Local delivery of simvastatin maintains tooth anchorage during mechanical tooth moving via anti-inflammation property and AMPK/MAPK/NF-κB inhibition

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Abstract
Simvastatin (SMV) could increase tooth anchorage during orthodontic tooth movement (OTM). However, previous studies on its bone-specific anabolic and anti-inflammatory properties were based on static in vitro and in vivo conditions. AMPK is a stress-activated kinase that protects tissue against serious damage from overloading inflammation. Rat periodontal ligament cells (PDLCs) were subjected to a serial of SMV concentrations to investigate the optimization that promoted osteogenic differentiation. The PDLCs in static and/or tensile culturing conditions then received the proper concentration SMV. Related factors expression was measured by the protein array, real-time PCR and Western blot. The 0.05UM SMV triggered osteogenic differentiation of PDLCs. The inhibition of AMPK activation through a pharmacological approach (Compound C) caused dramatic decrease in osteogenic/angiogenic gene expression and significant increase in inflammatory NF-κB phosphorylation. In contrast, pharmacological activation of AMPK by AICAR significantly inhibited inflammatory factors expression and activated ERK1/2, P38 MAPK phosphorylation. Moreover, AMPK activation induced by SMV delivery significantly attenuated the osteoclastogenesis and decreased the expression of pro-inflammatory TNF-α and NF-κB in a rodent model of OTM. The current studies suggested that SMV could intrigue intrinsic activation of AMPK in PDLCs that promote attenuate the inflammation which occurred under tensile irritation through AMPK/MAPK/NF-κB Inhibition.

Keywords
AMPK, animal study, MAPK, mechanical stress, periodontal ligament cells, simvastatin
mechanically induced physical movement raises great attention for shortening the treatment cycle and reducing the patients ‘ in- 
compliance caused by pain and discomfort, by either accelerating 
local movement rate or precisely increasing tooth anchorage in the 
alveolar.3 Existing biological therapy includes chemical methods 
(cytokines, hormone, drugs, growth factors and other biological sub-
stances) and gene therapy, among which the most potential transla-
tional approach is the locally administrated meditations.4

Simvastatin (SMV), as the HMG-CoA reductase inhibitor, is a cho-
lesterol-lowering statin drug.5 Recently, its protective bone anabolic 
and anti-resorptive properties have been unveiled and lead to a wide 
application in large-size bone defect regeneration, therapeu-
tic management of osteoporosis and periodontitis correction.6-8 According 
to clinical trials observation, SMV intake was beneficial for chronic 
periodontitis control for its function in inflammation limitation and 
periodontal tissues repair.9-11 Animal studies have shown that SMV 
when administered systemically resulted in increased tooth anchor-
age and reduced root resorption.12 To be mentioned, the pocket 
injection of 1.2% SMV gel decreased space re-opening after ortho-
dontic space closure in human anterior teeth.13 We thus assume a 
bone anabolic property of SMV on periodontal tissue under a bio-
mechanically induced inflammatory condition.

Adenylate-activated protein kinase (AMPK) is a silk and thre-
onine-protein kinase which mainly coordinates the metabolism of 
energy.14,15 Lately, researchers discovered its roles in autophagy inducement, inflammation control, cancer metastasis and autoim-
mune inhibition.16-22 Our group also found that AMPK-α1 knockout 
(AMPK-α1−/−) mice presented larger inflammatory periodontal bone 
defects and expressed higher levels of inflammatory cytokines than 
wild-type (WT) mice.23 Interestingly, OTM is a biomechanical pro-
cess accompanied by tension side bone regeneration and pressure 
side bone resorption, which indicates the AMPK activation in the in-
flammation-mediated osteogenesis under biomechanical irritation.24

Current in vitro observations grounded on the static-culturing 
periodontal ligament cells (PDLCs), which led to big inconsistency 
from the actual stress state of the PDLCs during OTM. In this study, 
we focused on the anti-inflammation effect of SMV on tensile side of 
the orthodontic moving tooth in a rodent model. An in vitro dynamic 
culturing system was also applied to mimic the stressing condition 
in OTM to explore the potential mechanism concerning the AMPK 
regulation.

2 | MATERIALS AND METHODS

2.1 | Animal for OTM model

Twelve-week-old male Sprague Dawley (SD) rats of 250 g ± 15 g 
were obtained from the Tongji Medical College Animal Center. 
All animal procedures were approved by the Animal Care and 
Experiment Committee of Tongji Hospital, Tongji medical college, 
Huazhong University of Science and Technology. Animals were 
housed in specific pathogen-free (SPF) condition at an ambient 
24-26°C temperature, 55%-60% humidity under a 12:12-hour 
light/dark cycle. Adequate measures were implemented to mini-
mize pain or discomfort of animals during all procedures. The study compliance with the ARRIVE guidelines.

2.2 | Acquisition and cultivation of human PDLCs

Ten human healthy pre-molars were collected from four patients (18-
25 years old) undergoing orthodontic treatment in the Department 
of Stomatology, Tongji Hospital. Informed consent was signed by all 
patients according to the Tongji Hospital, Huazhong University of 
Science and Technology Institutional Review Board Approval (IRB 
ID: 20140401) under the principles of the Declaration of Helsinki. 
After phosphate buffered saline (PBS) washing, the periodontal liga-
ment tissue was dissected from the mid-third portion of the roots 
using a sterile NO. 11 blade (Golden Circle Medical Supplies Co., 
Ltd.). The tissue explants were carefully placed on the bottom of a 
six-well culture plates and covered by a sterilized glass before add-
ing Dulbecco’s modified Eagle’s medium (DMEM, Gibco BRL) sup-
plemented with 10% foetal bovine serum (FBS, Gibco Biocult Co), 
100 U/mL penicillin and 100 mg/L streptomycin (Invitrogen Life 
Technologies).25 The explants were kept in a 5% CO2 incubator at 
37°C. Cells at passage 3-5 were adopted for the following tests.26

2.3 | The optimization of SMV concentration

Periodontal ligament cells were seeded into 24-well plates at a density 
of 4.0 × 104 cells/mL. One day later, six concentrations of SMV 
(0, 0.01, 0.05, 0.1, 0.5 and 1UM) (Sigma-Aldrich) in supplemented 
DMEM were offered. The cell numbers in each hole were evaluated 
on days 1, 4, 7 and 10 (Multisizer, Beckman Coulter Inc). PDLCs were 
seeded in 6-well plates at a density of 5 × 104 cells/well and cultured 
in the six concentrations prepared in osteogenic culture medium 
(supplemented DMEM with 50 mg/mL ascorbic acid, 10 mmol/L 
b-glycerophosphate and 10−8 mol/L dexamethasone, Sigma). After 
7- and 14-day inducement, ALP staining (Beyotime Biotech, Jiangsu, 
China) and semi-quantitative analysis (Sigma) were performed. The 
total protein content was determined using a protein assay kit (Bio-
Rad). The osteogenic gene expression of RUNX2, VEGF, OPN and 
OCN was measured by RT-PCR on days 1 and 7. Finally, after 4 weeks 
inducement, fixed PDLCs were marked by the Alizarin Red staining 
27 and the Von Kossa staining.28 Osteogenic medium (0UM SMV) 
was used as control. All experiments were performed in triplicate.

2.4 | Application of tension force system

External mechanical irritation was stimulated with a Flexcell FX-
5K® tension system (Flexcell International Corp.). PDLCs were 
plated onto six-well Bioflex plates (Flexcell International Corp.) at 
a density of 1 × 105 cells/well. When the culture reached 80%-90%
confluence, they were stretched with 10% equibiaxial strain at 0.1 Hz for 24 hours.29 PDLCs were assigned to six groups as:

(i) C-CON: No tension and no SMV (osteogenic culture medium only);
(ii) C-SMV: No tension + 0.05UM SMV;
(iii) T-CON: Tension only;
(iv) T-SMV: Tension + 0.05UM SMV (given 10 minutes before the stress loading);
(v) T-Compound C: Tension + 0.05UM SMV + 1UM Compound C (Compound C were given 10 minutes before the SMV dosing);
(vi) T-AICAR: Tension + 0.05UM SMV + 1UM AICAR (AICAR were given 10 minutes before the Compound C dosing).

2.5 | Protein array

The human protein membrane array (QAH-PDD-1, Ray Biotech) simultaneously profiles 20 proteins were in quadruplicate, and the supernatant of group i, ii, iii and iv was pulled together for measurement.30 The cytokine array membranes were blocked for 30 minutes and for 2 hours in the collected supernatants. After washing, the membranes were incubated with biotin-conjugated antibodies (1:250 dilution, 1 mL per array membrane) at room temperature for 2 hours and washed again. The horseradish peroxidase-conjugated streptavidin solution (1:1000, 2 mL) was added and incubated for 2 hours followed by a third wash step. Proteins were detected by the secondary antibodies (Abcam) incubation for 2 hours at 37°C, the membranes were subjected to the enhanced chemiluminescence for signal intensity quantification (Bio-Rad).

2.6 | Real-time Quantitative PCR

The total cellular RNA was extracted using the TRizol one-step method (Invitrogen Life Technologies) and reverse-transcribed to cDNA using a cDNA Synthesis kit (TaKaRa Bio). Primers were synthesized commercially (Shengong Co. Ltd) and were shown as follows: GAPDH, 5′-GACCTGACCTGCGTCTA-3′ and 5′-AGGAGTTGGTGCTGCGTGT-3′; OCN, 5′-CTCACAACCTGCGCCCTATT-3′ and 5′-GCCGGGCTTCCTCACAATT-3′; RUNX2, 5′-CATTCACTCTCGCTTTC -3′; OPN, 5′-ACTGATTTTCCCACGGACCT -3′; BMP-2, 5′-CAGAAAC GATTGGATGGCA -3′; TNF-κB, 5′-CGACTATCTCGACTTTGC -3′ and 5′-GGTTGAGGGTGTCTGAAGGA -3′; VEGF, 5′- AGGGCAGAAATCATACGGAAGT-3′ and 5′-AGGGTGTCCTGATTGATGGCA -3′; TGF-β, 5′-CCGACTATCTCGACTTTGC -3′ and 5′-GGTTGAGGGTGTCTGAAGGA -3′; NF-κB, 5′-GCACCTCAGAGACCTCTTACCT -3′; IL-6, 5′-GCCGGGCTCTCGCTTTCAC -3′; AMPK-α, 5′-ATGTTTGTGTTGATGGAACTTACGT -3′; ADP-α, 5′-GCCGGGCTCTCGCTTTCAC -3′; IL-6, 5′-GCCGGGCTCTCGCTTTCAC -3′.

The genes expression was evaluated using a real-time PCR kit (TaKaRa Bio). GAPDH was used as the housekeeping gene. The 2-ΔΔCT method was used to quantify the expression level. All tests were repeated in triplicate.

2.7 | Western blot analysis

Total proteins of the cells were extracted (RIPA lysis buffer, Sigma) and measured (BCA Protein Assay Kit, Pierce Biotechnology). Protein homogenates were run on SDS-PAGE gel (Beyotime) and transferred onto a PVDF membrane (Millipore). After 5% milk blocking, the membranes were incubated overnight in primary antibodies: AMPK (1:1000; Beyotime), P-AMPK (1:1000; Beyotime), p-p65 (1:2000), and p65 (1:2000), ERK1/2 (1:2000), p-ERK1/2, p38 MAPK and p-p38 (Abcam) at 4°C. GAPDH served as the internal reference. After secondary antibodies (Abcam) incubation for 2 hours at 37°C, the membranes were subjected to the enhanced chemiluminescence for signal intensity quantification (Bio-Rad).

2.8 | Pre-treatment of drug injection and Orthodontic model establishment

Four groups of pre-conditioning were randomly given to all animals:
(a) Control group, 70 mL saline once three days via submucosal injection 2 mm under the gingival papilla between the maxillary right first (M1) and second (M2) molar; (b) SMV group, 70 mL 100 mg/kg SMV; (c) Compound C group, 10 mg/kg Compound C was given 10 minutes before SMV administration; and (d) AICAR group, 10 minutes after the Compound C administration, AICAR (30 mg/kg) was given. A total of 3 times dosing were completed before orthodontic appliance fixation.31 The orthodontic wire was a titanium-nickel alloy measuring 0.228 mm in diameter and 14 mm in length.32 After intraperitoneal injection of 3.5 mg/100 g pentobarbital, the buccopalatal grooves of the M1 and M2 were enlarged with a diamond bur (NO. 145, Shofu) on the occlusal surfaces to deep enough to fit the spring wire. The site was dried, etched with 65% phosphoric acid (Shofu) for 20 seconds, rinsed with water and dried. The tips were brought together and maintained at a distance of 3 mm by a circular frame to deliver an initial force of 30 g. The springs were seated into the occlusal grooves and bonded with dental resin (Shofu). The frame was removed to activate the spring after bonding. The rats were then allowed to recover from anesthesia and returned to their cages.

At the end of 1 week active tooth movement, the orthodontic appliances were removed. Animals were euthanasia and perfused with 10% buffered formalin. The maxillae were carefully isolated and trimmed into single blocks together with three right upper molars before storage in 4% neutral formaldehyde solution at 4°C overnight.33
2.9 | Micro-CT measurement

The alveolar samples were assessed using a micro-CT system (μCT-80, Scanco Medical, Bassersdorf, Switzerland) as previously described. The microfocus of the X-ray source had a spot size 7 mm and maximum voltage 36 kV. A cylindrical region of interest (ROI) in all the samples mesial to M2 with an axis depth/length of 600 μm (100-700 μm below the M2 root furcation) and a diameter of 700 μm was selected for measurement of bone volume fraction (BV/TV, %), and the distance between the ROI and the mesial root of M2 was 200 μm. The amount of tooth movement was assessed on two-dimensional (2-D) sagittal sections taken through the centres of the M1 and M2 (the image plane that showed the most structure of the distobuccal and mesial root) and measured at the interproximal heights of contour between the most mesial point of the M2 crown and the most distal point of the M1 crown.

2.10 | Histological and histomorphometric assays

All samples were decalcified in 10% EDTA before embedding. The specimens were cut into 5 μm sections and prepared for H&E staining. The histomorphometric analysis was performed under the bright-field setting on a light microscope (Leica-microsystems) on six samples per group. On three randomly selected images (200 × magnification) on the tensile side of the distal root of the first molar per sample, the cell nuclei and Howship’s lacunae was calculated (image J software 5.0). Immunohistological staining of TNF-α and NF-κB (1:500; Abcam) was performed to locate the protein expression on six samples per group (n = 6). On three randomly selected images (40 × magnification) on the tensile side of the distal root of the first molar per sample, positive cell numbers were calculated. Finally, the sections were treated with a mixture of a tartaric acid solution and acid phosphatase substrates (Sigma). The TRAP labelled cells were calculated in the corresponding region of H&E-stained sections.

2.11 | Statistical analysis

The experimental data were presented as the mean ± standard derivation. Statistical analysis in this study was performed using one-way ANOVA and the SNK post hoc test based on the normality of the distribution with the SAS 8.2 statistical software package. Values of $P < .05$, $P < .01$ were considered statistically significant.
RESULTS

3.1 The 0.05UM SMV induced stronger osteogenic differentiation of rat PDLCs in vitro

Human PDLCs were successfully obtained for in vitro tests. To figure out a proper SMV concentration for in vitro tests, we first compared the PDLCs proliferation rate in an increasing concentration gradient of the drug. The cell number gradually increased in all groups as the inducement time prolonged. The proliferation rate of the low SMV concentration groups (0.01 and 0.05UM) was similar to the control group. The proliferation rate of the high SMV concentration groups (0.1, 0.5 and 1UM) slowed from day 4 and reached the platform period after day 7 (Figure 1A). Next, the ALP staining was performed on day 7. For the 0.05UM group, the purple-red colour stained area reached the maximum. As the SMV concentration increased, the colour of ALP became lighter (Figure 1B). Semi-quantitative detection of ALP was further performed on day 14. Simply, the ALP activity of 0.05UM group was significantly higher than the control group (*, \(P < .05\)), while 1UM group demonstrated the lowest ratio (*, \(P < .01\); Figure 1C).

We then selected two concentrations of 0.05 and 0.1UM for further assessments. The two-step RT-PCR showed that OCN expressions were similar among two testing groups, while the 0.05UM group stimulated higher expressions of RUNX2 (1-day induction) and OPN (7-day induction) (*, \(P < .05\); **, \(P < .01\)). The expression of angiogenic factor VEGF in both testing groups significantly increased on the 1st day and reached the highest level in the 0.05UM group on the 7th day (**, \(P < .01\)). The change of gene expression level suggested that 0.05UM SMV induced the strongest osteogenic and angiogenic differentiation in PDLCs (Figure 1D). Finally, we tested the calcified nodule formation in control and 0.05UM SMV groups. Both Alizarin Red and Von Kossa staining showed that the 0.05UM SMV induced more calcified nodules in larger size than the control group (Figure 1E).

3.2 SMV promoted osteogenesis on the tensile side of periodontium during OTM via anti-inflammatory effect

SMV shows protective bone anabolic and anti-resorptive properties.\(^6,11\) We here focused on identifying the intrinsic factor(s) responsible for the therapeutic effects of SMV in increasing tooth anchorage. We took advantage of protein array technology and scanned 20 pro- and anti-inflammatory factors in the co-stimuli of tension and SMV-sensitized PDLCs. We found that IL-6 was the most pronounced cytokine induced by PDLCs under the co-stimuli (Figure 2A,B). To detect the potential effect of AMPK during SMV-induced anti-inflammation, we subsequently measured the expression levels of p-AMPK/AMPK and p-P65/P65 (NF-\(\kappa\)B) in the cells. Our data demonstrated that increased AMPK was negatively connected with IL-6 secretion (Figure 2C). Also, we confirmed that IL-6 mRNA were correspondingly reduced after indicated treatment and AMPK mRNA levels were correspondingly elevated by the co-stimuli (Figure 2D,E).

The intervention of AMPK signalling was analysed by the administration of the AMPK inhibitor, Compound C and the AMPK activator, AICAR. RUNX2, BMP-2 and VEGF are important factors during osteogenesis. For BMP-2 and VEGF, the mechanical tension significantly increased their expression compared to the static culture condition (\(P < .01\)). Unlike VEGF expression, which tended to increase over time, BMP-2 expression remained high within 24 hours. For RUNX2, mechanical stress did not induce obvious augmentation of its expression level as did in BMP-2 and VEGF. However, the addition
of SMV significantly lifted the expression level in all the three osteogenic-related genes ($P < .05$; Figure 3A-C). For pro-inflammatory factors, the SMV administration increased its expression under a static environment ($P < .05$). This change was reversed under mechanical culturing condition. The pre-conditioning of Compound C and AICAR further proved this trend ($P < .01$; Figure 3D). For NF-κB, the inhibition of AMPK significantly elevated its expression in all the time-points ($P < .01$). This change was corrected after the administration of AMPK activator. However, the tensile culture condition did not show statistic influence on RUNX2 expression (Figure 3E). On the other hand, the expression level of AMPK significantly increased in tensile condition at 12 hours and 1 day. And SMV collaboratively increased the expression level at 6 hours ($^*, P < .05$) and 12 hours ($^{**}, P < .01$). This alteration could be inhibited by Compound C (Figure 3F). Meanwhile, there was no significant difference in the expression of AMPK-α ($P > .05$).

3.3 | SMV promoted osteogenesis on the tensile side of periodontium during OTM via AMPK/MAPK/NF-κB inhibition

To further clarify the effect of AMPK on SMV stimulated PDLCs in tensile culture condition, we detected the MAPK/NF-κB signalling by Western blot (Figure 4A). The results showed that 0.05UM SMV treatment down-regulated the folded expression of p-ERK, p-P38 and p-P65 ($^{##}, P < .01$ vs T-SMV group) (Figure 4B). This change would be altered by Compound C ($@@, P < .01$ vs T-COMP group) and further corrected by AICAR, which showed a negative correlation with the p-AMPK level. Finally, we detected the expressions of TNF-α and IL-1 in supernatants derived from indicated treatment by ELISA. The amounts of secreted TNF-α and IL-1 shown similar change patterns as the NF-κB p65 phosphorylation level as induced by the co-stimuli of tension and SMV (Figure 4C). Specifically, SMV decreased the inflammatory factors level ($^{**}, P < .01$ vs T-CON group).

3.4 | Surgical procedures and gross observation

In general, a total of 24 SD rats received orthodontic appliance ($n = 24$) placement. During the pre-conditioning period of 7 days, all rats were in good general health and showed any adverse reactions to the injection procedures. No orthodontic appliance was lost in the mechanical stimulation period (Figure 5). After the appliance installation, sites were observed with limited inflammatory responses.
Micro-CT assessment

The maxillae samples were examined by micro-CT before the following process. Massive bone resorption was noted around the distal root of the first maxillary molar in the control group, while the height of the bones around the distal roots in the SMV group was basically maintained at the bifurcation position (Figure 6A). The moved distance of first maxillary molar in control group was $497.13 \pm 63.28 \mu m$, which was significantly higher than the SMV group $27.56 \pm 31.15 \mu m$ and AICAR group $172.45 \pm 55.05 \mu m$ (**, $P < .01$). There was no statistical difference between the control group and the Compound C group $417.91 \pm 50.36 \mu m$ ($P > .05$; Figure 6B). The BV/TV percentage was consistent with 3D reconstruction observation. The control group $63.84 \pm 7.36\%$ was the lowest, while the SMV group $79.49 \pm 3.38\%$ the maximum (**, $P < .01$). The treatment of Compound C reduced the BV/TV to $60.55 \pm 4.56\%$, and the addition of AICAR significantly increased the ratio to $72.8 \pm 6.08\%$ (**, $P < .01$; Figure 6C).

3.6  Histological observation

Rest specimens were decalcified for paraffin sections preparing. H&E staining showed that the pressure side of the periodontium was narrower than the tension side (Figure 7A). Observing the periodontal ligament on the tension side of the distal root (within the yellow square), we found that the thickness of periodontal ligament was larger in the control group, their arrangement was more chaotic, and the nucleus appeared to be significantly longer than the other.
three groups. Meanwhile, the change of periodontal space in the SMV group was limited. There was no obvious bone resorption site on the tension side, as indicated by the yellow arrows in the other three groups. Little influx of inflammatory cells was detected in the SMV group. The histological findings of Compound C group and AICAR group were between the former two groups. In terms of cell arrangement, the Compound C group was more similar to the control group. This trend was proved by the histomorphometric assessment of cell nuclei numbers (Figure 7B). Specifically, there were more cells in the control group (311.67 ± 32.58) than the other three groups, as 162.17 ± 19.86 for SMV group, 252 ± 34.02 for Compound C group, and 182.5 ± 31.46 for AICAR group.

The immunohistological staining showed that the NF-κB and the TNF-α expression was strongly positive on the tension side of the moved molar in the control group. These two factors were both weakest expressed in the SMV group. However, their uniformly scattered coloration could be seen in the periodontal ligament of the Compound C group. The stained cells percentage of the two factors demonstrated identical changes among the four groups, which was similar to the histomorphometric results of the H&E staining (Figure 7B). Specifically, there were more cells in the control group (311.67 ± 32.58) than the other three groups, as 162.17 ± 19.86 for SMV group, 252 ± 34.02 for Compound C group, and 182.5 ± 31.46 for AICAR group.

The periodontal ligament is a layer of connective tissue between the root of the tooth and the alveolar bone. PDLCs contribute to the regeneration of periodontal ligament, cementum and alveolar bone and are recognized as the histological basis for OTM. However, previous studies explaining

4 | DISCUSSIONS

Control of stabilization during orthodontic treatment is considered of significant importance since it helps avoid undesirable tooth movements that may occur as a consequence of the reaction forces applied to move teeth. Pharmacological therapy of clinically accepted medicines is a shortcut to complement the orthodontic appliance which unavoidably leads to a higher level of pain and discomfort. Recent studies have found that SMV is effective in inhibition of AMPK/MAPK/NF-κB pathway by local administrated SMV could enhance tooth anchorage in alveolar during OTM.

The periodontal ligament is a layer of connective tissue between the root of the tooth and the alveolar bone. PDLCs contribute to the regeneration of periodontal ligament, cementum and alveolar bone and are recognized as the histological basis for OTM. However, previous studies explaining
the bone-specific anabolic property of SMV mostly based on observation of static-culturing osteoblasts.\textsuperscript{46,47} Researchers found that SMV could promote the expression of ALP and OCN from increased secretion of BMP-2.\textsuperscript{48,49} Concerning the cell origination for the metabolism of periodontium and alveolar during OTM, we applied human PDLCs for the in vitro tests. We found that the
low concentration SMV could elevate the expression levels of BMP-2 as well as VEGF in PDLCs under a tensile condition in vitro (Figures 1 and 3). This was consistent with previous findings that SMV could lift the expression of BMP-2 and VEGF mRNA in osteoblasts after local application of SMV.50 Since bone formation is a coupling process involving osteogenesis and angiogenesis, this result proves the bone anabolic property of SMV on PDLCs in the tensile condition. Interestingly, there were significant differences among the gene expression levels of BMP-2, VEGF, TNF-α and AMPK on most time-points between the tension and the static groups, which highlighted the importance of tension force system application.

NF-κB is a well-known transcription factor associated with inflammation and bone resorption.51 Many studies reported orthodontic mechanical stress could trigger the NF-κB pathway activation, which mediated the expression of pro-inflammatory genes such as TNF-α and IL-6, as proved in our histological findings.52 Therefore, inhibition of NF-κB activation is an important process in increasing tooth anchorage. On the other hand, ERK and p38 MAPK belong to the MAPKs signalling pathway, which played a central role involved in inflammatory response and cell survival.53,54 For certain, recent studies indicated the phosphorylation of p38 and ERK1/2 MAPKs was activated in stress-stimulated PDLCs.55 We then verified the effect of SMV on the MAPKs signalling pathway, and our results revealed SMV markedly inhibited stress-stimulated ERK and p38 MAPK phosphorylation in PDLCs, and these manifestations had significant changes after treatment with Compound C, the inhibitor of AMPK. The pre-condition of AICAR significantly reversed the inhibition trend and increased the pro-inflammatory factors expression level. Furthermore, we tested the expression level of TNF-α and IL-1. Similar changes were identified under the Compound C and AICAR treatment. AMPK is an energy sensor that is activated by several types of stresses and has been described as a negative regulator of NF-κB pathway mediating SMV bone-protective effect on the enhancement of tooth anchorage during OTM.

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CONFLICT OF INTEREST

The authors confirm that there are no known conflicts of interest associated with this publication.

AUTHOR CONTRIBUTIONS

Xu Qin: involved in conceptualization, supervision and project administration. Babak Baban involved in methodology and wrote—reviewed and edited. Guangxun Zhu helped in software. Jing Mao validated the study. Lianyi Xu involved in formal analysis. Xiaojuan Sun and Lianyi Xu involved in investigation, resources and data curation. Guangxun Zhu and Jing Mao wrote and prepared the original draft. Lianyi Xu and Xu Qin acquired funding.

DATA AVAILABILITY STATEMENT

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

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