Research Article

Synergistic Effects of Orbital Shear Stress on In Vitro Growth and Osteogenic Differentiation of Human Alveolar Bone-Derived Mesenchymal Stem Cells

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Cellular behavior is dependent on a variety of physical cues required for normal tissue function. In order to mimic native tissue environments, human alveolar bone-derived mesenchymal stem cells (hABMSCs) were exposed to orbital shear stress (OSS) in a low-speed orbital shaker. The synergistic effects of OSS on proliferation and differentiation of hABMSCs were investigated. In particular, we induced the osteoblastic differentiation of hABMSCs cultured in the absence of OM by exposing hABMSCs to OSS (0.86–1.51 dyne/cm²). Activation of Cx43 was associated with exposure of hABMSCs to OSS. The viability of cells stimulated for 10, 30, 60, 120, and 180 min/day increased by approximately 10% compared with that of control. The OSS groups with stimulation of 10, 30, and 60 min/day had more intense mineralized nodules compared with the control group. In quantification of vascular endothelial growth factor (VEGF) and bone morphogenetic protein-2 (BMP-2) protein, VEGF protein levels under stimulation for 10, 60, and 180 min/day and BMP-2 levels under stimulation for 60, 120, and 180 min/day were significantly different compared with those of the control. In conclusion, the results indicated that exposing hABMSCs to OSS enhanced their differentiation and maturation.

1. Introduction

The stem cell is a complex microenvironment combining an extracellular matrix, cell-to-cell interactions, and other factors such as growth factors, physical factors, and various cytokines. Stem cells are exposed to high Ca²⁺ concentrations and a variety of autocrine, paracrine, and endocrine signals (extrinsic factors) and they are attached to the ECM through integrin receptors [1–10]. Many researchers have already reported an influence of cell growth and differentiation with the use of physical stimulators. Also, we have previously reported the in vitro osteogenic effects of cell stimulation on human alveolar bone-derived mesenchymal stem cells (hABMSCs) using a simple rocking culture method [11].

Thereby, we ascertained that the shear stress on hABMSCs could significantly enhance cell migration, proliferation, and differentiation. Our previous study motivated us to identify other methods for simple cell stimulation.

Thus, we designed orbital shear stress (OSS), which considered another possible cell stimulation method with the concept that flow patterns within intraoral fluid in the mouth are circular. There have been several studies of the effects of OSS on cellular behaviors [2–12]. Steady laminar flow can induce the expression of many genes and proteins in stem cells. The physical forces have profound effects on the cytoskeleton and extracellular matrix. These cellular components are essential in maintaining the integrity of stem cells. In particular, gap junctions are membrane channels that mediate...
the cell-to-cell movement of ions and small metabolites [5, 6, 13]. Some studies have reported that the Cx43 which is involved in gap junction channel activity in cells, including stem cells, might be induced by OSS to regulate cell growth and differentiation [6–10]. It has been suggested that the mode of cell-cell communication might be of particular importance in the skeleton, where various signals mediate gap junction communication and connexin biology in the bone [8–10, 14, 15]. Above all, one mechanism of cell-cell interaction is direct cell-cell communication via gap junctions, which are transmembrane channels that allow for the continuity of cytoplasm between communicating cells [13–15]. Cellular signaling occurs through distinct events: binding of stimuli secreted from neighboring cells or cell junctions and release in response to stimuli. Such signals affect cellular migration, growth, and differentiation [16–18].

The purpose of our study, therefore, was to investigate the synergistic effects of OSS on in vitro growth and osteogenic differentiation of hABMSCs for tissue engineering applications.

2. Materials and Methods
2.1. Cell Culture. hABMSCs were collected at the Intellectual Biointerface Engineering Center, Dental Research Institute, College of Dentistry, Seoul National University. hABMSCs were placed in 35 mm culture dishes at a density of $1.0 \times 10^4$ cells/cm$^2$ and cultured for 5 and 10 days. Cells were cultured in α-minimum essential medium (α-MEM) containing 10% fetal bovine serum (FBS, Welgene Inc., Republic of Korea) and 10 mM ascorbic acid (L-ascorbic acid) and antibiotics (Antibiotic-Antimycotic solution, Gibco) at 37°C in a humidified atmosphere of 5% CO$_2$ (Steri-Cycle 370 Incubator, Thermo Fisher Scientific, USA). The cells were then incubated with osteogenic medium (100 nM dexamethasone, 50 μg/mL of ascorbic acid, and 10 nM of β-glycerophosphate; Sigma) for 10 days. The induction culture medium was
Figure 3: Continued.
changed every second or third day. The proliferation and osteogenic differentiation of the cells were examined after exposure to each OSS.

2.2. Stimulation Treatment of OSS and Experimental Device. OSS was applied to confluent cell cultures using a low-speed orbital shaker (Benchmark Scientific, USA). The OSS was calculated using the following equation (1) [2]:

$$\tau_w = a \times \sqrt{\rho \times u \times (2 \times \pi \times f)^3}, \quad (1)$$

where $\tau_w$ is shear stress, $a$ is the orbital radius of rotation of the shaker, $\rho$ is the density of the culture medium, $\mu$ is the viscosity of the medium, and $f$ is the frequency of rotation [2]. In this study, we calculated the values of shear stress at temporal points as shown in Figure 1 as 5, 10, 20, 30, and 40 rpm (revolutions per minute). The equation expresses constant magnitude of shear [17–19]. Figure 1 indicates temporal points for calculating values of OSS. The Reynolds number was calculated as $\omega R^2/\nu$, where $\omega$ is the rotational speed of the orbital shaker, $R$ is the radius of rotation of the orbital shaker (17.5 mm), and $\nu$ is the kinematic viscosity ($1.012 \times 10^{-6}$ m$^2$/s).

hABMSCs were exposed to OSS (0.86–1.51 dyn/cm$^2$) with plate on the orbital shaker (Reynolds number of 121). There were six treatment groups, stimulated for 10, 30, 60, 120, and 180 min/day.

2.3. Cell Viability, DNA Analysis, and In Vitro Cell Migration Assay. hABMSC proliferation was measured by WST-1 assay (EZ-Cytox Cell Viability Assay Kit, Daeillab Service Co., Ltd.). The formazan dye produced by viable cells was quantified by a multiwell spectrophotometer (Victor 3, Perkin Elmer, USA), measuring the absorbance of the dye solution at 460 nm. DNA concentration was quantified by fluorometry using the CyQUANT Cell Proliferation Assay Kit (Invitrogen), and the $\lambda$ Fluorescence was measured using a Cytofluor II fluorescence multiwell plate reader with excitation of 485 nm and emission of 530 nm. In vitro cell migration was assessed by the CytoSelect Wound Healing Assay according to the manufacturer’s protocols. Wound closure was measured by microscopy for up to 72 h, and photographs were taken. Cells were cultured with or without OSS, and cell morphology was observed by phase-contrast microscopy (Nikon TS100, Japan). hABMSCs were stimulated with exposure to OSS for 72 h, and the control was not exposed to OSS.

2.4. Measurement of Mineralized Nodule Formation. All cells except control cells were exposed to OSS for 10 days. Nodule formation was checked routinely by phase contrast microscopy. The presence of mineralized nodules (calcium deposition) was determined by staining with Alizarin red, as described [20]. The ethanol-fixed cells and matrix were stained for 1 h with 40 mM Alizarin red-S (pH 4.2) and extensively rinsed with water. After photography, the bound stain was eluted with 10% (wt/vol) cetylpyridinium chloride, and the Alizarin red staining in the samples was quantified by measuring absorbance at 544 nm (Victor 3, Perkin Elmer, USA). Cells were fixed with 4% (wt/vol) formaldehyde in PBS for 15 min. And the cells were incubated in 5% (wt/vol) silver nitrate (Sigma-Aldrich, USA) for 1 h under ultraviolet light condition, followed by incubation in 5% (wt/vol) sodium thiosulfate (Sigma-Aldrich, USA) for 5 min. Last, the wells were rinsed with distilled water twice and air-dried, and mineralization images were captured using an optical microscope.

2.5. Reverse Transcriptase-Polymerase Chain Reaction Analysis. Reverse transcriptase-polymerase chain reaction analysis (RT-PCR) was used to measure the expression of various osteogenic factors. After 10 days in OSS culture, total RNA was isolated with TRIzol reagent (Invitrogen) and used to

![Figure 3: In vitro cell migration as representative optical microscopic images with OSS groups compared to static culture (A), indicating that stimulation groups exposed at 10, 30, and 60 min/day were significantly different ($^*P < 0.05$) among groups (B) ($n = 3$).]
Figure 4: Representative optical fluorescence microscopy images of hABMSCs cultured for 5 days in static conditions (a1–d1) or at 10 min/day (a2–d2), 30 min/day (a3–d3), 60 min/day (a4–d4), 120 min/day (a5–d5), and 180 min/day (a6–d6) by OSS without OM; cell nuclei (a1–a6), actin filaments (b1–b6), gap junction (Cx43, c1–c6), and merged images (d1–d6) of the fluorescence stains. Fluorescence images showed more intense observation in OSS groups without OM compared to those in control (arrows: cell direction).
Figure 5: Representative optical fluorescence microscopy images of hABMSCs cultured for 5 days in static conditions (a1–d1) or at 10 min/day (a2–d2), 30 min/day (a3–d3), 60 min/day (a4–d4), 120 min/day (a5–d5), and 180 min/day (a6–d6) by OSS without OM; cell nuclei (a1–a6), actin filaments (b1–b6), OCN (c1–c6), and merged images (d1–d6) of the fluorescence stains. Fluorescence images showed more intense observation in OSS groups without OM compared to those of control.
synthesize cDNA using a first-strand cDNA synthesis kit (Invitrogen) according to the instructions of the manufacturer. The human primers used in this study are listed in Table 1. RNA was extracted from the cells 10 days after the addition of differentiation media. These extracts were subjected to RT-PCR analysis of Runx2 (runt-related transcription factor 2), COL1 (collagen type I), OCN (osteocalcin), OPN (osteopontin), SMAD1, and GAPDH (glyceraldehyde-3-phosphate dehydrogenase) as the positive control. The products were separated by electrophoresis on a 1% agarose gel (SeaKem ME; FMC Bioproducts) and visualized by ultraviolet-induced fluorescence. Expression levels of gene areas were measured using Image J 1.45s (National Institutes of Health).

2.6. Fluorescence Microscopy and Confocal Laser Scanning Analysis. Cells were washed in phosphate-buffered saline (PBS, Sigma-Aldrich, Milwaukee, WI, USA), fixed in a 4% paraformaldehyde solution (Sigma-Aldrich, Milwaukee, WI, USA) for 20 min, and permeabilized with 0.2% Triton X-100 (Sigma-Aldrich, Milwaukee, USA) for 15 min. Cells were incubated with TRITC-conjugated phalloidin, anti-osteocalcin, its secondary antibody (Cat. no. AB10911, Millipore), and DAPI (Millipore, Billerica, MA, USA) for 1 h to stain actin filaments, focal contacts, and nuclei, respectively. Cytoskeleton organization was visualized using an actin cytoskeleton and focal adhesion staining kit (FAK100; Millipore, Billerica, MA) according to the manufacturer’s instruction. Cells were mounted in glycerol/buffer on a glass slide after extensive washing with PBS. Images of labeled cells were obtained by a Confocal Laser Scanning Microscope (Carl Zeiss, LSM710).

2.7. ELISA Assay. To measure the levels of vascular endothelial growth factor (VEGF) and bone morphogenetic protein-2 (BMP-2), we used an ELISA kit with specific antibodies (Quantikine human VEGF and Quantikine human BMP-2 immunoassays, R&D Systems, USA). The culture supernatants were collected to quantify the levels of VEGF and BMP-2 produced from hABMSCs in vitro after 5 days. The assay protocol was performed according to the instructions of the manufacturer. Each sample was measured in triplicate.
Figure 7: RT-PCR analysis of cell cultures between stimulus conditions (from 10 min/day to 180 min/day) and static culture for 10 days (A). RNA was extracted from the cell cultures at 10 days after the addition of differentiation media. These extracts were subjected to RT-PCR analysis with Runx2, COL1, OCN, OPN, SMAD1, and GAPDH as the positive control. Expression levels (B) of COL-1 (a), Runx2 (b), OPN (c), and OCN (d) at 10 days were significantly higher in OSS stimulus conditions on cells than those in control. OSS groups exposed at 10, 30, 60, and 120 min/day were significantly different (* P < 0.05 and ** P < 0.001) among groups.
Figure 8: Continued.
2.8. Statistical Analysis. Statistical analysis was carried out using the SAS Statistical Analysis System for Windows v9.3 (SAS Institute, Inc., Cary, NC, USA). Statistical significance between control and treatment groups was compared with t-test, two-way ANOVA, and Duncan’s multiple range tests at *P < 0.05. The data are reported as the mean ± standard deviation.

3. Results and Discussion

3.1. Cell Viability and Growth Are Enhanced by OSS. Cell metabolic viability of hABMSCs was measured using optical density and WST-1 according to Figure 1. The cell viability of the 40rpm group when exposed at 10 min/day increased more than 10% over those of 10 and 20rpm groups (Figure 2(a)). DNA concentration (Figure 2(b)) as a percentage of initial hABMSC measured using the CyQuant cell proliferation with OSS stimulation (40rpm). Specifically, we observed that 40 rpm and OSS stimulation of 10, 30, and 60 min/day induced greater cell metabolic activity. OSS groups had higher cell metabolic viability than control group. We have indicated that OSS in short term stimulated the cell growth and proliferation in vitro whereas hABMSCs proliferation was associated with decrease with exposure to laminar shear stress for long term. The in vitro hABMSCs migration result was shown in Figure 3. The in vitro cell migration shown in the optical microscopic images (A) showed that the difference between OSS and static culture groups was significant, and the OSS groups exposed at 10, 30, and 60 min/day also showed significant differences (*P < 0.05) (B). Based on the cell growth, migration assay, and DNA proliferation, hABMSCs proliferated significantly (about 20%) under OSS condition of 10 and 30 min/day when compared with that of control. We could consider that OSS does produce laminar shear stress on the cell-seeded culture dish, which is related to increased proliferation.

3.2. Enhanced Gap Junction (Cx43) and OCN in the Absence of Osteogenic Media (OM). Figure 4 showed representative confocal images of hABMSCs cultured for 5 days in static conditions (a1–d1) or at 10 min/day (a2–d2), 30 min/day (a3–d3), 60 min/day (a4–d4), 120 min/day (a5–d5), and 180 min/day (a6–d6) by OSS in the absence of OM; cell nuclei (a1–a6), actin filaments (b1–b6), gap junctions (Cx43, c1–c6), and merged images (d1–d6) of the fluorescence stains. The Cx43 indicated more intense staining in OSS groups in the absence of OM compared with the control. Gap junction communication is important in bone cells [21], where the channels are involved in mechanical transmission [22–24], induction of cytokines in osteoblasts [25], and coordination of hormonal responses [26, 27]. In osteoblast-like cells in vitro, Cx43 is the dominant connexin subtype and likely plays an important role in normal skeletal development [28–30]. Many studies have demonstrated a mutual relationship between cell growth and the expression of tissue-specific genes during mineralization [31–33]. In this respect, we could assure that gap junction (Cx43) was accelerated by OSS compared with that of control in the absence of OM. Figure 5 presented representative optical fluorescence microscopy images of hABMSCs cultured for 5 days in static conditions (a1–d1) or...
Figure 9: Continued.
at 10 min/day (a2–d2), 30 min/day (a3–d3), 60 min/day (a4–d4), 120 min/day (a5–d5), and 180 min/day (a6–d6) by OSS with OM; cell nuclei (a1–a6), actin filaments (b1–b6), OCN (osteocalcin, c1–c6), and merged images (d1–d6) of the fluorescence stains. Fluorescence images of OCN ascertained that gap junction (Cx43) was affected by OSS even in the absence of OM compared to those of the control, suggesting that the shear stress stimulates the cells mechanically and alters cellular functions.

3.3. Gene Expression of Osteoblastic Differentiation Markers. We investigated alkaline phosphatase activity (ALP) of hABMSCs stimulated with OSS for 7 days (Figure 6). To induce osteoblast differentiation in MSCs, the culture medium was supplemented with osteogenic agents, including L-ascorbic acid, β-glycerophosphate, and dexamethasone [34–37]. L-Ascorbic acid enhances collagen synthesis and upregulates adenosine triphosphatase and ALP activity, and β-glycerophosphate serves primarily as a source of inorganic phosphate ions [2, 12, 34–40]. The results of RT-PCR analysis of the cell cultures between stimulus conditions (from 10 min/day to 180 min/day) and static culture for 10 days (A) were shown in Figure 7. Expression of genes associated with the osteoblastic differentiation was examined using RT-PCR to investigate the effect of the stimulation with OSS on gene expression at 10 days. Expression levels (B) of COL-1 (a), Runx2 (b), OPN (c), and OCN (d) at 10 days were higher in OSS stimulation conditions on cells than those in control. Stimulation groups of 10, 30, 60, and 120 min/day were significantly different ($^* P < 0.05$ and $^{**} P < 0.001$) among groups. In particular, the 30 min/day group showed high expression levels of OPN and OCN.

3.4. Osteoinduction of hABMSCs by OSS in the Absence of OM. Figure 8 indicated representative optical microscopic images of osteoinduction of hABMSCs after Alizarin red staining of cells treated with static conditions (a1, b1) or by stimulation for 10 min/day (a2, b2), 30 min/day (a3, b3), 60 min/day (a4, b4), 120 min/day (a5, b5), and 180 min/day (a6, b6) on 5 days or 10 days, respectively. The cells induced with OSS treatment during 10, 30, and 60 min/day were intense compared with those of control. Representative microscopic images after von-Kossa staining are also shown (Figure 8) for static condition (c1, d1) or for stimulation for 10 min/day (c2, d2), 30 min/day (c3, d3), 60 min/day (c4, d4), 120 min/day (c5, d5), and 180 min/day (c6, d6). The cells stimulated with OSS (C) showed significant differences in osteoinduction ($^* P < 0.05$ and $^{**} P < 0.001$) among groups. Interestingly, mineral induction via OSS indicated the cells were moving outwards. We considered that one of the migration roles of hABMSCs used in this study could be controlled to the desired migration direction as well as external force on cells. Ultimately, osteogenic differentiation promotion on hABMSCs was induced by the simple orbital shear shaker as physical cues of the microenvironment.

3.5. Effects of OSS Induction with OM on Osteogenic Differentiation. Figure 9 shows representative optical microscopic images of osteogenic differentiation of hABMSCs after
Figure 10: Representative optical fluorescence microscopy images of hABMSCs cultured for 5 days in static conditions (a1–d1) or at 10 min/day (a2–d2), 30 min/day (a3–d3), 60 min/day (a4–d4), 120 min/day (a5–d5), and 180 min/day (a6–d6) by OSS with OM; cell nuclei (a1–a6), actin filaments (b1–b6), gap junction (Cx43, c1–c6), and merged images (d1–d6) of the fluorescence stains. Fluorescence images showed more intense observation at OSS groups compared to those of control (arrows: cell direction).
Figure 11: Representative optical fluorescence microscopy images of hABMSCs cultured for 5 days in static conditions (a1–d1) or at 10 min/day (a2–d2), 30 min/day (a3–d3), 60 min/day (a4–d4), 120 min/day (a5–d5), and 180 min/day (a6–d6) by OSS with OM; cell nuclei (a1–a6), actin filaments (b1–b6), (OCN, c1–c6), and merged images (d1–d6) of the fluorescence stains. Fluorescence images showed more intense observation in OSS groups with OM compared to those of control (arrows: cell direction).
Figure 12: Quantitative analysis of VEGF and BMP-2 proteins was performed with conditioned medium between OSS treatment and control group. VEGF protein of OSS group was significantly different (30 and 120 min/day; *P < 0.05, 10, 60, and 180 min/day; **P < 0.001). BMP-2 protein of OSS induction group also showed significant differences (60, 120, and 180 min/day; *P < 0.05). Overhead brackets with asterisks indicated significant differences between groups.

Alizarin red staining of cells treated with static conditions (a1, b1) or by stimulation for 10 min/day (a2, b2), 30 min/day (a3, b3), 60 min/day (a4, b4), 120 min/day (a5, b5), and 180 min/day (a6, b6) on 5 or 10 days, respectively. Cells treated with OSS for 10, 30, and 60 min/day were intense compared to those of control. Representative images of von Kossa-stained hABMSCs treated with static conditions (c1, d1) or by stimulation for 10 min/day (c2, d2), 30 min/day (c3, d3), 60 min/day (c4, d4), 120 min/day (c5, d5), and 180 min/day (c6, d6) were also shown. Cells treated with OSS induction for 10, 30, and 120 min/day showed significant differences (C, *P < 0.05) among groups.

Several studies have shown effects of OSS on in vitro growth of cells and experimental apparatus that can provide quantifiable shear stress, involving inducing a rotating flow. In particular, endothelial cells could experience shear exerted by the flow of blood, causing them to become aligned and elongated with the direction of flow and to undergo other biochemical changes [2, 41]. More importantly, orbital shakers provide oscillatory flow, somewhat like the pulsing fluid movement in the human vasculature system [41–43]. In the biopharmaceutical development, agitation varies from simple mixing of components to increasing mass transfer, to deliberate introduction of agitation-related stresses and to accelerate protein degradation in screening experiments [44]. Based on these facts, we ascertained that the cell stimulation could mediate a strong effect on cell proliferation and differentiation.

3.6. Analysis of Gap Junction (Cx43) and OCN with OM. Figure 10 shows representative optical fluorescence images of hABMSCs cultured for 5 days under static condition (a1–d1) or under stimulation for 10 min/day (a2–d2), 30 min/day (a3–d3), 60 min/day (a4–d4), 120 min/day (a5–d5), and 180 min/day (a6–d6) by OSS with OM; cell nuclei (a1–a6), actin filaments (b1–b6), gap junction (Cx43, c1–c6), and merged images (d1–d6) of the fluorescence stains. Fluorescence images indicated more intense staining in cells treated with OSS compared with those of controls. Interestingly, the gap junction (Cx43) fluorescence stains of cells cultured with PM (proliferation media) were more strong compared with cells cultured with OM.

Figure 11 demonstrates representative optical fluorescence images of hABMSCs cultured for 5 days under static conditions (a1–d1) or under stimulation for 10 min/day (a2–d2), 30 min/day (a3–d3), 60 min/day (a4–d4), 120 min/day (a5–d5), and 180 min/day (a6–d6) by OSS with OM; cell nuclei (a1–a6), actin filaments (b1–b6), OCN (osteocalcin, c1–c6), and merged images (d1–d6) of the fluorescence stains. The fluorescence images presented more deep staining in the cells stimulated by OSS with OM compared to those of control.

Gap junction intercellular communication is the most direct way of achieving such signaling, and gap junction communication through connexin-mediated junctions, in particular connexin 43 (Cx43), plays a major role bone development [45]. Given the important role of Cx43 in controlling development and differentiation, especially in bone cells, controlling the expression of Cx43 may provide control over cell-to-cell communication and may help overcome some of the challenges in craniofacial tissue engineering [45–47].

Connexins play a major role in response to many mechanical, electrical, chemical, and hormonal stimuli and help regulate cell homeostasis as well as calcium signaling and differentiation [26, 48, 49]. Therefore, controlling fluid flow like OSS can also potentially induce the opening of Cx43 channels in osteocytes and other bone cells allowing for
enhanced cell-cell communication and bone formation [50–52]. The major premise of functional tissue engineering is to provide physical cues to cells as a means of enhancing proliferation, differentiation, and tissue formation. Physical stimulation of cells in monolayer enhances gap junction function [23, 49, 50, 53, 54]. Thus, the mechanisms with relation to the enhanced tissue regeneration that are subjected to physical stimulation may be gap junction mediated [55].

3.7. Quantitative Analysis of BMP-2 and VEGF Proteins. Quantitative analysis of VEGF and BMP-2 proteins was performed with conditioned medium. VEGF protein of cells in the OSS induction group showed significant differences, as shown in Figure 12 (30 and 120 min/day; \( P < 0.05 \), 10, 60, and 180 min/day; \( P < 0.001 \)). BMP-2 protein in the OSS induction group also indicated significant differences (60, 120, and 180 min/day; \( P < 0.05 \)). The interaction between VEGF and BMP-2 is dependent on the ratios of angiogenic and osteogenic factors. Osteogenic factors such as BMP-2 can stimulate osteoblasts, and VEGF can modulate vascularization [56–58].

4. Conclusions

In this study, we investigated the synergistic effects of OSS on in vitro growth and osteogenic differentiation of hABMSCs. The results indicated that OSS stimulation treatment has an important effect on the activation of mechanotransduction. Cell viability stimulated for 10, 30, and 60 min/day increased by about 10% compared with that of the control. We also found an effect of OSS on the osteogenic differentiation of hABMSCs with OM and without OM, respectively. The OSS groups with OM and without OM that underwent stimulation for 10, 30, and 60 min/day showed more intense staining compared with the control. We also quantified VEGF and BMP-2 protein expression levels after stimulation for 10, 60, and 180 min/day and found that VEGF protein levels and BMP-2 protein levels after 60, 120, and 180 min/day were significantly different from levels measured in the control. In conclusion, this study showed that exposing hABMSCs to OSS stimulation enhanced cell differentiation and maturation.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

Acknowledgments

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