Post microtextures accelerate cell proliferation and osteogenesis

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Abstract

The influence of surface microtexture on osteogenesis was investigated in vitro by examining the proliferation and differentiation characteristics of a class of adult stem cells and their progeny, collectively known as connective tissue progenitor cells (CTPs). Human bone marrow-derived CTPs were cultured for up to 60 days on smooth polydimethylsiloxane (PDMS) surfaces and on PDMS with post microtextures that were 10 μm in diameter and 6 μm in height, with 10 μm separation. DNA quantification revealed that the numbers of CTPs initially attached to both substrates were similar. However, cells on microtextured PDMS transitioned from lag phase after 4 days of culture, in contrast to 6 days for cells on smooth surfaces. By day 9 cells on the smooth surfaces exhibited arbitrary flattened shapes and migrated without any preferred orientation. In contrast, cells on the microtextured PDMS grew along the array of posts in an orthogonal manner. By days 30 and 60 cells grew and covered all surfaces with extracellular matrix. Western blot analysis revealed that the expression of integrin αvβ5 was greater on the microtextured PDMS compared with smooth surfaces. Real time reverse transcription-polymerase chain reaction revealed that gene expression of alkaline phosphatase had decreased by days 30 and 60, compared with that on day 9, for both substrates. Gene expression of collagen I and osteocalcin was consistently greater on post microtextures relative to smooth surfaces at all time points.

Keywords:
Polydimethylsiloxane
Microfabrication
Soft lithography
Connective tissue progenitor cells
Adult stem cells

1. Introduction

The response of cells to a substrate surface is strongly influenced by its properties, such as chemical composition and physical topography. These surface characteristics have important implications in the rational design and optimization of biological implants and bioreactors [1–5]. Recent advances in microelectromechanical systems (MEMS) technology provide new opportunities for the investigation of a variety of biological phenomena [6]. For example, soft lithography has been used to pattern the distribution of various chemicals on a material surface to explore selective cellular responses to specific biomolecular species [5]. Microfabrication and related MEMS techniques also enable precise production of surface topographical features to investigate the effects of physical cues on cellular behavior [4–7]. For tissue engineering applications these techniques could be conceptually combined with conventional scaffold processing strategies to ultimately provide scaffolds that possess precise topographical, spatial and chemical properties to optimize control over cellular behavior.

Numerous studies have been conducted to show that different surface topographies influence bone cell behavior [2–10]. For example, Hamilton et al. [7] reported increased cell proliferation and enhanced osteoblast differentiation on discontinuous edge surfaces compared with smooth ones and Mata et al. [8] reported an apparent increase in the proliferation of the progeny of bone marrow-derived connective tissue progenitors (CTPs) on polydimethylsiloxane (PDMS) micro-posts compared with identical cells cultured on smooth PDMS surfaces. CTPs are heterogeneous population of stem and progenitor cells that are resident in native tissue. These cells are capable of proliferating and giving rise to progeny which contribute directly to the formation of one or more connective tissues [11]. Harvest and transplantation, and even concentration, of CTPs from native bone marrow have been known to improve bone graft efficiency [12,13]. A characteristic of many marrow-derived CTPs is their ability to give rise to progeny that are capable of differentiating along a number of mesenchymal lineages, including bone, cartilage, muscle and fat [11–13].

Osteoblastic differentiation is particularly relevant to clinical bone repair strategies. Fig. 1 illustrates a generally accepted...
pattern of gene expression that reflects three major periods of cell and tissue development in bone formation [14,15]. The differentiated osteoblast is typically characterized by a specific pattern of gene expression, such as alkaline phosphatase (AP), collagen and osteocalcin (OC), and in vitro mineralization capacity [1,14,15].

In our previous study [8], human bone marrow-derived CTPs were cultured for 9 days on smooth PDMS surfaces and on PDMS with post microtextures that were 6 μm high and either 5, 10, 20 or 40 μm in diameter. We discovered that cells on microtextured PDMS exhibited a different morphology and increased cell count relative to those on a smooth PDMS surface. In particular, our investigations showed that 10 μm diameter post textures significantly enhanced CTP growth. This result suggested that, despite identical surface chemistries, the substrate topography had a significant effect on the biological performance of CTPs and suggested a potential role for microtextured materials in bone tissue engineering applications. In the current study, we investigate the influence of microtextured surfaces on proliferation and osteogenic differentiation of CTP progeny in long-term culture. Human marrow-derived CTPs were plated in primary culture on PDMS substrates presenting either post microtextures or a control (smooth) surface for up to 60 days, as an in vitro model to investigate the potential response of CTPs to different surface textures in a bone healing environment in vivo.

2. Materials and methods

2.1. Substrate preparation

The microfabricated PDMS substrate was produced by soft lithography techniques [8]. Briefly, a 6 μm thick layer of SU-8 2010 negative photoresist was first coated on top of a silicon (Si) wafer. Using an ultraviolet (UV) photolithography process, the 10 μm diameter and 6 μm high micro-post pattern, with 10 μm separation between posts, was then transferred from a photomask to the photoresist. Afterwards, this SU-8 mold was coated with H1H1H2H2H-perfluoro-decyltrichlorosilane (Lancaster Synthesis, Pelham, NH) to aid the release of PDMS in the final step of substrate production. The liquid PDMS base and curing agent (Sylgard 184, Dow Corning, Midland, MI) components were subsequently mixed at a ratio of 10:1, degassed for 20 min and then poured uniformly on top of the patterned SU-8 mold. After additional degassing for 10 min, the PDMS was cured at 85 °C for 2 h or at room temperature for 1 day (Fig. 2a). The cured PDMS cast was finally released from the mold and sectioned into 2 × 2 cm samples. Representative samples were inspected by scanning electron microscopy (SEM) (JSM-5310, JEOL USA, Peabody, MA). An unpatterned SU-8 mold was used to produce the smooth PDMS surface, which served as the control substrate for our study (Fig. 2b).

2.2. Cell culture

As described by Muschler et al. [16], bone marrow aspirates were harvested from the anterior iliac crest with informed consent from three patients immediately prior to elective orthopedic procedures. Briefly, a 2 ml sample of bone marrow was aspirated from the anterior iliac crest into 1 ml saline containing 1000 U heparin (Vector Labs, Burlingame, CA). The heparinized marrow sample was suspended in 20 ml heparinized carrier media (α-minimal essential medium (MEM) + 2 U ml⁻¹ of Na-heparin) (Gibco, Grand Island, NY) and centrifuged at 1500 rpm (400g) for 10 min. The buffy coat was collected, resuspended in 20 ml 0.3% bovine serum albumin–MEM (Gibco) and the number of nucleated cells was counted. The PDMS substrates were sterilized for 30 min with 70% ethanol and placed inside standard tissue culture dishes (Chamber Slide System, Lab-Tek, Naperville, IL). Cells were then plated on day 0 at a seeding concentration of 1 × 10⁶ cells per well (2 × 2 cm) and cultured for up to 60 days under conditions promoting osteoblastic differentiation [16].

In previous studies [11,16] we established that CTPs exhibit an osteoblastic phenotype after 9 days of culture. In order to determine possible effects of surface topography on the transition of cells from the lag to the log phase, cell counts were performed daily for the first 10 days and subsequently on days 30 and 60.

2.3. Cell morphology

On days 9, 30 and 60 the medium was removed and the plated substrates were placed in a solution containing 2% glutaraldehyde (Electron Microscopy Sciences, Fort Washington, PA), 3% sucrose (Sigma–Aldrich Co., Irvine, UK) and 0.1 M phosphate-buffered saline (PBS) (Cambrex BioScience, Walkersville, MD) at 4 °C and pH 7.4. After 1 h the substrates were rinsed twice with PBS for 30 min at 4 °C and washed with distilled water for 5 min. Dehydration was achieved by placing the plated substrates in 50% ethanol for 15 min while increasing the concentration of ethanol sequentially to 60%, 70%, 80%, 90% and finally 100%. Dehydrated samples were then critical point dried, mounted on aluminum stubs, sputter-coated with gold–palladium and examined using SEM (JSM-5310, JEOL USA).

2.4. Cell proliferation and differentiation

2.4.1. PicoGreen DNA quantification

CTP progeny seeded substrates were resuspended with 50 μl lysis buffer (1% sodium dodecyl sulfate, 10 mM ethylenediaminetetraacetic acid (EDTA) and 50 mM Tris–HCl, pH 8.1) (Sigma–Aldrich Co.) to lyse the membranes of adherent CTP progeny. After 60 min the samples were centrifuged at 14,000 rpm for 5 min and the supernatant was removed for analysis. A 40 μl sample of aqueous supernatant containing DNA was added to 0.96 ml TE buffer (10 mM Tris adjusted to pH 7.0 with HCl,
1 mM EDTA) (Molecular Probes, Eugene, OR). As per the manufacturer’s instructions (Molecular Probes), stock PicoGreen reagent was diluted 1:200 in TE buffer and 1 ml of that was added to each DNA-containing sample. The tubes were capped, vortexed and incubated at room temperature in a dark room for 3 min. The fluorescence was measured with a SpectraMax Gemini fluorescence microplate reader (Molecular Devices Co., Sunnyvale, CA), with excitation and emission wavelengths of 480 and 520 nm, respectively. All calibration samples were assayed four times and a fresh calibration curve was generated for each 96-well plate. Baseline fluorescence was determined with a TE blank, the average of which was subtracted from the averaged fluorescence of other samples. Using this analysis, we determined a DNA concentration of \( \frac{4.5}{10^6} \) adherent CTPs. Thus, we estimated total cell number by assuming that 4.5 pg of DNA represents one cell. Also, we performed a calculation to determine that, for identical projected surface areas, the actual surface area of post microtextures was 1.47 times greater than that of the smooth (control) surface. Consequently, we divided the total cell number (estimated via DNA quantification) from the post microtextures by 1.47 to enable a meaningful comparison with the cell number from smooth surfaces.

2.4.2. DAPI stain

Cell nuclei were stained with 6-diamidino-2-phenylindole dihydrochloride hydrate (DAPI). Ethanol-fixed cells were rinsed three times with PBS and then a 10 \( \mu \)l drop of DAPI-containing Vectashield mounting medium (Vector Labs) was placed on the substrates. Immediately thereafter the edges of the coverslips were sealed with three coats of clear nail polish and viewed under a fluorescence microscope (BX510F, Olympus Optical Co., Japan).

2.4.3. Alkaline phosphatase (AP) stain

Cells were stained for AP using Vector Red substrate working solution (5 ml 100 mM Tris–HCl with the addition of 2 drops of reagents 1, 2 and 3) (Vector Labs) for 30 min at room temperature in the dark, and then washed in distilled water. The positively stained cells with AP activity appeared red when viewed under a fluorescent microscope.

2.4.4. von Kossa stain

Cells were rinsed with PBS and fixed in 4% paraformaldehyde (Electron Microscopy Sciences) for 1 h. They were then incubated in 5% silver nitrate (Sigma–Aldrich Co.) for 30 min in a dark room, rinsed with distilled water and exposed to UV light for 1 h. Secretion of calcified extracellular matrix (ECM) was confirmed as deep blue-purple nodules with von Kossa staining under a phase contrast microscope.

2.5. Integrin expression

Integrin expression was confirmed by Western blot analysis. Cell extracts were prepared by adding lysis buffer containing 10 mM Tris–HCl, 5 mM EDTA, 150 mM NaCl, 30 mM sodium pyrophosphate, 50 mM NaF, 10% glycerol and a cocktail of protease inhibitors (Sigma–Aldrich Co.). Lysates were clarified by centrifugation at 13,000 rpm for 15 min at 4 °C and the protein content of supernatants was determined using a modified Bradford assay (Bio–Rad, Hercules, CA). Diluted 15 \( \mu \)g protein samples were loaded onto a 10% Tris–HCl ready gel (Bio–Rad) and electrophoresed in SDS running buffer at 130 V for 2 h. The proteins were electrophoretically transferred onto a Hybond–P membrane (Amersham Pharmacia Biotech, Piscataway, NJ) in 1× transfer buffer at 100V for 2 h in a cold room (4 °C).
Membranes were blocked with 5% skimmed milk and PBS containing 0.1% Tween (TPBS) at room temperature for 1 h. Primary antibodies (integrins α1, α2, α5 and β1 and GAPDH) (Chemicon, Temecula, CA) were diluted 1/200–1/1000 in 5% skimmed milk and incubated with the membranes overnight. After washing the membranes three times in TPBS they were incubated for 1 h with 1/1000 horseradish peroxidase (HRP)-conjugated secondary antibody (Chemicon). The membrane was washed again three times in TPBS and the signals were visualized using ECL detection reagents (Amersham Pharmacia) for 3 min and exposed to radiographic film (Eastman Kodak, Rochester, NY) for 30 s to 10 min. The intensity of the exposed bands was measured using the Gel-Pro program (Gel-Pro Analyzer version 3.1., Silver Spring, MD) to quantify protein expression.

2.6. Gene expression

2.6.1. Real time reverse transcription-polymerase chain reaction (real time RT-PCR)

Expression of osteoblast-specific genes, such as AP, type I collagen (Col I) and OC, were detected by real time RT-PCR. Total cellular RNA was isolated using a RNeasy kit (Qiagen Inc., Valencia, CA) and reverse transcribed by conventional protocols with a Sensiscript Reverse Transcription kit (Qiagen Inc.). Expression of AP, Col I, OC and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was quantified using real time RT-PCR analysis with a Power SYBR Green PCR Master Mix kit (Applied Biosystems, Foster City, CA). GAPDH is an enzyme utilized in cellular metabolism and is assumed to be expressed at the same level in most cells, therefore gene expression of GAPDH was used as an internal control to normalize for any differences in amount of total RNA isolated [16]. Real time quantitative PCR was performed on a 7500 Real Time PCR system (Applied Biosystems). Data analysis was carried out using 7500 System Sequence Detection software (Applied Biosystems) [17].

2.7. Statistical analysis

The PicoGreen DNA quantification and real time RT-PCR analyses were each performed a total of nine times (three replicates for each of the three patients, i.e. n = 9 for each substrate) as per our standard laboratory protocols. The mean and standard deviation (SD) values were subsequently calculated using the data of all groups. Statistical analysis (n = 9) was performed by a one-way analysis of variance (ANOVA) with Tukey’s multiple comparison test (SPSS Version 10.0., SPSS Inc., Chicago, IL) with a significance level of P < 0.05 (the 95% confidence interval).

3. Results

3.1. Cell morphology

The SEM images revealed that human CTPs attached to PDMS substrate surfaces showed varying cell morphologies (Fig. 2c). CTP progeny on the smooth surfaces exhibited arbitrary flattened shapes and migrated without any preferred orientation for up to 60 days. In contrast, on day 9 CTP progeny on PDMS post microtextures mostly tended to attach next to the posts and spread between them while directing their processes towards other posts and cells. On day 30 increased cell growth was observed on the post microtexture substrates. By day 60 numerous cells had grown and spread over the top of the post microtextures and covered most of the surface with ECM.

3.2. Early cell growth

CTP cultures from all three donors expanded with characteristic lag and log phases (Fig. 3a). DNA quantification analysis revealed that the number of cells initially attached (day 1) to the PDMS substrates were almost identical (391 cells cm⁻² for smooth and 361 cells cm⁻² for microtextured). Furthermore, there were minimal changes in daily cell numbers for both substrates through day 4. On day 5 CTPs on PDMS post microtextures transitioned to log phase, as demonstrated by a 3.6-fold increase in cell number. In contrast, the lag phase of CTPs on smooth surfaces lasted longer, with a 2.1-fold increase in cell number occurring on day 7 (Fig. 3b). After exiting from lag phase the rate of increase in cell number in log phase was similar for both PDMS post microtextures and smooth surfaces.

3.3. Cell proliferation

The CTPs on substrates were viewed in situ on days 9, 30 and 60 by DAPI staining (Fig. 4). The stain revealed that there were qualitatively more cells on PDMS post microtextures than smooth surfaces throughout the culture period. This observation was confirmed by the DNA quantification analysis (Fig. 4). On day 9 PDMS post microtextures supported more than twice the number of cells supported on smooth surfaces. Furthermore, on day 30 the PDMS post microtextures supported over three times the number of cells supported on control surfaces (P < 0.05). Finally, on day 60 PDMS post microtextures continued to support almost twice the number of cells compared with control surfaces.

3.4. Integrin expression

The expression of all integrins on day 9 was low, most probably due to the relatively low number of cells (data not shown). On days 30 and 60 Western blot analysis revealed that integrins α1, α2, α5 and β1 were expressed by the cells on all surfaces, although integrin α1 was expressed at generally lower levels (Fig. 5). On day 30 integrins α1, α2 and β1 demonstrated comparable expression levels between post microtextures and smooth surfaces. In contrast, integrin α5 exhibited an almost 3-fold greater expression level on the PDMS post microtextures. On day 60 integrin α5 still exhibited increased expression on PDMS post microtextures compared with smooth surfaces, although the difference had decreased to 1.5-fold.

On day 60 integrins α1, α2 and β1 again demonstrated comparable expression levels between substrates. However, compared with day 30 the actual expression level of α2 had increased over 2-fold, while decreasing very slightly for integrin α1 and increasing slightly for β1. Integrin α5 still exhibited a greater expression level (1.5-fold) on the PDMS post microtextures compared with smooth surfaces. However, compared with day 30 the actual expression on smooth surfaces had increased by almost 2.5-fold, while increasing very slightly for the post microtextures.

3.5. AP expression and mineralization

CTPs on the various PDMS substrates were also viewed in situ on days 9, 30 and 60 using AP and von Kossa staining. In general, cells on the PDMS post microtextures stained more intensely for AP compared with smooth surfaces on day 9 (Fig. 6a). Almost all cells expressed AP on both substrates on days 30 and 60. In contrast, the von Kossa staining was minimal on all substrates on day 9, but increased on day 30, and even further on day 60 (Fig. 6c). The spatial distribution and intensity of the von Kossa stain appeared to be consistently greater on PDMS post microtextures than on the smooth surfaces.

3.6. Gene expression

Expression of key osteoblastic bone markers, such as AP, Col I and OC, was assessed in the CTP progeny on substrates using real
time RT-PCR. The results revealed that AP expression was higher on PDMS post microtextures on day 9, but decreased on days 30 and 60 (Fig. 6a). Col I expression increased over time on both microtextured and control surfaces, but was consistently greater on microtextured surfaces than on control surfaces (Fig. 6b). Trends in OC expression were similar to Col I, with consistently greater expression on microtextured surfaces, particularly on day 60 (Fig. 6c).

4. Discussion

Understanding and optimizing cell–ECM–substrate surface interactions is critical to the rational design of biological implant materials and tissue engineering scaffolds, particularly for bone healing applications, where the concentration and prevalence of local osteogenic CTPs in native tissues are generally suboptimal [18–22]. We hypothesized that the biological performance of CTPs and their progeny with respect to proliferation, migration and osteoblastic differentiation would be significantly modified by interaction with PDMS post microtextures when compared with control (smooth PDMS) surfaces, as assessed by cell retention, expression of integrins, AP activity, matrix deposition and mineralization associated with osteoblastic differentiation.

Our study shows that the transition of CTP progeny from lag to log phase and associated indicators, such AP expression, collagen and osteocalcin synthesis, mineralization and the expression of integrin α5, were all increased or accelerated on PDMS post microtextures when compared with the smooth surfaces. The choice of PDMS as the construction material for both substrates, and the similarities in processing and handling and identical culture conditions, allow us to gain insights into the role of surface topography on cell growth, since the underlying surface chemistry is identical.

Cellular responses are generally attributed to the surface-adsorbed ECM, which comes from the surrounding medium and can also be produced by cells themselves [20–22]. Cells are known to attach to ECM via integrin receptors, and integrin-mediated attachment affects cell behavior, such as adhesion, migration, proliferation and differentiation [23]. Integrin α1β1 and α2β1 bind to type I collagen, which is the dominant bone matrix protein, and this binding has been reported to regulate osteoblastic differentiation.

Fig. 3. Time dependent (a) growth curves and (b) fold changes of CTPs attached to PDMS substrates. DNA quantification analysis revealed that the number of CTPs initially attached to PDMS substrates were almost identical. However, the greatest fold change was observed between day 4 and day 5, as the cells on PDMS post microtextures were about to exit from lag phase. Cells on the control surfaces were about to exit from lag phase on days 6 and 7. After exiting from the lag phase, the cell growth curves were similar on PDMS post microtextures and control surfaces.
Binding of integrin $\alpha_5\beta_1$ to ECM is known to regulate osteoblast survival, proliferation, bone-specific gene expression and matrix mineralization [24,25]. In our Western blot results integrin $\alpha_5$ and $\beta_1$ were expressed similarly over time, while the expression of integrin $\alpha_5$ was greater on the PDMS post microtextures compared with control surfaces. Thus, these data suggest that the increased integrin $\alpha_5$ expression of CTPs on post microtextures resulted in increasing ECM production and subsequent osteoblast-specific gene expression by cells on post microtextures (Fig. 6).

Fig. 4. CTP number on PDMS post microtextures and corresponding control surfaces. On day 9 the number of CTPs on the post microtextures was greater than on the control surfaces. The post microtextures exhibited a significant increase in CTPs on day 30. On day 60 the number of CTPs increased on post microtextures compared with the control surfaces. Fluorescent images show cell nuclei stained with DAPI and reveal more cells on post microtextures than control surfaces. (Note: the original color images were converted to grayscale and reversed to provide visual clarity). The PicoGreen DNA quantification was performed a total of nine times (three replicates for each of the three patients, i.e., $n = 9$ for each substrate) as per our standard laboratory protocol. Statistically significant compared with control surfaces on day 30 ($P < 0.05$).

Fig. 5. Integrins and GAPDH expression in CTPs after 30 and 60 days culture on PDMS post microtextures and control surfaces. Integrins $\alpha_1$, $\alpha_2$, $\alpha_5$ and $\beta_1$ were expressed by the cells on all surfaces, although integrin $\alpha_1$ was expressed at generally lower levels. On days 30 and 60, integrins $\alpha_1$, $\alpha_2$, and $\beta_1$ showed comparable expression levels between post microtextures and smooth surfaces. In contrast, integrin $\alpha_5$ exhibited greater expression level on the PDMS post microtextures compared with smooth surfaces.

[24].
account for the difference in actual surface areas between post microtextures and smooth surfaces. Furthermore, according to our previous experiments [8] and Fig. 3 the numbers of CTPs attached initially to the PDMS post microtextures and smooth surfaces were almost identical. Therefore, it appears that the enhanced cell growth on post microtextures resulted primarily from the shortened lag phase, which is evident in Fig. 3. The earlier onset of proliferation would lead to increased cell number on days 9, 30 and 60. Presumably, the higher cell number resulted in their early confluence and stimulation of osteogenic differentiation.

![Gene expression of (a) AP, (b) collagen type I (Col I) and (c) osteocalcin (OC) by CTPs after 9, 30 and 60 day on PDMS post microtextures and control surfaces. (a) AP mRNA expression during development, with the highest levels present on day 9 on post microtextures, while it was expressed more highly on the control surface on day 60. Fluorescent images show cells on the post microtextures stained more intensely for AP compared with control surfaces on day 9, and AP increased on all surfaces by days 30 and 60. (b) mRNA of Col I expression at all time points, with slightly higher expression on day 30 and significantly higher on day 60. (c) Compared with day 9, OC mRNA expression showed a significant increase in cells grown on post microtextures on day 60 compared with cells grown on the control surface. Phase contrast images show von Kossa stain and the intensity of this stain on post microtextures increased with time compared with the control surfaces. The real time RT-PCR was performed a total of nine times (three replicates for each of the three patients, i.e. n = 9 per substrate) as per our standard laboratory protocols. Statistically significant compared with control surfaces on same day (P < 0.05).]
(ECM production, osteoblast–ECM related integrin expression and subsequent osteoblast-specific gene expression) on post microtextures. Larson et al. [26] have investigated the growth of passaged human bone marrow-derived multipotent stromal cells in low density (50 cells cm$^{-2}$) cultures on standard tissue culture substrates for up to 10 days. In this time frame their cultures expanded with characteristic lag, log and near stationary phases, with the greatest fold change observed on days 2 and 3, when the cells transitioned to log phase. In contrast, our CTPs remained in lag phase for 6 and 4 days for the smooth and microtextured PDMS, respectively. Furthermore, the CTPs continued to proliferate for over 30 days without achieving stationary phase. Possible reasons for the differences in our results from the previous reports could arise from our choice of substrate material (PDMS), culture conditions (high density) and the heterogeneity of the CTP population. Additional investigations will be required to gain further insights into the differences in growth characteristics between CTPs and conventional mesenchymal stem cells.

Numerous groups have examined the effect of surface topographies on cell behavior [27–38]. Table 1 presents a summary of results from previous work and a comparison with the current study. In general, our results are consistent with previous reports which suggest that surface microtopography influences osteogenesis. However, the current study has focused on CTPs, which have not been examined similarly in the past. In addition, we have demonstrated that, in spite of the identical surface chemistry, cells cultured on post microtextures appear to exhibit a shortened lag phase relative to smooth surfaces. Thus, even when the initial numbers of cells attached to the substrates were similar, post microtextures ultimately resulted in increased cell number and appeared to accelerate osteogenesis.

There are many factors contributing to the increased cell numbers and expression of osteogenic markers on different surface topographies. The incorporation of micro- and nano-scale topographies at the cell–substrate interface might provide an attractive approach to enhancing specific cell behavior without destabilizing the delicate biochemical environment. However, the mechanisms by which cell behavior changes in response to different geometrical and biochemical stimuli remain unclear. Further investigation is needed to elucidate all of the possible factors and establish definitive mechanistic links between cell–surface interactions and cell differentiation.

Moreover, demonstrating the feasibility of using CTP surface topography constructs for bone tissue formation will require experiments in an animal model for performance assessment in vivo. These experiments could initially be performed with 3-dimensional scaffolds with different surface topographies to verify enhanced CTP growth characteristics. Afterwards, the performance of 3-dimensional scaffolds should be investigated in an animal model to provide insights into the behavior of these for human bone graft applications.

5. Conclusion

Post microtextures accelerate proliferation and osteogenic differentiation of CTP cultures compared with those on control (smooth) surfaces. This difference in overall cell growth can be attributed to a reduction in lag time between initial cell contact.
with the substrate surface and cell spreading. Cells proliferate in greater numbers, express higher amounts of ECM genes and secrete greater amounts of minerals on the post microtextures than on the control surfaces after long-term culture. The earlier onset of cell proliferation on microtextures results in increased cell numbers, their early confluence and stimulation of osteogenic differentiation. This study demonstrates a valuable in vitro model, based on the precise and reproducible patterning capabilities of microfabrication and related MEMS techniques, in which to explore the relationship between topographical features of bone tissue engineering scaffolds and the likely response of human adult stem cells and progenitor cells in the setting of bone repair in vivo.

Appendix. Figures with essential colour discrimination

Certain figures in this article, particularly Figs. 3, 4 and 6, are difficult to interpret in black and white. The full colour images can be found in the on-line version, at doi:10.1016/j.actbio.2009.06.016.

References

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Table 1

Effect of surface topography on cells.

<table>
<thead>
<tr>
<th>Reference</th>
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<th>Time</th>
<th>Surface topography</th>
<th>Investigation/results</th>
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<td>Rough Ti surfaces with different morphologies (SLA Ra = 3.3 μm and TPS Ra = 0.22 μm)</td>
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<td>[28]</td>
<td>Mgf3 osteoblast-like cells</td>
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<td>Human embryonic palatal mesenchymal cells</td>
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<td>600 Grit (grooved) or sandblasted (roughened) cpTi implant discs</td>
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<td>Rat osteoblasts</td>
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<td>The effect of substratum topography on osteoblast adhesion mediated signal transduction and phosphorylation</td>
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<td>[36]</td>
<td>Epithelial cells and osteoblasts</td>
<td>4 Weeks</td>
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<td>Tapered pits stimulate osteoblast mineral deposition in vitro and in vivo, but do not prevent epithelial sheet migration</td>
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<td>[37]</td>
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<td>[38]</td>
<td>Human mesenchymal stem cell</td>
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<td>Our data</td>
<td>Human bone marrow-derived CTs</td>
<td>60 Days</td>
<td>PMDS post microtextures 10 μm diameter and 6 μm height (effective Ra = 3 μm)</td>
<td>PMDS post microtextures accelerate proliferation and osteogenic differentiation of CTs</td>
</tr>
</tbody>
</table>


