Nanopit-induced osteoprogenitor cell differentiation: The effect of nanopit depth

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Abstract
We aimed to assess osteogenesis in osteoprogenitor cells by nanopits and to assess optimal feature depth. Topographies of depth 80, 220 and 333 nm were embossed onto polycaprolactone discs. Bone marrow–derived mesenchymal stromal cells were seeded onto polycaprolactone discs, suspended in media and incubated. Samples were fixed after 3 and 28 days. Cells were stained for the adhesion molecule vinculin and the osteogenic transcription factor RUNX2 after 3 days. Adhesion was lowest on planar controls and it was the shallowest, and 80-nm-deep pits supported optimal adhesion formation. Deep pits (80 and 220 nm) induced most RUNX2 accumulation. After 28 days, osteocalcin and osteopontin expression were used as markers of osteoblastic differentiation. Deep pits (220 nm) produced cells with the highest concentrations of osteopontin and osteocalcin. All topographies induced higher expression levels than controls. We demonstrated stimulation of osteogenesis in a heterogeneous population of mesenchymal stromal cells. All nanopit depths gave promising results with an optimum depth of 220 nm after 28 days. Nanoscale modification of implant surfaces could optimise fracture union or osteointegration.

Keywords
Osteointegration, nanotopography, osteoprogenitor

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Introduction
The control and modification of surfaces on a micro and nanoscale has been shown to affect cell interaction with the material. Adherence,1–7 metabolic activation,4,8 alignment,9 gene expression10–12 and differentiation2,7,11,13 can all be controlled by topographical cues detected by the cells in contact with the surface. Osteoprogenitor cells and selected mesenchymal stem cells have been shown to respond to certain topographical patterns by preferentially differentiating into mature osteoblasts and exhibit higher levels of osteogenesis compared with planar control substrates. Furthermore, levels of osteogenic differentiation in response to topographic features alone have been observed to be similar to those using osteogenic differentiation supplements.2,14

In orthopaedic surgery, osteogenesis is key to fracture healing and osteointegration of implanted material. In un cemented arthroplasty implants, such as acetabular components of hip replacements, a stable implant with bony ingrowth correlates with longer implant survivorship and better function.15 Commonly, however, the implant–bone interface can produce a host response that results in fibrous encapsulation of the foreign material leading to early loosening and a threefold increase in pain.15 Because topography does not rely on changes in chemistry
and substrate mechanical properties, micro/nanoscale modification of the implant surface is a potential strategy for future implants to encourage a shift towards osteointegration preferentially instead of fibrosis.

A previous report has suggested that diameters of 30 μm are strongly osteogenic through mimicry of topographic features such as those left by osteoclasts in resorption pits. This ties in with previous observations of 40-μm-diameter pits being osteogenic. However, in these reports, linking feature depth to osteogenesis was largely unexplored and cell lines rather than primary cells were mainly used. Thus, here, we use a heterogeneous bone marrow–derived mesenchymal stromal cell mix which is present around an uncemented arthroplasty component inserted into cancellous bone. Furthermore, we fix diameter at 30 μm and focus on assessment of nanopit depth (ranging from 80 to 333 nm) and osteoinductive potential.

In order to achieve this, we first study cell adhesion to the features. Cell adhesion and derived intracellular tension is key to osteogenesis. Mesenchymal stem cells and osteoblasts actively expressing bone markers (differentiation is key to osteogenesis. Mesenchymal stem cells to mature osteoblasts. A total of 10 mL of aspartate was added to transfer media (10 mL phosphate-buffered saline (PBS), 0.03 g EDTA sterilised in autoclave). This was then washed with basal media (Dulbecco’s modified Eagle’s medium (DMEM), 10% foetal bovine serum, sodium pyruvate and non-essential amino acids) and centrifuged at 300g for 6 min.

This cell population was isolated from contaminating erythrocytes and plasma by Ficoll-Paque medium. Cells were cultured in basal media at 37°C and media were changed twice weekly. After two passages, and once samples were confluent, cells were seeded onto the PCL discs containing the topographies at a density of 1 × 10⁴ cells in 1 mL of basal media in 24 well flasks. A total of 12 identical wells were prepared for each topography and control, to allow each of the four markers to be stained in triplicate. Again, all samples were incubated at 37°C and media changed twice weekly.

Immunofluorescence

After 3 days, half of the samples were fixed using 4% formaldehyde/PBS, with 1% sucrose at 37°C for 15 min. When fixed, the samples were washed with PBS and a permeabilising buffer (10.3 g sucrose, 0.292 g NaCl, 0.06 g MgCl₂, 0.476 g HEPES buffer, 0.5 mL Triton X, in 100 mL water, pH 7.2) added at 4°C for 5 min. Samples had anti-vinculin (1:150 in 1% bovine serum albumin (BSA)/PBS, Sigma, UK), rhodamine-conjugated Phalloidin (1:50% BSA/PBS, Invitrogen, UK), or anti-RUNX2 (raised in rabbit, Insight Biotechnology, UK) added for 1 h at 37°C. Samples were then washed with 0.5% Tween 20/PBS (5 min × 3). Secondary biotin-conjugated antibodies (either anti-rabbit or anti-mouse, 1:50 in 1% BSA/PBS, Vector Laboratories, UK) were added to transfer media (10 mL) and samples incubated for 1 h at 37°C prior to washing. The tertiary, fluorescein isothiocyanate (FITC)-conjugated streptavidin, layer was then added (1:50 in 1% BSA/PBS, Vector Laboratories) and samples incubated at 4°C for 30 min. Discs, with fixed and stained cells on their surface, were then mounted on slides with Vectorshield mounting medium with

**Materials and methods**

**Fabrication**

Polycaprolactone (PCL) discs (Aldrich, average Mn 45,000) with the specific surface topographies (all 30 μm diameter, with a depth of 80, 220 or 330 nm) were created using the nickel shims and a hot embossing technique at 80°C (see Supplementary Information). Discs were trimmed to 1 cm diameter. Samples embossed with each topography, as well a control samples melted on a planar surface, were examined by scanning electron microscopy (SEM) to validate the technique. Samples were treated with three 10-s 70% ethanol emersions followed by serial emersions in HEPES saline prior to use.

**Cell culture**

Ethical approval for use of discarded human tissue was in place through NHS Greater Glasgow and Clyde. After obtaining informed consent, a bone aspirate was obtained from the femoral neck of a healthy adult at the time of total hip replacement for osteoarthritis. Samples obtained in this manner have been shown to contain a mixed cell population of osteoprogenitor cells, ranging from mesenchymal stem cells to mature osteoblasts. A total of 10 mL of aspartate was added to transfer media (10 mL phosphate-buffered saline (PBS), 0.03 g EDTA sterilised in autoclave). This was then washed with basal media (Dulbecco’s modified Eagle’s medium (DMEM), 10% foetal bovine serum, sodium pyruvate and non-essential amino acids) and centrifuged at 300g for 6 min.

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4',6-diamidino-2-phenylindole (DAPI) (Vector Laboratories). After 28 days, the remaining live samples were fixed and stained for either OPN or OC (both 1:150 raised in mouse, Insight Biotechnology, UK) using the above protocol. Samples were viewed under fluorescence microscope (Zeiss Axiovert 200M – 10–40× magnification, NA 0.5). Images were analysed using Photoshop CS (Adobe) and ImageJ, analysing 40 individual cells in each group for staining intensity and morphology.

Results

Materials

SEM of the PCL discs showed successful embossing of the topographies with a consistent 30 µm pit diameter with 90 µm centre–centre pit spacing in all samples in a square arrangement. Planar controls lacked any significant irregularities or patterning and were essentially smooth (Figure 1).

Cytoskeleton and morphology

After 3 days, the cells were seen to be well spread on all surfaces with well developed, abundant actin stress fibres (Figure 2). The stress fibres were particularly noteworthy on the 80-nm-deep pitted samples.

Cell adhesions

Vinculin expression was assessed after 3 days of culture. Individual cells, which had no identifiable contact with other cells, were selected for analysis to eliminate the influence of ‘cell to cell’ interaction instead of ‘cell to surface’ adhesion. A total of 40 cells in each group were analysed. Lower concentrations of vinculin were seen around the periphery of cells in the planar control group. Cells on the shallowest pits, 80 nm, had the highest levels of vinculin expression with notably larger and more distinct adhesions. However, it is noted that all the pitted surfaces supported more mature adhesions than the planar control (Figure 2).

Early osteoblastic differentiation

Expression of RUNX2, a transcription factor involved in osteoblastic differentiation, was assessed after 3 days. The data show that while RUNX2 could be noted in the nuclei of cells on the planar surface, more intense nuclear staining and also cytoplasmic staining of the transcription factor was noted in cells on the topographies, particularly the 80- and 220-nm-deep pits (Figure 2).

Expression of osteoblast phenotype

OC and OPN expression after 28 days were used as markers of cell differentiation into an osteoblastic phenotype. Using microscopy, cell populations were found to be well established on all surfaces and controls with large aggregates of cells spread through the discs. OPN in particular showed high levels of cytoplasmic and extracellular staining on topographies compared with planar controls (Figure 3).
Cells cultured on the 220-nm-deep pits were consistently observed to contain the highest concentrations of OPN. In contrast, only a perinuclear blush of OPN staining was visible on control samples, even when using high contrast settings. Using ImageJ analysis, multiple slides consistently showed higher staining intensity for 220-nm-deep pits compared with controls or other topographies. Average OPN staining intensity per slide for 220 nm pits was compared to control, 80 nm and 333 nm pits using Student’s t-tests ($p=0.017$, 0.029 and 0.045, respectively). Mean staining intensity of OPN per cell was also calculated for all images captured to control the variable of cell density.

Figure 2. Cells fixed after 3 days showing actin (cytoskeleton), vinculin (adhesion) and RUNX2 (osteoblastic transcription factor) staining. Vinculin formed large, distinct adhesion complexes at the peripheries of cells particularly on the 80-nm-deep features and the other topographies compared to control. Concomitantly, actin stress fibres were also more organised. RUNX2 had increased nuclear and cytoplasmic concentrations on the topographies compared with controls (arrows indicate nuclear localisation and arrowheads indicate cytoplasmic localisation).
when comparing different slides (Figure 4). Increased levels of OPN were again observed with 220-nm-deep topographies. Staining intensity of OC after 28 days also suggested superior osteoblastic differentiation on 220-nm-deep pits (Figure 3). The other topographies, 80 and 333 nm, also showed a trend towards increased OPN and OC expression compared to controls, but this did not reach statistical significance.

**Discussion**

Honing of surface nanopatterning to optimise and steer targeted cell response is dependent on a number of factors. Size, shape, spacing and configuration of the nanostructures (islands, grooves or pits) as well as the material’s physicochemical properties play an important role. The chemical properties of the surface have been shown to have an independent influence on osteoblast activation compared with topographic modification. A limiting factor can be the feasibility and reproducibility of nanofabrication in the chosen material.

In terms of osteogenesis, cells of an osteoblastic lineage must be stimulated to adhere, undergo preferential differentiation and metabolic activation with the final desired outcome of osteoid matrix synthesis and deposition. Adherence can initiate a cascade of intracellular
signalling and essentially acts to transfer information from the physical environment to make the cell ‘spatially aware’. Several theories are being explored to explain the exact nature of the topography–cell interaction. Adhesion formation and filipodia play part in contact guidance as part of this process.

It is perhaps not such a large surprise that shallower pits, 80 and 220 nm, showed superior adhesion and osteogenic induction than the deeper pits (333 nm). Osteoblasts have been shown to interact with nanogrooves with a depth of only 17 nm. Human foetal osteoblasts have been shown to exhibit superior adhesion formation on 14- and 29-nm-deep pits compared with 45-nm-deep pits. In terms of the smallest topographical features, the threshold for contact guidance of fibroblasts to occur has been shown to be 35 nm. Even smaller features, for example, 15 nm titanium nanopillar structures, had been shown to have the most osteoinductive potential compared with larger features and Mesenchymal stem cells (MSCs) have been shown to interact with features just 8 nm high. All these suggest that all features used in this study are of a size that can potentially interact and guide contacting cells.

On day 3, increased adhesion sizes and levels of cytoskeletal organisation could be noted on the features, particularly the 80-nm-deep pits. This is important in osteogenesis as recruitment of focal adhesion kinase (FAK) and activation of G-proteins is considered key to osteogenic stimulation. Indeed, RUNX2 expression is seen to intensify on these surfaces and is noted to be present in the cytoplasm as well as the nuclei. This perhaps suggests shuttling into the cytoplasm to allow phosphorylation and activation. It is thus logical that these particular topographies go on to induce the highest levels of OPN and OC expression. The 220-nm-deep features were particularly potent at initiating the deposition of these mature osteoblast markers (OC is totally exclusive to maturing osteoblasts).

We have demonstrated induction of osteogenesis in a cell population that is a suitable model for the cell mix present around joint replacements. Nanopits (220-nm deep) with a diameter of 30 µm had the greatest inductive potential. This would be a suitable nanopattern to develop using implantable orthopaedic materials, such as metals and ceramics, and continued use for further in vitro experimentation. There has been great interest in the development of such modified surfaces and materials with scope for a wide range of future orthopaedic applications. The expanding body of evidence and research exploring the use of nanotechnology in orthopaedic surgery may herald a new generation of implants and material.

**Supplementary Information**

**Fabrication of nickel shims**

Silicon wafers (Compart Technologies, UK) were cleaned under acetone in an ultrasonic bath for 5 min. They were rinsed thoroughly in reverse osmosis water (ROH2O) and blow dried with an air gun. Next, they were spun with primer (Shipley AZ Coupler, Shipley, UK) for 30 s at 4000 r/min, then spun with S1818 photoresist (Shipley AZ Coupler) for 30 s at 4000 r/min and baked for 30 min at 90°C. The resulting layer was measured to be 1.8 µm thick. The photoresist layer was exposed to UV light through a chrome mask on a Karl Suss MA6 mask aligner for 3.8 s. Then, the resist layer was developed for 75 s in 50:50, Microposit developer (Shipley AZ Coupler):ROH2O.

The developed circle patterns were then used as a mask for reactive ion etching. The silicon substrate was etched in the silicon tetrachloride gas plasma of a Plasmalab System 100 machine (gas flow = 18 sccm, pressure = 9 mTorr, rf power = 250 W, DC bias = −300 V). Each wafer was etched individually at 18 min at a nominal etch rate of 18 nm/min. All wafers were stripped of resist in an acetone ultrasound bath for 5 min, followed by a 5-min soak in concentrated sulphuric acid/hydrogen peroxide mixture before rinsing in ROH2O and drying in an air gun.

Nickel dies were made directly from the patterned resist samples. A thin (50 nm) layer of Ni-V was sputter coated on the samples. This layer acted as an electrode in the subsequent electroplating process. The dies were plated to a thickness of ca 300 µm. Once returned from the plater, the nickel shims were cleaned by first stripping the protective polyurethane coating using chloroform in an ultrasound bath for 10–15 min. Second, silicon residue was stripped by being wet etched in 25% potassium hydroxide at 80°C for 1 h. Shims were rinsed thoroughly in ROH2O and then
air gun dried. The shims were finally trimmed to approxi-
mately 30 mm × 30 mm sizes using a metal guillotine.

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