A simple rocker-induced mechanical stimulus upregulates mineralization by human osteoprogenitor cells in fibrous scaffolds

**Short title:** Rocking stimulates mineralization by osteogenic cells in 3D

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Abstract:

Biodegradable electrospun polycaprolactone (PCL) scaffolds can be used to support bone-forming cells and could fill a thin bony defect, such as in cleft palate. Oscillatory fluid flow (OFF) has been shown to stimulate bone production in human progenitor cells in monolayer culture. The aim of this study was to examine whether bone matrix production by primary human mesenchymal stem cells from bone marrow (hBMSC) or jaw periosteal tissue (HJP) could be stimulated using OFF supplied by a standard see-saw rocker. This was investigated for cells in 2D culture and within electrospun PCL scaffolds. From day 4 of culture onwards, samples were rocked at 45 cycles/min for 1 hour per day, 5 days per week (rocking group). Cell viability, calcium deposition, collagen production, alkaline phosphatase activity and vascular endothelial growth factor (VEGF) secretion were evaluated to assess the ability of the cells to undergo bone differentiation and induce vascularisation. Both cell types produced more mineralized tissue when subjected to rocking and supplemented with dexamethasone. Mesenchymal progenitors (hESMP) and hBMSC in 3D scaffolds up-regulated mineral deposition after rocking culture as assessed by micro-computed tomography (µCT) and alizarin red staining. Interestingly, VEGF secretion which has previously been shown to be mechanically sensitive was not altered by rocking in this system and was inhibited by dexamethasone.

Rocker culture may be a cost effective, simple pre-treatment for bone tissue engineering for small defects such as cleft palate.

Key word (up to 8 words)
marrow stromal cells; human jaw periosteal cells; fluid flow; osteogenesis; micro computed tomography; flow cytometry; electrospinning; cleft palate
1. Introduction

Cleft lip and/or cleft palate is a common oral and facial malformation birth defect with a worldwide incidence of approximately 1.7 per 1000 live births (Mossey et al., 2009). The current treatment necessitates multiple operations over several years and involves bone harvesting from the iliac crest (hip bone), resulting in donor site morbidity. The alveolar ridge region is usually filled with an autologous bone graft to support permanent tooth eruption. Many researchers in the field have suggested investigating potential alternatives to bone graft as a treatment for the reconstruction of cleft palate (Lohberger et al., 2013). Moy et al., (1993) found that bone graft substitute (BGS) materials implanted alone instead of autologous bone graft resulted in a lower success rate compared with the combination of cells and BGS. This suggests that tissue engineering strategies using cells combined with a supportive biomaterial scaffold may be a more effective treatment for cleft palate repair than biomaterials alone (Behnia et al., 2012, Khojasteh et al., 2008).

Mesenchymal stem cells (MSC) have been widely used for bone regeneration due to their ability to self-renew and differentiate into osteoblastic cells, using specific culture media. Harvesting bone marrow MSC (hBMSC) from the iliac crest incurs less pain at the operation site, fewer arterial and nerve injuries, and a lower risk of infection compared to harvesting of traditional iliac crest bone grafts (Samee et al., 2008). Tissue engineering strategies using MSC from patients have shown promise for reconstruction of a cleft defect at the alveolar ridge region; resulting in a successful repair and support of tooth eruption over the following two years (Hibi et al., 2006). Another interesting cell source for bone repair is human periosteal cells. Periosteal cells reside in a connective tissue membrane that covers the outer surface of all bones, except at the joints of long bones, they can be extracted from human jaw periosteum (HJP) and appear to contain a sub-population of MSCs. Studies have shown that HJP derived cells show promise for bone tissue engineering due to reduced donor site morbidity and time of operation compared to autologous bone graft from iliac crest (Trautvetter et al., 2011). HJP tissue is easy to harvest during wisdom tooth removal or other routine maxillofacial surgical procedures. For patients with cleft palate, HJP tissue could be harvested from the palate at the
time of palatal closure which typically takes place at 6-12 months of age (National-Health-Service, 2010).

Polycaprolactone (PCL) electrospun scaffolds are a promising support for bone forming cells as they show low biodegradation rates, good mechanical properties (Jha et al, 2011) and PCL is approved for medical use by the U.S. FDA (Hutmacher et al., 2001). Electrospinning is a straightforward fabrication method able to produce fibres of various diameters from 5 nanometers to several micrometres (Zargarian and Haddadi-Asl, 2010). The non-woven fibrous sheets formed are suitable for tissue engineering a thin region of bone, such as the palate.

Several studies have shown that fluid flow induced mechanical stress can enhance the osteogenic differentiation of MSCs, as previously reviewed (Delaine-Smith et al., 2012, 2014b). Oscillatory fluid flow (OFF) is similar to the flow found in the canalicular system in mature bone structure and flow in the bone marrow (Gurkan and Akkus, 2008). Previous work from our group, demonstrated that OFF applied using a standard see-saw rocker stimulated mineralized matrix production by osteoblasts and embryonic derived osteoprogenitor cells in monolayer culture (Delaine-Smith et al, 2012). Based on these findings it was hypothesised that OFF could increase osteogenic differentiation and mineralization of primary osteoprogenitor cells to precondition a construct for cleft palate patients. The aims of this study were firstly to determine if clinically applicable primary adult osteoprogenitor cells respond to rocker culture in a similar way to the cell lines previously examined and to determine if the angiogenic factor VEGF was upregulated by OFF culture. Secondly, we aimed to apply the mechanical stimulation technique to a tissue engineered construct suitable for cleft palate: an electrospun PCL scaffold seeded with mesenchymal progenitor cells.
2. Materials and methods

All chemicals and culture consumables were obtained from Sigma-Aldrich (Dorset, UK) unless otherwise stated and used as supplied.

2.1. PCL scaffold fabrication and characterization

PCL pellets (molecular number 80,000) were dissolved in dichloromethane (Fisher Scientific, Loughborough, UK) at a concentration of 10 wt%. Solutions were stirred for a minimum of 24 h at room temperature. Scaffolds were fabricated using an electrospinning rig as previously described (Bye et al., 2013, Delaine-Smith et al., 2014a). Sheets of PCL microfibers were fabricated at 17 kV, a flow rate of 40 μl/min, a working distance of 17 cm, and a drum rotation speed of 300 rounds per minute at room temperature. The solution was delivered via 4 needles to a rotating drum collector and the resulting scaffolds dried at room temperature and sterilized with 0.1% v/v peracetic acid before use.

Scaffold characterization was performed by scanning electron microscopy (SEM, Phillips XL-20 SEM, Amsterdam, The Netherlands) of scaffolds coated with a gold ultrathin layer. Mean fibre diameter and their distribution were measured over 100 randomly selected fibres from four recorded SEM micrographs using image analysis software (ImageJ, National Institute of Health, USA). Each picture was randomly overlaid with a square grid size 2500 μm² and 25 points were selected for fibre diameter measurements.

2.2. Cell Culture

Three different cell types were used in this study: 1) primary human jaw periosteal cells (HJPC) isolated from the periosteum tissue which was harvested from the jaw of patients undergoing maxillofacial surgery at the Charles Clifford Dental Hospital, Sheffield (with written, informed consent). This waste tissue collection was conducted under ethical approval 09/H1308/66 from the NRES Committee Yorkshire and The Humber, Sheffield, 2) human MSC (hBMSCs), isolated from bone marrow mononuclear cells from three individual donors (Lonza®, Castleford, UK) and 3) human embryonic cell derived mesenchymal progenitor cells hES-MP 002.5 (hESMP) (Cellartis, Gothenburg, Sweden) A schematic diagram detailing which methods were used with each cell type
can be found in supplementary figure 1. HJPs were not used for scaffold culture due to there not being sufficient numbers at low passages. hESMPs were the only cell type used for the μCT study as they secreted more mineral on average, and therefore mineral deposition was more likely to be detectable by μCT.

All cell types were cultured in an expansion media (EM) which consisted of α-MEM culture medium (Lonza®, Basel, Switzerland), supplemented with 10% fetal calf serum (FCS; v/v), 2 mM L-glutamine, 100 mg/ml penicilin and streptomycin (P/S) (basal culture medium). hESMP were cultured in pre-coated 1% gelatin 75 cm² tissue-culture flasks. For osteogenesis induction media (OIM), EM was supplemented with 50 µg/ml ascorbic acid-2-phosphate (AA) and 5 mM beta-glycerolphosphate (β-GP), or without the addition of dexamethasone (Dex) which was termed supplemented media (SM). 10 nM Dex was added to HJPC and hBMSC cultures and 100 nM Dex to hESMP cultures based on previous experiments in our laboratory to establish the best Dex concentration for osteogenic induction. All cells were cultured at 37ºC in 5% CO₂ in a humidified atmosphere. Media was changed every 2-3 days. HJPC and hESMPs were used between passages 3-8, as suggested by (De Bari et al., 2006) and hBMSC cells were used between passage 2-3.

2.3. Cell isolation

HJPC were isolated from freshly isolated periostium tissue of 2 different donors referred to as HJPC-1 and HJPC-2. Briefly, removed tissue of approximately 1x1 cm² was rinsed with phosphate buffered saline (PBS; OXOID Limited, Hamshire, England) containing 100 mg/ml P/S, cut into smaller pieces and added to 0.25% collagenase type II in EM without FCS and incubated at 37°C for 3 h (Samee et al., 2008). After this time, the cells were centrifuged, the supernatant removed and the cells placed into a 25 cm² tissue-culture flask in 2 ml of fresh EM. Fresh media was added to the flask every 2-3 days for 7 days. After 7 days the culture media and non-adherent cells were removed (Dominici et al., 2006, Chao et al., 2012). hBMSCs were isolated from mononuclear cells (Lonza®, Castleford, UK). Briefly, mononuclear cells were treated with pre-warmed basal culture media containing 0.1 mg/ml DNaseI (Stemcell™ technology, Grenoble, France). Cells were cultured at a density of 1.2 x 10⁵
cells/cm² in 25 cm² tissue-culture flasks. After 7 days, the non-adherent cells were removed by washing and the adherent population termed passage 0.

2.4 Cell phenotyping

hBMSC, hESMP and HJP were detached from culture flasks using non-enzymatic cell disassociation solution, washed in PBS and resuspended in 1 ml of FACS buffer (0.1% bovine serum albumin, 0.1% sodium azide, PBS 100 mM). Cell number was adjusted to 1x10^5 ml⁻¹ and cells stained with anti-CD90, anti-CD105, anti-CD146, and anti-CD45 (human MSC multi-colour flow cytometry kit, R&D system, Abingdon, UK) as per the manufacturer’s instructions. Samples were analysed using a LSR II flow cytometry (BD Biosciences, Oxford, UK).

2.5 Application of fluid shear stress by rocking

For cell monolayers, HJP and hBMSC were seeded in standard 6-well plates (Corning Inc., Amsterdam, The Netherlands) at a density of 10,000 cells per well in basal culture media. Cells were allowed to adhere for 24 h after media was replaced with osteogenic inducing media. For 3D cultures, hBMSC and hESMP were seeded onto the PCL scaffold (width 15 mm x length 35 mm x thickness 390µm) and placed in the standard 6-well plates with a sterile stainless steel ring on top to hold them in place. Scaffolds were immersed in 1 ml basal culture medium overnight before cell seeding. 1x10^5 cells in 100 µl were seeded onto the scaffold and left for 24 h after which the cell seeded scaffolds were transferred to fresh plates and a coil of sterilized dental wire placed on top to submerge the scaffolds in the culture media. Media was replaced with SM or OIM after a further 24 h.

For all conditions, samples were divided into a static condition (no-flow) or subjected to rocking which would induce OFF (termed rocking group) using a standard see-saw rocker (STR6 platform, Stuart Equipment, Stone, UK) starting from day 4 at 45 rpm for 1 h per day, for 5 days per week, at room temperature, based on previous published studies (Delaine-Smith et al., 2012, Lim et al., 2013, Delaine-Smith, 2013). The static group were sham treated by placing on the laboratory bench for the same length of time but not subjected to rocking. Medium samples were collected on day 7, 14, 21 for monolayer cultures, and 28 days for 3D cultures to evaluate vascular endothelial growth factor
(VEGF). In each case the media were changed 48 h prior to the collection time point so that the media collected represented 48 h of cell secretion.

2.6 Resazurin reduction test for cell viability

The resazurin reduction test was used to measure cell viability as previously described (O'Brien et al., 2000). Briefly, 0.1 mM resazurin solution salt solution in basal culture media was added to each sample and incubated for in the dark for 4 h at 37°C. 200 μL of pink resorufin product was transferred to a microtitre plate, and measured spectrofluorometrically (FLx800 Fluorescence Reader, BioTek, Potten, UK) at an excitation wavelength of 540/35 nm, and emission wavelength of 630/32 nm.

2.7 Total DNA measurement and alkaline phosphatase activity

Total DNA was assessed using fluorescent QuantiT™ PicoGreen® dsDNA reagent assay kit (Invitrogen, Paisley, UK) as per the manufacturer’s instructions. Briefly, 500 μL of carbonate buffer was added to the cells for 30 min at 37°C before scraping to remove cell lysate. Cell lysates were freeze-thawed three times, vortexed and centrifuged. Cell lysate and PicoGreen® solution were mixed and the fluorescence intensity measured spectrofluorometrically with an excitation of 480 nm and emission of 520 nm (Oliveira et al., 2006). The total DNA was calculated from the fluorescence emission to nano-gram per ml using a standard curve. To determine alkaline phosphatase (ALP) activity, cell lysate was mixed with an ‘Alkaline Phosphatase Yellow Liquid Substrate’ Sigma-Aldrich (Dorset, UK) based on p-nitrophenol phosphate and the absorbance measured using a spectrometer (ELx800, BioTeK, Potten, UK) at 405 nm every minute for 30 min. The enzyme activity was calculated as nM of para-nitrophenol per min (nmol pNP/min) and normalized to the total DNA from the same sample.

2.8 Collagen and calcium staining

Collagen and calcium staining was performed on day 21 and 28 for monolayer and 3D cultures, respectively. Total collagen production was measured by staining with 0.1% pico-sirius red solution for 18 h. All unbounded dye was removed with distilled water (dH₂O) and samples left to air-dry. The samples were destained with sodium hydroxide (0.2 M) and methanol (ratio 1:1) with
shaking at 50 rpm for 15 min. The absorbance of the resulting solution was measured in a spectrometer at 490 nm.

Total calcium deposition was measured by staining with 1 mg/ml alizarin red in dH₂O, adjusted to pH 4.1 with ammonium hydroxide for 20 min at room temperature. All unstained dye was removed with dH₂O and left to air-dry. The stained samples were dissolved by adding 500 µl perchloric acid (5% v/v) for 30 min at room temperature and the absorbance of the eluted stain measured at 405 nm using a spectrometer.

2.9 Micro-computed tomography imaging

Mineralized matrix was also evaluated using micro-computed tomography (μ-CT) (Skyscan 1172, Kontich, Belgium). Samples were stacked and placed on a 20 mm diameter brass tray. The μ-CT parameters were set at 17.5 µm/voxel resolution, two images for each point were merged, the filter was 1 mm of aluminium and a rotation step of 0.8º, 360º was applied. Approximately 750 slices were obtained. CTan and CTvol analysis software (Skyscan 1172, Kontich, Belgium) were used to quantify bone volume percentage (%BV) to evaluate mineral deposition on scaffolds and to reconstruct the 3D structure of the samples, respectively. The region of interest (ROI) for measuring %BV, a diameter of 1.2 mm (inner diameter of metal ring) and height of 390 µm (approximate scaffold thickness) was selected and sectioned equally into top, mid, and bottom portions (130 µm/region). Threshold values were calculated by analysing the grey scale index (GSI) distribution.

2.10 Assessment of vascular endothelial growth factor secretion

VEGF secretion into culture conditioned media was measured using a standard human VEGF ELISA Kit (PeproTech®, London, UK), following the manufacturer’s instruction. Absorbance values were converted to VEGF concentration (pg/ml) using a standard curve and normalised to cell number determined from the resazurin reduction assay.

2.11 Statistics
Data are expressed as mean values ± standard error of mean (SEM). Numbers of replicates are stated in the figure legend. ‘N’ represents a biological repeat (separate experiment) and ‘n’ represents a technical repeat (different samples within one experiment). Statistical analysis was performed using SPSS (IBM SPSS statistics 21). Cell viability, DNA quantification, ALP activity, calcium deposition, collagen production, and VEGF secretion were analyzed using a Mann-Whitney U test. The differences were considered to be statistically significant when the p-value was less than or equal to 0.05 (p ≤ 0.05).
3. Results

3.1 Rocking in combination with dexamethasone supplementation upregulates mineralization in monolayer cultures of primary adult hBMSC and HJPCs.

In monolayer culture (Figure 1a) the number of hBMSC, as measured by total DNA, increased over time in all conditions, irrespective of the supplements added (SM versus OIM) from day 7 to 21 while under rocking conditions cells in both media showed an increase to day 14 then no further increase by day 21.

With respect to ALP (Figure 1b), the highest ALP activity was observed in the OIM conditions with no difference between the rocking and static groups. In the absence of dexamethasone there was very little ALP production.

Collagen production was relatively low in all cells under all conditions (Figure 1c). Total calcium deposition however (Figure 1d) was stimulated in the OIM media group compared to the SM only group and further stimulated by rocking by day 21.

HJPC from two donors exhibited a fusiform mesenchymal stem cell-like morphology similar to hBMSC however hBMSC appeared visibly larger than HJPC (supplementary figure 2). The cell number, as measured by total DNA, of both HJPC-1 and HJPC-2 in monolayer culture increased under all conditions (SM and OIM groups) from 7 to 21 days (Figure 2a). The SM groups were similar to the OIM groups under both rocking and static conditions.

For HJPC, ALP activity normalized to DNA (Figure 2b) was higher in the presence of dexamethasone for both donors at all time-points. There was no significant difference between static and rocking conditions. For collagen production, the SM and OIM groups produced similar amounts of collagen in both static and rocking groups by day 21 (Figure 2c and e). However, total calcium deposition by the OIM rocking group was higher than for any other groups by day 21 (Figure 2d and f).
3.2 Rocking in combination with dexamethasone upregulates mineral deposition in cells cultured in electrospun scaffolds:

Electrospinning of PCL produced fibres with a smooth fibre morphology without bead formation. Representative SEM images of the 10wt% PCL randomised electrospun scaffolds spun using DCM as a solvent are shown in Figure 3a. The average fibre diameter of PCL was 2.7µm (Figure 3b). In electrospun scaffolds, the cell viability of the hBMSCs increased in all conditions (SM and OIM) with no difference between these groups from day 7 to 28 (Figure 3c). There were no differences in collagen production between any conditions (Figure 3d and f). Total calcium deposition was greatest for cells cultured in OIM –this was significantly higher than for the other groups ($P<0.05$) (Figure 3e and g).

The cell viability of hESMPs (mesenchymal progenitors) seeded in electrospun scaffolds increased over time in all conditions from day 7 to 28 (Figure 4a). There were no differences in collagen production between cells cultured under any conditions (Figure 4b and d). Total calcium deposition by the rocking OIM group was significantly higher compared to the other groups ($P<0.05$) (Figure 4c and e). The SM groups deposited negligible calcium compared to the OIM groups irrespective of whether they were rocked or not.

Mineralization was also studied using µ-CT analysis as described in Figure 5. Figure 5a describes the approach in which scaffolds were divided into 3 layers (Figure 5b) and examined using a Grey scale index (GSI) threshold for mineral detection. GSI in the range of 30-70 was found for PCL scaffolds without cells while for PCL scaffolds plus cells (hESMPs) cultured in OIM with rocking produced calcium deposits which could be detected with a GSI of 70 and greater as shown in Figure 5c. This group was chosen for thresholding as calcium deposition was demonstrated in these constructs based
on alizarin red staining (Figure 4c). The GSI values for the scaffolds without any cells were subtracted from GSI values for scaffolds plus cells to determine mineral deposition in the samples.

It was evident that mineralized matrix was not evenly distributed throughout the scaffolds. A side view image of a transverse section throughout the scaffold from the rocking OIM group shows that the top region contains more mineral than the other layers (white arrow head) and the top layer of the static OIM group (black arrow head) (Figure 5e). For the rocking OIM group there was significantly higher matrix in the upper region than in the middle and bottom regions ($p<0.05$) (Figure 5d). The middle region of the rocking OIM group contains more mineral than the middle region of the static OIM group.

3.3 Effect of dexamethasone and rocking on VEGF secretion

The secretion of VEGF (sampled by collecting media over a 48 h collection period) by hBMSC monolayers increased overtime under all conditions (Figure 6a). There was no significant effect of media or of rocking on VEGF secretion for these cells.

For cells cultured in 3D scaffolds VEGF secretion was detected from day 7 through day 28 at similar concentrations. VEGF levels in the absence of dexamethasone were slightly greater but this was not statistically significant. There was no effect of rocking on VEGF secretion (Figure 6b).
4. Discussion

In this study we have shown that primary bone marrow and jaw periosteal osteoprogenitor cells (HBMSCs and HJPCs) respond to rocking induced OFF as a mechanical stimulus with an increase in mineralization, as predicted by previous data on the embryonic cell line hESMPs. Subsequently we used hBMSCs and hESMPs, as cell sources that yield high numbers, to assess whether this affect could be used to stimulate the cells while in the 3D environment of an electrospun scaffold and showed that indeed mineralization in 3D can be stimulated by rocker culture.

Bone tissue engineering aims to provide a biocompatible material capable of substituting for autologous bone grafts for surgical reconstruction. The challenge for cleft palate repair is to fabricate scaffolds for the hard part of the palate to allow normal facial development. Further requirements are that the polymer scaffold is biodegradable and has the appropriate mechanical strength and biological properties to behave as a temporary extracellular matrix (ECM) until the supported cells are able to synthesize their own ECM to enable regeneration. In order to do this electrospun PCL was chosen as the test material because of its biocompatibility, high porosity, slow degradation rate, and ability to support bone forming cells. For example an in vivo study found that PCL scaffolds degraded by about 39±1% after 28 days of implantation in mice more slowly than polylactic-glycolic acid co-polymer (PLGA, 50:50) (Sung et al., 2004). Moreover, PCL electrospun scaffolds partially mimic the fibrous architecture of collagenous ECM and support good cell adhesion, proliferation, and osteogenic differentiation (Hutmacher et al., 2001).

All cell types used in this study were analysed for their surface antigen expression. Both hESMP and hBMSC were confirmed to be MSCs by expression of CD146, CD105, and CD90 and the lack of expression of CD45 (supplementary data 3) which are key makers for MSCs (Tormin et al., 2011). CD45 would indicate the presence of haematopoetic cells which may contaminate the osteoprogenitor cells. However we found no evidence of CD146 in the HJPC group. With the caveat that this data was obtained from passaged cells and should be verified in freshly isolated cells we suggest that the lack of CD146 relates to the cell’s origins. CD146 is known as a melanoma cell adhesion molecule and it
has been found to be present in human bone marrow cells that contribute to the vascular niche but not on cells that contribute to the osteoblastic niche (Sloan and Waddington, 2009). This may be expected as the periosteum is a membrane beneath which bone formation occurs whereas hBMSC are derived from bone marrow which also contains a vascular niche.

OFF can enhance osteogenic differentiation and therefore may be of use as a pre-treatment in cleft palate bone tissue engineering. We previously showed that OFF applied using similar conditions slightly up-regulates ALP activity and collagen production and strongly upregulates calcium production in monolayer culture by hESMPs, (Delaine-Smith et al., 2012). Others also applied OFF to monolayer cultures of hBMSC or human fetal osteoblastic cells using a parallel plate flow chamber at a peak shear stresses of up to 2 Pa (Li et al., 2004). Fluid flow under an orbital shaker has also been shown to enhance chondrogenesis in HJP (Ferretti and Mattioli-Belmonte, 2014, Tarng et al., 2010). To our knowledge this is the first study to stimulate the osteogenesis of HJP using fluid flow and here we demonstrate that this stimulus enhances mineralized matrix production in a similar way as to that previously demonstrated for other osteogenic progenitors. HJP are derived from the patient jaw periosteum tissue which is easily removed from the donor site while the patient is undergoing palatal closure surgery. In the other hand, hBMSCs were collected from another harvesting region (hip bone), therefore patients may suffer from donor site morbidity and more time consuming surgical treatment to source these cells. While this data indicates some promising characteristics of HJP there were fewer cells than would be obtained from a typical bone marrow sample and so it will be important to understand whether faster proliferating cells or larger tissue samples can be obtained from younger donors.

There were some small but important differences between the previously demonstrated response of hESMP cells to the rocking stimulus and that presented here for primary cells. For example: here we did not see an effect of OFF on ALP activity or collagen production whereas we reported these to be slightly higher after rocking in hES-MP cells (Delaine-Smith et al., 2012). However it has been well demonstrated that there is high donor variability in the ALP response of individual human donors to an osteogenic stimulus (Sittichokechawut et al., 2010, Osyczka et al., 2009), as can be also seen by...
the error bars in figure 1b and 2b, and that ALP peaks during osteogenic differentiation (Lian and Stein, 1995), and therefore the apparent ALP response is highly dependent on the analysis time-points. Here ALP did not appear to have reached a peak before 21 days. Our result is consistent with the study of Nuaman et al., (2001) who found no effect of fluid flow on ALP activity measured on day 14.

Collagen deposition can be stimulated by a mechanical stimulus under certain conditions for example bone marrow MSCs cultured on 3D scaffolds and stimulated in a bioreactors (Zhou et al., 2011, Sikavitsas et al., 2003, Sittichokechaiwut et al., 2010). However in this study we did not demonstrate an effect of OFF in the cell types examined. This result is similar to the study by Li et al., (2004) who subject MSC to OFF in a parallel plate flow chamber and found the gene expression of collagen type I was not affected. Filipowska et al (2016) found that perfusion flow of hBMSC in porous polyurethane scaffolds could enhance osteogenic potential but had no effect on collagen production.

Our study found that calcium deposition, indicative of bone-like matrix, in the rocking groups under the OIM medium in both hBMSC and HJP was higher than in the SM medium. The combination of OFF and Dex appears to further enhance osteoblastic differentiation especially at the late stages. Interestingly, when we compare multiple modes of mechanical stimulation of MSCs in our laboratory we consistently find that calcium deposition is the most strongly affected by the mechanical stimulus (Sittichokechaiwut et al., 2010, Delaine-Smith et al., 2012).

Having established the suitability of rocker culture to stimulate mineralization in monolayer in cell lines and primary cells we proceeded to test whether this culture method could be applied to 3D scaffolds. To the best of our knowledge, no study has used a standard see-saw rocker as a simple method for mechanically stimulating cells with OFF for cell culture on electrospun scaffolds. Here we demonstrated that subjecting cell-seeded electrospun PCL scaffolds to short bouts of OFF on a rocker in the presence of Dex caused greater calcium deposition compared to static cultures. The effect was very similar in both embryonic derived hESMP, used here as a model cell line, due to their capacity to reliably mineralize in 3D culture, and primary adult hBMSC which are well-known osteoprogenitor
cells suitable for clinical study. The results presented here suggest that the combination of OFF and Dex during culture synergistically stimulate the osteogenic differentiation of mesenchymal progenitors on PCL scaffolds. Although there were not sufficient mesenchymal progenitors available from the periosteal source for 3D culture of HJPCs we predict that HJPCs could also be induced to mineralize in electrospun scaffolds by this method.

Fluid flow has been shown to enhance osteogenic differentiation of osteoprogenitor cells and calcium deposition by several pathways such as the stretch-activated ion channels, gap junctions, focal adhesion complexes, paracrine signaling such as prostaglandin E2 (PGE₂) and the primary cilia (Janmey and McCulloch, 2007, Delaine-Smith et al., 2014b). Lastly, OFF can stimulate the mineralization of cells by enhanced nutrient distribution and oxygen transportation to cells in scaffolds compared to static culture (McCoy and O'Brien, 2010). The electrospun PCL scaffolds in our study are thin (300µm thickness) compared to typical porous scaffolds subjected to bioreactor culture therefore nutrient diffusion should be less of a limitation to static culture compared to other studies. However it has been suggested that cells should not be more than 200 µm from the nutrient source therefore cells in the centre of the electrospun scaffolds may still have experienced limited diffusion, especially as matrix forms and fills up the pores in longer term static cultures (Muschler et al., 2004).

For hBMSC the effect of rocking in electrospun scaffolds was very similar to in monolayer, with the rocking group under Dex supplemented medium having the highest calcium deposition. It is interesting that the effect of Dex in the 3D culture is very small and not statistically significant (Figure 3e) compared to in 2D culture (Figure 1d). And therefore the difference that rocking makes in 3D is much greater, leading to 10 fold more calcium deposition compared to Dex alone. Kale et al., (2000) and Yamaguchi et al., (2014) reported that bone cells in 3D culture deposited more calcium and non-collagenous protein such as ALP and osteonectin than in monolayer culture. This might because the higher density of bone cells facilitates greater cell-cell communication and interaction compared with 2D culture.
µ-CT has advantages as a tool for real-time monitoring for tissue engineering as it is a non-destructive imaging technique used to evaluate quantitative mineralization in the scaffolds (Thimm et al., 2013, Cartmell et al., 2004, Hagenmueller et al., 2007). In addition it has been used widely in biomaterials for analysis of animal models because there is no need to biopsy the bone tissue for evaluation. The characterization of mineral deposition on electrospun scaffolds using µ-CT has not been reported in many studies. The threshold for quantitative µ-CT analysis of bone usually used is the global threshold or automate global threshold such as Otsu’s method (1979). However, this method has disadvantages as small amount of mineralized deposition would not be detected (Parkinson et al., 2008). Individual threshold determinations were used in this study. As demonstrated by alizarin red staining the combination of both Dex and rocking produced significantly higher mineral deposition on the electrospun scaffolds. However Dex alone did not increase the amount of mineral visualized by the µ-CT technique and the differences in mineralization where much smaller than those detected by alizarin red staining. Therefore it appears that the µ-CT was not able to detect small mineral deposits that were detectable by alizarin red. Interestingly the ability of µ-CT to localize the deposited mineral demonstrated that there was a higher percentage of mineral deposition at the top level of the rocked scaffolds than the middle and bottom sections (Figure 5d). The reasons for this were probably because in the rocker the fluid would have flowed over the scaffold surface and the flow magnitude would gradually decrease with scaffold depth assuming the scaffold is firmly attached to the well base (scaffolds remained in the same orientation with respect to the well base throughout the culture period). A more even distribution of mineral may be obtained by turning the scaffold over during rocking or by adapting the culture system such that the scaffold is suspended in flowing fluid. One would need a computational model of fluid flow through a porous medium to calculate what the shear stress is on the walls of the electrospun fibres, such as those used for other types of porous scaffolds (Birmingham et al., 2013, Marin and Lacroix, 2015).

Vascularisation is an important process during bone repair initiated by recruiting endothelial cells for blood vessel formation to provide nutrients and excrete waste from the regenerating bony matrix. In this study cells secreted VEGF in all conditions and although VEGF was slightly inhibited by Dex
this recovered at later time-points. Work by others has also indicated Dex may inhibit VEGF (Heiss et al., 1996). The rising levels of VEGF indicate that if implanted, the cells in the scaffolds would support recruitment of endothelial cells. However, there were no differences in VEGF secretion by hBMSC between the rocking and static groups in either monolayer or 3D culture. This did not support our hypothesis that VEGF would be upregulated by rocker culture. VEGF secretion has been shown to be markedly stimulated by pulsatile fluid flow (Thi et al.’s (2010), Yuan et al. (2013), in non Dex containing media. However in another study human MSC subjected to OFF in rocker culture showed only a small (19%) increase in VEGF secretion. This indicates there is a complex relationship between the flow stimulus, Dex and VEGF secretion. In general, VEGF has been shown to be upregulated in parallel to mineralization, the rocker regimen used in this study which strongly supported mineral deposition did not appear to be correlated with higher VEGF. However, VEGF was measured in the media as a product of all the cell’s activity and the CT scan results indicate that mineralization was localized more strongly to the upper surface, therefore it would be interesting to observe whether there were regional differences in VEGF secretion throughout the scaffolds.

5. Conclusions

Our study indicated that a rocking stimulus previously described as supporting mineralization of embryonic derived progenitors (hESMPs) in monolayer can be applied to primary more clinically relevant osteoprogenitor cells in both monolayer and 3D to stimulate mineralization. Furthermore we confirmed that electrospun PCL is a suitable scaffold material for bone tissue engineering as it showed good biocompatibility, cell attachment and allowed osteogenic differentiation of human osteogenic progenitor cells. HJPC responded to OIM and rocker stimulation in a similar manner to hBMSC, suggesting that these cells have potential as an autologous osteogenic progenitor cells. HJPC might be a new cell source for cleft palate repair which could be collected from palate closure surgery, at a patient age of about 1 year. The introduction of OFF induced by a standard see-saw rocker further increased the mineralization of cells in Dex supplemented culture conditions in both monolayer and 3D scaffolds, particularly on the upper surface of the scaffolds. This would be a cost-effective and easy to scale-up method for pre-stimulation of tissue engineered constructs. We
conclude that an oscillatory flow stimulus could stimulate hard tissue formation from human osteogenic progenitor cells on thin tissue engineered constructs for cleft palate repair and other oral maxillofacial reconstructions such as alveolar ridge augmentation for tooth implantation and guided bone tissue regeneration for periodontal surgery.

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Figure 1: The effect of OFF on hBMSC matrix formation in monolayer culture in the presence (OIM) and absence of dexamethasone (SM) after 21 days. DNA (a), ALP activity normalised to DNA (µg/ml) (b), collagen production by picro-sirius red staining (c), and total calcium deposition by alizarin red staining (d). The photoimages show representative sets of picro-sirius red staining (e) and alizarin red staining (f) on monolayer cultures of hBMSC. Data presented as mean ± SEM, (N=3, n=3), *p<0.05 comparison between the SM and OIM groups, θ=p<0.05 comparison between the static and rocking group.
Figure 2

Figure 2: The effect of OFF on HJPC monolayer culture, for 28 (HJP-1) days and 21 days (HJP-2), in the absence (SM) or presence of Dex (OIM). Different time-points are shown as the two groups of donor cells had different growth rates as seen in (a). Total DNA (a), ALP activity normalised to DNA (b), total collagen production by picro-sirius red staining (c), and total calcium deposition by alizarin red staining (d). The photoimages show representative set of picro-sirius red (e) and alizarin red staining (f). Data is presented as mean ± SE, (N=2, n=3), *=p<0.05 comparison between the SM and OIM groups, θ=p<0.05 comparison between the rocking and static groups in the same medium.
Figure 3: Top: (a) 10 wt% PCL electrospun scaffold characterisation and schematic of fibre diameter measurement from SEM images, (b) fibre diameter distribution and average fibre diameter of electrospun scaffolds. Scale bar 50 µm, mean ± SEM (n=100). Below: The effects of OFF on osteogenic differentiation of hBMSC cultured on 3D PCL electrospun scaffolds in the absence (SM) or presence of dex (OIM). The viability of hBMSC was measured using a resazurin reduction test (c) for 28 days. Total collagen production was measured using picro-sirius red staining (d) and total calcium deposition using alizarin red staining (e), after 28 days of culture. The photoimages show representative sets of picro-sirius (f) and alizarin red (g) staining of hBMSC. Data presented as mean±SEM, (N=2, n=3), * = p<0.05 comparison between the SM and OIM in the same condition, θ = p<0.05 comparison between the rocking and static groups in the same medium.
Figure 4: The effects of OFF on hESMP cultured on PCL scaffolds in the absence (SM) or presence of Dex (OIM) for 28 days. The viability of hESMP were measured using a reszasurin reduction test (a), total collagen production by picro-sirius red staining (b) and total calcium deposition was measured by alizarin red staining (c). The photoimages show representative sets of picro-sirius (d) and alizarin red (e) staining of hESMP. Data presented as mean ± SEM, (N=2, n=3), *= p<0.05 comparison between SM and OIM, θ= p<0.05 comparison between the rocking and static conditions.
Figure 5

Figure 5: Top: (a) A window from CTanalyze software presenting the grey scale distribution histogram (red circle). (b) Schematic of the section of the electrospun scaffold that was used to analyse calcium deposition (bone volume percentage). The cylindrical ROI, diameter 1.2 mm. The scaffold was divided into 3 layers with 130 µm in each layer: top, middle, and bottom. The grey scale distribution histogram of standard PCL and calcium deposition on the scaffolds was used for setting the thresholds (c). The range between 30-70 GSI served to identify standard PCL scaffolds. The range between 70-125 GSI was determined to identify calcium deposition (N=1, n=3). Below: The effects of OFF on hESMP calcium deposition cultured on PCL scaffolds in the absence (SM) or presence of Dex (OIM) for 28 days. The top, middle, and bottom of percentile bone volume (%BV) with subtraction of standard PCL scaffolds measured using micro-CT analyser (d). Data presented as mean ±SEM, (N=2, n=3), * = p<0.05 comparison between SM and OIM. θ= p<0.05 comparison between the rocking and static conditions in the same level and medium. Microcomputed tomograph images of calcium deposition (orange area) in the scaffolds within the cylindrical region of interest (e). The circle images show the top surface of the scaffolds and half circle images showed the cross-section of scaffolds. The top level of the rocking-OIM group contained more calcium than the static-OIM group (white arrow heads).
Figure 6

Figure 6: The effects of OFF of VEGF secretion in hBMSC cultured as monolayers or 2D culture (a) or in PCL electrospun scaffolds or 3D culture (b) in the absence (SM) or presence of dex (OIM). VEGF secretion over 28 days (during a 48 hour collection period) normalised to viable cell number by resazurin assay. Data presented as mean±SEM, (N=2, n=3), *=p<0.05 comparison between the SM and OIM in the same condition.