Modulating OPG and TGF-β1 mRNA expression via bioelectrical stimulation

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ABSTRACT

Background: Bone remodeling is a lifelong process that ranges from orthodontic tooth movement/alignment to bone damage/healing, to overall bone health. Osteoprotegerin (OPG) and transforming growth factor β1 (TGF-β1) are secreted by osteoblasts and participate in bone remodeling. OPG promotes bone remineralization and stabilization prominent in post-mechanical repositioning of the teeth in the dental alveolus. TGF-β1 participates in regulatory processes to promote osteoblast and osteoclast equilibrium. In the context of orthodontic tooth movement, post-treatment fixation requires additional, exogenous, stabilization support. Recent research showcases supplementary solutions, in conjunction to standard tooth fixation techniques, such as OPG injections into gum and periodontal tissues to accelerate tooth anchorage; however, injections are prone to post-procedure complications and discomfort. This study utilizes non-invasive bioelectric stimulation (BES) to modulate OPG and TGF-β1 as a novel solution to regulate bone remineralization specifically in the context of post-orthodontic tooth movement.

Purpose: The aim of this study was to investigate a spectrum of BES parameters that would modulate OPG and TGF-β1 expression in osteoblasts.

Methods: Osteoblasts were cultured and stimulated using frequencies from 25 Hz to 3 MHz. RT-qPCR was used to quantify changes in OPG and TGFβ-1 mRNA expression.

Results: OPG mRNA expression was significantly increased at frequencies above 10,000 Hz with a maximum expression increase of 332 ± 8% at 100 kHz. Conversely, OPG mRNA expression was downregulated at frequencies lower than 1000 Hz. TGF-β1 mRNA expression increased throughout all stimulation frequencies with a peak of 332 ± 72% at 250 kHz. Alizarin Red tests for calcium, indicated that mineralization of stimulated osteoblasts in vitro increased 28% after 6 weeks in culture.

Discussion: Results support the working hypothesis that OPG and TGF-β1 mRNA expression can be modulated through BES. Noninvasive BES approaches have the potential to accelerate bone remineralization by providing a novel tool to supplement the anchorage process, reduce complications, and promote patient compliance and reduce post-treatment relapse. Noninvasive BES may be applicable to other clinical applications as a novel therapeutic tool to modulate bone remodeling.

1. Introduction

Bioelectrical stimulation (BES) has been associated with tissue remodeling and repair within the body (Tyler, 2017) via distinct mechanisms (Levin, 2013) that make it attractive as a noninvasive therapy for many diseases. BES has the ability to produce specific tissue and cellular responses via manipulation of key electrical parameters like signal strength, frequency, pulse form, and duration, allowing for a broad depth of therapeutic possibilities.

Exogenous bioelectric stimulation has been implicated in the up- or down-regulation of various growth factors that, subsequently, elicit various cell-specific responses. This phenomenon, although still not clearly understood (Vander Molen et al., 2000), is particularly relevant in non-excitable cells where ion channels are highly involved in tissue homeostasis (Cervera et al., 2016). It has, therefore, been suggested by many clinical laboratories that BES may be a non-invasive approach for enhancing bone repair and reducing healing times following bone damage (Aleem et al., 2016; Bhavsar et al., 2019; Kuzyk and Schemitsch,
One particular treatment in which bone remodeling is heavily involved is orthodontic tooth movement (OTM). In OTM, attached or removable dental appliances are used to apply mechanical forces to reposition teeth. Applying mechanical forces to the teeth initiates bone remodeling cascades in alveolar bone and periodontal tissues (Davidovitch, 1991; Krishnan and Davidovitch, 2006). These cascades induce osteoclastic activity along the leading (compression) edge of the tooth that instigate bone resorption (Yokoya et al., 1997). Similarly, osteoclastic activity is induced along the trailing (tension) edge of the tooth—promoting bone deposition or remineralization (Garlet et al., 2007). The interrelationship between osteoclastic and osteoblastic activities can be characterized by, and correlated to, observed changes in three specific bone-modulating growth factors—receptor activator of nuclear factor-kappa B (RANK), its ligand (RANKL), and osteoprotegerin (OPG) (Walsh and Choi, 2014; Yamaguchi, 2009).

OPG is one of the key factors responsible for maintaining bone homeostasis (Theoleyre et al., 2004). Its concentration in relation to RANK, RANKL, and other molecules influences bone metabolism and governs orthodontic tooth movement by determining bone deposition and resorption patterns (Ikebuchi et al., 2018). RANK is a type I transmembrane protein expressed along osteoclast precursor membranes (Idriss and Naismith, 2000; Ito and Hata, 2004). When activated by RANKL, an osteoclastogenic factor is released by surrounding osteoblasts and these pre-osteoclastic cells differentiate into mature osteoclasts to promote bone resorption. OPG, a secreted osteoblast-derived RANK decoy receptor, can inhibit osteoclastogenesis by binding RANKL and preventing RANK activation (Baud-huin et al., 2013).

Transforming growth factor β1 (TGF-β1) is another molecule that is highly relevant and highly involved in regulating bone remodeling. TGF-β1 has multiple roles in bone formation. It enhances osteoblast proliferation (Kassem et al., 2000) and recruits osteoblastic precursors, or matrix-producing osteoblasts, to the region via chemotactic attraction (Lucas, 1989). During initial phases of osteoblastic differentiation, TGF-β1 enhances the production of extracellular bone matrix protein (Alliston et al., 2001) and cooperates with metalloproteins to regulate the differentiation of osteoblasts (Canalis et al., 2003).

While the exploration of TGF-β1 is still relatively nascent in OTM, the exploration of OPG has seen increased interest. For example, it has been found that the local injection of OPG directly into the gum and periodontal ligament can reduce molar movement and osteoclast numbers (Dunn et al., 2007). The injection delivery of OPG and its resultant tooth stabilizing effects, while not free from complications (Baxter et al., 2020), may provide novel pharmacological approaches to prevent undesirable tooth relapse following appliance or aligner removal post-OTM (Dunn et al., 2007; Li and Tang, 2009).

BES provides an alternative, noninvasive approach to tooth stabilization. By delivering key electrical sequences within biological tissues, physiological responses are triggered that mimic, enhance, and/or modulate biomolecular processes. Several studies demonstrate the ability of BES to improve bone remodeling and regeneration (Hess et al., 2012; Srirussamee et al., 2019; Zhou et al., 2019b) or to modify cell growth patterns in vitro (Eischen-Loges et al., 2018; Spadari et al., 2017). In vitro, the mineralization process is active in cultured osteoblasts, and mineral plaques can be observed after 4 weeks in culture (Aboushady et al., 2018; Blair et al., 2017). This cellular physiological activity in vitro could be correlated with the mineralization in bone, and we hypothesize that it can be enhanced during the induction of expression of OPG during BES.

Considering its therapeutic potential, we hypothesize that BES can be successfully used in vitro to upregulate or modulate OPG and TGF-β1, and that it can be also applied in local periodontal tissues to modify gene expression. Successful modulation may lead to improved, noninvasive, and novel orthodontic treatments. Furthermore, we hypothesize that modulation would be dependent on both frequency and stimulation strength. BES protocols of varying frequencies and voltages were applied to osteoblasts in culture to determine the effects on OPG and TGF-β1 mRNA expression.

2. Materials and methods

2.1. Cell culture

Murine osteoblasts (MmOsteo; MC3T3-E1 Subclone 4) were acquired and cultured in alpha Minimum Essential Medium (αMEM; Gibco, Gaithersburg, MD, USA) supplemented with 10% FBS. Cells were maintained between six and nine total passages per supplier protocols (ATCC, Manassas, VA, USA). Osteoblast dissociation was performed with Trypsin-0.25% EDTA (Caisson, Smithfield, USA) and cells were plated on Nunclon™ Delta 6-well MultiDishes (ThermoFisher, Waltham, MA, USA) with 5% Non-Essential Amino Acids (NEAA; Quality Biologicals, Gaithersburg, MD, USA) and 5% Penicillin-Streptomycin (ThermoFisher, Waltham, MA, USA). All cultures were maintained in a 5% CO₂ incubator at 37 °C.
2.2. Experimental design

Osteoblasts were grown on 6-well plates until they reached between 80% and 100% confluency. Cells were electrically stimulated for 30 min, immediately washed 3 times with PBS, and dissociated with trypsin for real-time polymerase chain reaction (RT-qPCR) analysis. After trypsinization, cells were collected, centrifuged, resuspended in PBS, and re-centrifuged. Supernatant was aspirated and the remaining cell pellet was immediately stored at -80 °C for subsequent storage.

2.3. Stimulation process

Bioelectric stimulation was applied to cultured osteoblasts in vitro using a commercially available constant voltage waveform generator RIGOL LXi 1022Z (Beaverton, OR, USA) via a 6-well stimulating plate interface (IONOPTIX, Westwood, MA, USA). To induce uniform electric fields in all stimulation chambers, 1.3 mL of DMEM solution was added to each well prior to BES signal application.

2.4. Cell viability and proliferation

To determine the effect of BES on cells, and to evaluate cell health, osteoblasts were plated, stimulated, and assessed for cell growth, shape, and viability. Density was controlled at 600,000 cells per well. Three wells received a 2.0 V, 500,000 Hz, biphasic stimulation at 50% duty for 30 min while the other three wells remained unstimulated as controls. After stimulation, micrographs were taken, and cells were observed for 24 h. After the observation period, cells were split using Trypsin-0.25% EDTA and 7000 cells were plated in 24-well plates. Cells were counted at 3 and 5 days after stimulation.

2.5. Gene expression

Gene expression was determined by extracting mRNA from osteoblasts and applying RT-qPCR. mRNA was extracted using Invitrogen PureLink RNA Mini Kit (Invitrogen, Carlsbad, CA, USA) to produce highly pure, intact mRNA. An ultraviolet absorbance ratio at 260:280 nm was measured using a NanoDrop (ThermoFisher, Waltham, MA, USA) to ensure high-quality mRNA and achieve optimal reaction performance. mRNA was stored at -80 °C and good laboratory practices were followed to prevent degradation by exogenous ribonucleases prior to RT-qPCR analysis.

For complementary DNA (cDNA) generation, mRNA was reverse transcribed using ThermoFisher Scientific Maxima H Minus First Strand cDNA synthesis kit (ThermoFisher, Waltham, MA USA). Samples were processed following the manufacturer’s guidelines. Genomic DNA was eliminated to ensure optimal gene expression profiling by incubating template RNA, 10× dsDNAse Buffer, dsDNAse, and nuclease-free water at 37 °C for 2 min. First-strand cDNA synthesis was performed by adding a mixture of oligo (dT) primer and random hexamer primer, 10 mM dNTP mix, nuclease-free water, 5× RT Buffer, and Maxima H Minus Enzyme mix into a reaction tube. The tube was incubated for 10 min at 25 °C followed by 15 min at 50 °C and 5 min. The synthesized cDNA was then stored on ice.

TaqMan qPCR primers and probes from Applied Biosystems® (Applied Biosystems, Foster City, CA, USA) for OPG and TGF-β1 (targets) and GAPDH (reference), were used to determine mRNA concentrations. GAPDH primers were chosen as endogenous controls based on relative expression levels (Chapman and Waldenstrom, 2015). The reaction mixture contained 2 μL of cDNA, 10 μL of TaqMan Fast Advanced Master Mix, 1.0 μL of TaqMan assay, and 7.0 μL of nuclease-free water. 2 μL of gene-specific primer were transferred into each well of a 384-well plate. 18 μL of the TaqMan assay mix were added into each cDNA-filled well. The PCR plate was then sealed with optical adhesive film, briefly centrifuged, and loaded into an Applied Biosystems QuantStudio3 Real-time PCR system (Applied Biosystems, Foster City, CA, USA). Denaturation took place at 95 °C for 10 min followed by 40 annealing cycles at 95 °C for 10s and an extension cycle of 60 °C for 20s. The measurement of fluorescence at each cycle allowed for direct detection of PCR products. Control experiments were performed and processed in parallel throughout the protocol and used within individual experiment sets to calculate change due to treatment (2^ΔΔCt). All RT-qPCR measurements were performed in triplicate and effectively used in analysis if the triplicate standard deviation was less than 0.50 Ct.

2.6. Mineralization

To determine mineralization changes in osteoblasts in vitro resulting from BES, we seeded (80% confluence) on 13 mm glass coverslips previously covered with collagen (US Biological, Collagen Type I, Rat C7510-18; 0.5 mg/mL in PBS) for 1 h and then washed out with PBS. Two coverslips were placed inside each well of a 6-well plate and three wells of the dish were used for control (not simulated). Alpha MEM complete media was replaced every 3-5 days for 6 weeks. After one week of seeding, cells were stimulated with 1000 kHz for 30 min. This

Fig. 2. Cell Growth Curves. Cell growth comparison between control (squares) and bioelectrically stimulated (circles) osteoblasts. Cells were counted 3 and 5 days after stimulation. No significant changes were observed in cell growth rates.
stimulation was repeated, two and four weeks after seeding. At the end of 6 weeks, calcium deposits were quantified using Alizarin Red S, an anthraquinone derivative, used to identify calcium in tissue sections. Calcium forms an Alizarin Red S-calcium complex in a chelation process, and the end-product is orange-red and birefringent.

Cells were fixed with alcohol 70% in deionized (DI) Water for 5 min. Control cells were not stimulated but followed the same feeding process for 6 weeks. Cells were washed with DI Water 3 times and Alizarin Red (2 g in 100 mL; pH 4.2) was added for 5 min. Coverslips were washed with Di Water, dried, and dehydrated with Acetone 100% Glass slides were placed in 24 well plates and covered with glycerin. Alizarin Red absorbance (405 nm) was measured for each coverslip using a Perkin Elmer EnSpire 2300 Multilabel reader.

2.7. Statistical analysis

A multiple factor polynomial regression was used to evaluate the main effect of frequency and voltage on log fold change in OPG and TGF-β1 expression due to treatment ($2^{\Delta\Delta Ct}$). Prior to gene expression analysis, outliers were identified and removed using scatter plot analyses. If significant, a one-sample $t$-test was performed to evaluate the log fold change due to treatment and determine its significance. The multiple comparisons were corrected using the Benjamini and Hochberg (1995) method. The generalized additive model was used to evaluate the modulation of the log fold change in OPG and TGF-β1 expression due to treatment ($2^{\Delta\Delta Ct}$) by either frequency or voltage and required cubic splines anchored by knots selected between data inflection points. All analyses were performed using RStudio (version 1.1.44, Boston, MA, USA) and R (version 3.4.4, Vienna, Austria) with packages: ggplot2,
Rmisc, ggplot2, forcats, splines. Alpha was set at 0.05 for all statistical tests.

3. Results

3.1. Effect of bioelectrical stimulation on viability and proliferation of osteoblasts

Post-stimulation findings indicated that phenotype appeared to be unchanged, adhesion characteristics remained consistent with controls, and cell shape did not vary when compared to pre-experiment observations. Under identical growth conditions, no observable changes were noted in cell viability after 30 min of stimulation at 1 mV and 500 Hz. Growth rates were not significantly different after 3 and 5 days, post-cell stimulation (Fig. 2).

3.2. Effect of bioelectrical stimulation on OPG mRNA expression

RT-qPCR analysis was used to determine changes in OPG expression via mRNA concentration analysis. OPG mRNA expression responded to changes in the frequency of electrical stimulations (4th order polynomial, \( p < 0.029 \)) by exhibiting consistent upregulation or down-regulation trends under controlled frequency conditions. Similar trends were not observed with voltage changes.

Fig. 3A shows mRNA fold change of OPG in response to specific stimulation signals. Fig. 3A (left axis) shows the logarithmic fold change of OPG. Values above zero reflect an increase in OPG mRNA expression compared to the basal expression level in control experiments while values below zero (an inverted bar element) reflect a decrease in mRNA expression. Fig. 3A (right axis) provides an alternative perspective by juxtaposing linear fold increases/decreases within the construct of the logarithmic scale. Circles along the curve indicate the predicted values for each frequency based on a general additive model with knots chosen between inflection points within the frequency domain defined at 20,
90, 250, 700, 2500 and 200,000 Hz (model fit adjusted $R^2 = 0.36$, $P < 0.001$) for OPG mRNA expression.

OPG study outcomes, shown in Fig. 3A, can be broadly divided into two regions—frequencies that downregulate mRNA expression and those that upregulate it. OPG mRNA expression is reduced at frequencies between 100 Hz and 1000 Hz, with a negative peak close to 500 Hz. Frequencies between 10,000 Hz and 500,000 Hz increase OPG mRNA expression with a peak at 100,000 Hz. A cut score was defined at 2500 Hz (sensitivity: 0.67; specificity: 0.86; area under the curve 66%) where a transition point was identified between OPG mRNA up- and down-regulation. Mixed results were observed for stimulations at and above 750,000 Hz. It is worth noting that the x-axis is not linear. Frequency selections were made to span the broad range of signals used in clinical practice and not on a strictly mathematical construct.

Voltages commonly applied in the clinical realm were chosen to determine the effect of voltage on OPG mRNA expression. Variability in OPG mRNA expression was observed, however, trends in upregulation and downregulation arising from different voltage applications showed no voltage dependence.

3.3. Effect of bioelectrical stimulation on TGF-β1 mRNA expression

RT-qPCR analysis was also used to determine changes in TGF-β1 expression via mRNA concentration analysis (Fig. 3B). TGF-β1 mRNA expression concomitantly responded to changes in signal frequency by, similarly, exhibiting consistent modulations in the upregulation throughout the frequency ranges tested (general additive model adjusted $R^2 = 0.29$, $P < 0.001$, with inflection points, knots at 750, 2500, 75,000 and 500,000 Hz). In contrast to OPG expression, there are no frequencies in the test range that lead to a downregulation of mRNA (i.e., values below the zero line). Only results from 0.1 V stimulations were considered as it had been previously determined, during OPG experiments, that gene expression regulation was not voltage dependent.

TGF-β1 study outcomes, shown in Fig. 3B, can also be broadly divided into two regions, albeit slightly different than those observed with OPG. TGF-β1 mRNA expression is increased at frequencies between 75 Hz and 10,000 Hz, with a positive peak close to 500 Hz and again at higher frequencies between 100,000 Hz and 1000,000 Hz with a peak at 250,000 Hz thereby producing a positive double-peak response to frequency changes. As with Fig. 3A, the x-axis of Fig. 3B is not linear and frequencies were selected to span the broad ranges used in clinical practice.

3.4. Effects of BES over mineralisation

Mineralization is a common phenomenon observed after maintaining osteoblast in culture (Aboushady et al., 2018; Blair et al., 2017; Magloire and Joffre, 1979). These calcium deposits can be observed after two or three weeks (Fig. 4B). After stimulating the cells for 30 min with $2 \text{ V-100 kHz}$ pulses, one week, two weeks and four weeks after plated, the absorbance of Alizarin Red for the stimulated cells ($n = 4$ cover slips) was 28% higher than in the control group after 6 weeks in culture (Fig. 4D) indicating an increase in calcium deposits. Unpaired t-student test indicates a significant difference with a confidence interval of 95%. ($n = 4$, $df = 6$ and $t = 2.80$, with a two-tailed $P = 0.0310$). As a control, calcium chloride and Alizarin red were added to well 1-4 to identify the red-orange reaction.

4. Discussion

There have been multiple studies investigating the effects of various OTM techniques that have resulted in the elucidation of several molecular mechanisms explaining the process and physiology of tooth movement (Huang et al., 2014). On the one hand, it has been clearly demonstrated that the expression of RANK and RANKL promote bone resorption via osteoclastic differentiation—initiating bone softening and subsequent tooth movement (Ikebuchi et al., 2018). On the other hand, the induction of tooth fixation, by way of increased OPG expression, stops bone softening by reducing osteoclastogenesis and promoting bone hardening (Baud’huin et al., 2013). Previous studies involving the injection of OPG directly into dental tissues have also proven successful (Baxter et al., 2020; Li et al., 2019; Li and Tang, 2009), although application can become stressful and painful to the patient during multiple treatments. This study documents that an alternative method, bioelectric stimulation, can increase the concentration of OPG and TGF-β1 mRNA expression within osteoblasts, leading to a potentially non-invasive alternative to increasing OPG expression without the associated risk and discomfort of repeat OPG injections. This developing approach may accelerate bone remineralization and mechanical tooth repositioning treatments that use attached or removable appliances.

Osteoblasts were stimulated in vitro with signals of varying voltage and frequency. Stimulated cells did not experience changes in growth rate nor were there significant alterations in cell morphology within 5 days (Fig. 2). This observation implies that BES targeted to bone remodeling specific pathways does not trigger deleterious effects such as programmed cell death within stimulated cells.

The results of this study demonstrate, for the first time, that OPG and TGF-β1 expression, as measured by mRNA concentration, can be directly modulated by bioelectric stimulation. Upregulation for OPG was consistently and significantly induced at frequencies between 10,000 Hz and 500,000 Hz while downregulation was influenced by frequencies in the range of 100 Hz to 1000 Hz. Contrary to OPG, TGF-β1 displayed consistent upregulation where specific frequency ranges were more significant than others i.e., 75 Hz – 1000 Hz, 100,000 Hz – 1000,000 Hz. These consistent phenomena indicate a novel, and potentially more compliant, therapy for tooth movement/stabilization protocols.

Outcomes also demonstrated that these results are not a general initiation of cellular gene expression, but that there are specific frequencies that can be applied to osteoblasts whereby the expression of OPG and TGF-β1 mRNA can be controlled, manipulated, and optimized. This novel finding implies that there are molecular pathways within osteoblasts, osteoclasts, and likely other cells that predictably respond to specific electrical stimuli.

Direct current stimulation has been shown to induce gene modulation (Mobini et al., 2017) where stimulation was provided for one to fourteen days. In this case, some molecules like Osteopontin and COL1A2 were upregulated. In other cases, bioelectrical stimulation of cultured cardiocytes induced changes in cellular phenotype (Genovese et al., 2008). Using microarray assays, multiple mRNAs were reported to be either upregulated or downregulated, but without distinguishing which genes were specifically involved in bone repair (Caputo et al., 2014). Comparatively, modulation of gene expression has also been reported after electromagnetic field stimulation (Chang et al., 2005; Hinsenkamp and Collard, 2011; Zhou et al., 2019a) with different results. More specifically, the release of insulin has been controlled through electrical stimulation in genetically engineered cells (Krawczyk et al., 2020) highlighting the therapeutic potential of bioelectric stimulation.

It is important to note the down regulation of OPG mRNA. Gene expression is primarily controlled at the level of transcription, largely because of binding of proteins to specific sites on DNA. Regulation of protein production is largely achieved by modulating access of RNA polymerase to the structural gene being transcribed.

At the DNA level, osteoblast gene expression can be upregulated by enhancers, which in turn are modulated by proteinic activators that increase the transcription of a particular gene expression (GM, 2000; Phillips, 2008). Specifically in osteoblasts, gene expression is regulated after differentiation by Runx2, the master regulator, as it can be regulated by phosphorylation, acetylation, and ubiquitination and some of its domains can mediate either transcriptional activation or repression through associations with co-activators and co-repressors (Jensen et al., 2010).
Runx2 is also regulated by parathyroid hormone, highly important in skeletal physiology (Bellido et al., 2003). Expression of mRNA for OPG can also be downregulated by testosterone and upregulated by estrogen. Therefore, there are multiple paths to follow for the reduction in expression of OPG as protein or its transcript mRNA.

There have been several attempts to describe the cellular mechanisms behind protein expression through BES. Calcium transmembrane relocation (Xu et al., 2009) and enzyme activation (e.g. alkaline phosphatase; Caputo et al., 2014) can all be influenced by BES. Furthermore, bioelectric regulation of gene expression seems to be indicative of a multifactorial phenomenon that depends on the cell type and intracellular milieu, thus, requiring future cellular and molecular investigation (Caputo et al., 2014; Srirussamee et al., 2019; Thrivikraman et al., 2018; Wang et al., 2020).

Regardless of the underlying mechanisms, OPG and TGF-β1 mRNA expression, in response to BES, revealed a promising trend, namely, that BES can be used to consistently and significantly alter mRNA expression (in terms of both upregulation and downregulation) specifically in osteoblasts. Despite this general trend, some variability was observed, particularly regarding low voltages in OPG. Variability in the 0.1 V signals may be attributed to impedance within the stimulation system, which include connections, electrodes, and conductive media within each well. Such impedance likely reduced the signal to noise ratio of the stimulus that reached the cell monolayer. The standard error from sister control dishes ranged from 0.06 to 0.15 Log 2 stimulus that reached the cell monolayer. The standard error from sister each well. Such impedance likely reduced the signal to noise ratio of the variability given that each sample was compared directly to its own control and not against the aggregate of all control values of the study.

Our results clearly indicate that bioelectric stimulation can induce both upregulation and downregulation of OPG mRNA expression and two upregulation regions of TGF-β1 mRNA expression in osteoblasts—mostly influenced by a signal frequency. This trend indicates that OPG and TGF-β1, both key bone remodeling and regulatory proteins, can be manipulated via bioelectric stimulation to potentially influence bone softening for teeth alignment and bone hardening for post-alignment tooth anchorage. In terms of mineralization, our mineralization data indicates that osteoblasts can respond to electrical stimulation with a physiological response resultant in more calcium deposits that directly instigate bone hardening in patients. Furthermore, this discovery opens an entirely new methodology for efficient, non-invasive, therapeutic treatment—particularly in orthodontic tooth movement.

In summary, this study demonstrated that BES selectively modulates mRNA expression of OPG and TGF-β1 in osteoblasts within a specific range of bioelectric frequency and voltage parameters. Frequency parameters produced predictable and consistent control of both upregulation and downregulation of OPG, and upregulation of TGF-β1 at low and high frequencies. It is feasible that such findings can be translated into clinical scenarios. Incorporating findings into clinical application could result in an effective, more aesthetic, and noninvasive method, leading to improved patient compliance in orthodontic treatment.

5. Conclusions

Orthodontic tooth movement involves long and painful periods of mechanical manipulation of teeth in patients’ mouths. The ability to modulate bone remodeling molecules like OPG and RANKL via BES creates new treatment paradigms to improve patients’ conditions by better ameliorating their stress, improving patient compliance, and reducing treatment time during orthodontic procedures.

Bioelectric stimulation has been used in other tissues to induce gene expression. This manuscript presents significant data that points towards the controlled expression of OPG, an important molecule to strengthen bone, and TGF-β1 which participates in inflammation mechanisms. It is remarkable, that these two molecules have a different optimal pulse frequency to improve their expression. This goes in

Table 1
Fold and log fold change due to treatment (2 ΔΔCt) of mRNA expression from bioelectric stimulations.

| Frequency (V) | Volts | Fold Change (ΔΔCt) | Log Fold Change (Log 2ΔΔCt) | Fold Change (2 ΔΔCt) | Log Fold Change (Log 2ΔΔCt) |
|--------------|-------|-------------------|-----------------------------|---------------------|-----------------------------|
| 25           | 2.0   | 1.09 ± 0.07       | 0.08 ± 0.07                 |                     |                             |
| 50           | 2.0   | 0.97 ± 0.10       | -0.04 ± 0.11                |                     |                             |
| 75            | 0.1   | 1.15 ± 0.17       | 0.12 ± 0.15                 | 1.54 ± 0.25         | 0.41 ± 0.16 *              |
| 75            | 2.0   | 1.04 ± 0.11       | 0.02 ± 0.10                 |                     |                             |
| 100           | 0.1   | 0.76 ± 0.47       | -0.73 ± 0.51                | 1.77 ± 0.34         | 0.53 ± 0.20 *              |
| 100           | 2.0   | 0.37 ± 0.09       | -1.06 ± 0.25                |                     |                             |
| 250           | 1.0   | 0.36 ± 0.01       | -1.02 ± 0.04                |                     |                             |
| 500           | 0.1   | 0.25 ± 0.07       | -1.44 ± 0.28                | 2.61 ± 0.40         | 0.94 ± 0.15 *              |
| 500           | 1.0   | 1.21 ± 0.24       | 0.15 ± 0.22                 |                     |                             |
| 500           | 2.0   | 0.85 ± 0.21       | -0.22 ± 0.26                |                     |                             |
| 750           | 0.1   | 0.86 ± 0.15       | -0.18 ± 0.19                | 1.95 ± 0.30         | 0.65 ± 0.15 *              |
| 750           | 2.0   | 0.81 ± 0.10       | -0.22 ± 0.12                |                     |                             |
| 1000          | 0.1   | 0.78 ± 0.19       | -0.28 ± 0.25                | 1.24 ± 0.10         | 0.21 ± 0.01 *              |
| 1000          | 1.0   | 0.80 ± 0.25       | -0.32 ± 0.31                |                     |                             |
| 2500          | 0.1   | 1.05 ± 0.04       | 0.04 ± 0.09                 | 2.08 ± 0.22         | 0.72 ± 0.11 *              |
| 2500          | 1.0   | 1.26 ± 0.09       | 0.19 ± 0.22                 |                     |                             |
| 5000          | 0.1   | 1.11 ± 0.11       | 0.09 ± 0.10                 | 1.93 ± 0.50         | 0.6 ± 0.24 *               |
| 5000          | 2.0   | 1.20 ± 0.07       | 0.18 ± 0.06                 |                     |                             |
| 7500          | 0.1   | 1.16 ± 0.15       | 0.13 ± 0.14                 | 1.68 ± 0.03         | 0.52 ± 0.02 *              |
| 7500          | 2.0   | 1.29 ± 0.16       | 0.24 ± 0.12                 |                     |                             |
| 10000         | 0.1   | 1.00 ± 0.09       | -0.01 ± 0.09                | 1.47 ± 0.17         | 0.38 ± 0.12 *              |
| 10000         | 2.0   | 1.30 ± 0.06       | 0.26 ± 0.05                 |                     |                             |
| 25000         | 0.1   | 1.49 ± 0.09       | 0.38 ± 0.06                 | 1.85 ± 0.51         | 0.51 ± 0.35                |
| 25000         | 2.0   | 1.38 ± 0.02       | 1.35 ± 0.04                 |                     |                             |
| 50000         | 0.1   | 1.26 ± 0.16       | 0.19 ± 0.13                 | 1.35 ± 0.09         | 0.48 ± 0.25                |
| 50000         | 2.0   | 1.27 ± 0.06       | 0.23 ± 0.04                 |                     |                             |
| 75000         | 0.1   | 1.98 ± 0.14       | 0.67 ± 0.08                 | 1.51 ± 0.38         | 0.51 ± 0.34                |
| 75000         | 2.0   | 1.60 ± 0.12       | 0.46 ± 0.07                 |                     |                             |
| 100000        | 0.1   | 1.41 ± 0.24       | 0.29 ± 0.14                 | 2.97 ± 0.44         | 1.07 ± 0.15 *              |
| 100000        | 2.0   | 3.32 ± 0.09       | 1.20 ± 0.02                 |                     |                             |
| 250000        | 0.1   | 1.30 ± 0.16       | 0.21 ± 0.14                 | 3.32 ± 0.72         | 1.15 ± 0.23 *              |
| 250000        | 2.0   | 1.34 ± 0.02       | 0.29 ± 0.01                 |                     |                             |
| 500000        | 0.1   | 2.72 ± 0.11       | 0.99 ± 0.04                 | 1.67 ± 0.01         | 0.51 ± 0.01 *              |
| 500000        | 2.0   | 2.24 ± 0.03       | 1.51 ± 0.03                 |                     |                             |

(continued on next page)
growth factor modulation. Naturally, the functionality of the proteins necessary to understanding the clinical relevance of non-invasive treatment will allow better elucidation of different proteins and/or mechanisms that respond to electrical fields and intracellular pathways.

Conversely, the responses at low frequencies are totally opposite, whereby the expression of TGF-β1 and OPG in osteoblasts. However, in osteoblast treatment, OPG and TGF-β1 expression changes of other mRNAs and proteins involved in bone remodeling process, such as the receptor activator of nuclear factor-

Table 1 (continued)

| Frequency | Volts | Fold Change (2**ΔΔCt) | Log Fold Change (Log 2) | Fold Change (2**ΔΔCt) | Log Fold Change (Log 2) |
|-----------|-------|-----------------------|------------------------|-----------------------|------------------------|
| 750,000   | 0.1   | 1.27 ± 0.04            | 0.17 ± 0.22            | 2.31 ± 0.12           | 0.84 ± 0.05            |
| 750,000   | 2.0   | 1.04 ± 0.21            | 0.34 ± 0.19            | 1.68 ± 0.21           | 0.50 ± 0.12            |
| 1000,000  | 0.1   | 0.16 ± 0.10            | 0.14 ± 0.08            | 0.70 ± 0.06           | 0.08 ± 0.08            |
| 1000,000  | 1.0   | 0.10 ± 0.10            | 0.12 ± 0.10            | 0.09 ± 0.09           | 0.16 ± 0.12            |

* Indicates significant change due to treatment, p < 0.05. OPG; Osteoprotegerin.

accordance to reported data that suggests that other biological gene/molecules possess their own optimal frequency for expression. The use of bioelectric signals is a remarkable improvement to reduce discomfort, accelerate mineralization, and improve tooth alignment.

6. Future studies

The possibility that the expression of TGF-β1 enhances the expression of OPG has been previously suggested (Yang et al., 2015). Nonetheless, in our cellular system, we indeed see that OPG, and TGF-β1 respond similarly to high frequencies, although they could be easily differentiated by several hundred Hertz in terms of response to the stimuli, suggesting that their expression may be interdependent. Conversely, the responses at low frequencies are totally opposite, strongly suggesting that TGF-β1 may not be directly involved in the expression of OPG in osteoblast in vitro, nonetheless, it would be necessary to explore the nature of possible interactions between the expression of these two molecules.

The clinical use of bioelectric stimulation will demand the discovery of electrical signature protocols that selectively induce the expression of distinct and specific molecules and growth factors. We have determined these electrical signatures for TGF-β1 and OPG in osteoblasts. However, such discovery warrants additional exploration in determining the expression changes of other mRNAs and proteins involved in bone remodeling process, such as the receptor activator of nuclear factor-kappa B (RANK) and bone morphogenetic protein (BMP).

In addition to the clinical relevance of ES, another important goal of ES exploration is to determine the mechanism of action by which gene expression is influenced by specific electrical stimuli. Understanding the mechanism will allow better elucidation of different proteins and/or receptors that respond to electrical fields and intracellular pathways which lead to differential expression.

Lastly, the overarching influence of ES on physiological processes is necessary to understanding the clinical relevance of non-invasive growth factor modulation. Naturally, the functionality of the proteins expressed in response to bioelectric stimulation must be explored to demonstrate that the genes are still functional. Bioassays will be used to demonstrate growth factor viability and to show that bone does not overexpress other genes that can compromise the function of a specific protein studied (e.g., OPG and TGF-β1) (Table 1).

Credit authorship contribution statement

Chaudhari S.D., Conceptualization, Methodology, Investigation, Validation, Data Curation, Writing-Review & Editing. Sharma K.S, Conceptualization, Methodology, Writing-Review & Editing. Hyden H., Formal Analysis, Software, Data Curation, Writing-Review & Editing. Marchetto, J.J., Funding Acquisition, Data Curation, Clinical Methodology, Writing-Review & Editing. Burton M.B., Conceptualization, Methodology, Writing-Review and Editing. Moreno A.P., Writing Original Draft, Methodology, Conceptualization, Data Curation, Investigation, Writing-Review and Editing.

All authors gave their final approval and agreed to be accountable for all aspects of the work.

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