activator protein 1 and reduces the associated inflammatory cytokine expression (70,71). The results of the present study also indicate that the levels of ac-H3 and ac-H4 were lower with DEX treatment alone than with the addition of VPA. Although DEX lowered the expression of HDAC1 and -2, as VPA did, the present results indicated that DEX did not have the same inhibitory effect on HDAC activity as VPA did. This may be the reason for DEX inhibiting osteogenic differentiation, and the addition of VPA may improve or reverse this inhibitory effect. As stated above, DEX inhibits osteogenic differentiation by binding to the promoter of Runx2 with GR, recruiting HDACs and bringing transcription to an end. Consequently, VPA has a beneficial effect on osteogenic differentiation inhibited by DEX, which may be explained by the inhibition of HDACs and rendering of histone acetylation, giving rise to initiation of Runx2 expression.

The present study hypothesized that VPA has a beneficial effect against GC-induced ONFH. In a previous study, Xu et al (35) reported that pre-treatment of mouse adipose-derived stem cells with VPA for 2 days produced an increased amount of bone formation after injection into bone defect areas. Rashid et al (39) demonstrated that intraperitoneal injection of VPA for 1 week accelerated the healing of maxillary bone cavities. To the best of our knowledge, no previous in vivo experiment has been performed to assess the preventive effect of VPA against GC-induced ONFH. GC-induced ONFH was successfully achieved in rats by injection of 20 mg/kg MP on 3 consecutive days over 3 weeks. Previously, animal models of ONFH have been established using a combination of 20 mg/kg MP and 2 mg/kg lipopolysaccharide (72-74), as LPS causes vasculitis, which occurs in most autoimmune diseases (75). However, not all patients receiving GC treatment are diagnosed with autoimmune diseases, e.g. those with spinal cord injury. Accordingly, it is better to establish an animal model of ONFH by induction with MP only to study GC-induced ONFH. The GC-induced rat model of ONFH has been successfully established in previous studies (49,50). In the present study, this ONFH rat model exhibited characteristics of classically-defined necrosis on histology with karyolytic and karyorrhectic osteocytes; it also featured necrotic marrow filled with acellular substance or fibroblasts invasion, and sinusoidal vessels devoid of red blood cells, demonstrating that devascularisation and necrosis had occurred in the femoral head. Of the rats that had received MP injection, a randomly selected 50% were simultaneously administered 300 mg/kg VPA once a day for 3 weeks with no intermittence, as administration of 250-300 mg/kg VPA has been reported to cause HDAC inhibition and improve functional recovery in stroke (76), nerve injury (77), retinal ischaemic injury (78) or survival of haemorrhagic shock (79). However, it remains elusive whether 300 mg/kg of VPA is the optimal dose, e.g. whether larger doses would increase the efficacy or whether smaller doses would suffice, particularly in long-term treatment periods of ≥3 weeks. Therefore, the dose-dependent effect on the prevention of GC-induced ONFH will be assessed in a future study.

In the present study, blood samples were collected from the experimental rats for measurement of serum ALP activity, which reflects the process of bone formation in general. The results indicated that MP+VPA attenuated the suppression of ALP activity caused by MP during the second and third weeks. VPA treatment enhanced the serum ALP levels, which may explain for how VPA attenuates the inhibitory effect of MP on ALP activity (80). However, serum ALP is an indicator of bone formation rather than osteonecrosis progression (81-83). The bone density of the femoral head was then measured, and histological and angiographic analyses were performed. Overall, 11 out of 15 rats that had received MP treatment presented with ONFH, but it only occurred in 2 out of 15 rats subjected to MP+VPA treatment. The results of the micro-CT scanning indicated that VPA yielded a greater preservation of trabeculae and bone volume in animals treated with MP (MP+VPA group). On the contrary, no normal-shaped trabeculae were observed in the subchondral area, which was only filled with a silt-like substance in single treatment of MP. On histological analysis, the femoral heads exhibited no apparent necrotic area in the group subjected to VPA treatment, with evenly distributed expression of OcN along the trabeculae, indicating that VPA treatment preserves a relatively large amount of mature bone under MP challenge. The major blood supply into the femoral head was also preserved in the VPA group, and as in the control group, an area of VEGF-expressing tissue was observed in the marrow cavity, indicating that VPA may be able to protect vessels as well. A further study on the effect of VPA against MP- or DEX-induced vessel deficits will be performed in the future. Finally, BMSCs were collected from rats in each group and western blot analysis indicated that the protein expression of

Figure 12. Expression of proteins associated with osteogenesis in rat BMSCs. The protein expression of Runx2 and ALP in rat BMSCs was detected by western blot and image analyses. Values are expressed as the mean ± standard deviation (n=3). *P<0.05 vs. control group; **P<0.05 vs. MP group. VPA, valproic acid; MP, methylprednisolone; BMSCs, bone marrow mesenchymal stem cells; Runx2, runt-related transcription factor 2; ALP, alkaline phosphatase.
Runx2 and ALP was higher in the MP+VPA group compared with that in the MP group, indicating that the osteogenic capacity may have been improved by co-treatment with VPA, as indicated in the in vitro experiments. It may also be possible to prevent ONFH by improvement of the osteogenic capacity of BMSCs.

In summary, the present study reported that VPA enhanced the osteogenic differentiation of BMSCs, and attenuated the negative effect of GCs on BMSC proliferation, apoptosis and osteogenic differentiation. To the best of our knowledge, the present study was also the first to apply VPA as a preventive agent against GC-induced ONFH. The present results demonstrate that VPA is a promising drug for preventing ONFH, possibly via the enhancement of the osteogenic differentiation of BMSCs.

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Availability of data and materials

The analyzed data sets generated during the study are available from the corresponding author on reasonable request.

Authors' contributions

DZ was responsible for designing the concept, acquisition of data, analysis and interpretation and manuscript writing. CZ and YF contributed to the design, analysis and interpretation. YC analysed and interpreted the data and animal experiments. JY, ST and SG were involved in the experimental method optimisation. ZW contributed to the Micro-CT scanning and analyses. Each author read and approved the final version of the manuscript.

Ethical approval and consent to participate

The present study was performed according to the principles of the Helsinki Declaration, and written consent was obtained from each patient. The experimental procedures were approved by the Ethical Review Board of Shanghai Jiaotong University Affiliated Sixth People’s Hospital (Shanghai, China). Patient consent was provided before any procedures were performed.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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