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A genomics approach in determining nanotopographical effects on MSC phenotype

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1. Introduction

Current developments in the application of nanotopography have provided us with promising results in the field of regenerative medicine. Major results with mesenchymal stem cells, a key regenerative cell target given their indicated immune-privilege and availability as autologous cells, have included the ability to target osteogenesis using controlled disorder, NSQ50 (pits of 120 nm average diameter, 100 nm deep with a near square arrangement – average 300 nm centre–centre with up to ±50 nm offset in X and Y) indicating that implant modifications may be possible to improve clinical outcome [1–4]. More recently, it was shown that nanostructured surfaces with tightly controlled arrangement, SQ (similar to NSQ50 but with no offset) can retain stem cell phenotype and maintain stem cell growth with implications therein for provision of high quality stem cells to clinic [5]. Furthermore, recent literature has also highlighted the potential for modifying embryonic stem cell response with nanotopography [6,7].

Understanding the mechanism of the physiological processes that control cell–biomaterial interactions and the influence of nanotopography on cell adhesion and phenotype is fundamental to understanding stem cell differentiation. In this study isolated multipotential bone marrow skeletal stem cells also known as mesenchymal stem cells (MSCs), with the potential to differentiate along the stromal lineages, were examined. MSCs can give rise to

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Mechanotransduction
Gene expression
Cell signalling
Molecular biology

Topography and its effects on cell adhesion, morphology, growth and differentiation are well documented. Thus, current advances with the use of nanotopographies offer promising results in the field of regenerative medicine. Studies have also shown nanotopographies to have strong effects on stem cell self-renewal and differentiation. What is less clear however is what mechanotransductive mechanisms are employed by the cells to facilitate such changes. In fastidious cell types, it has been suggested that direct mechanotransduction producing morphological changes in the nucleus, nucleoskeleton and chromosomes themselves may be central to cell responses to topography. In this report we move these studies into human skeletal or mesenchymal stem cells and propose that direct (mechanical) signalling is important in the early stages of tuning stem cell fate to nanotopography. Using fluorescence in situ hybridization (FISH) and Affymetrix arrays we have evidence that nanotopography stimulates changes in nuclear organisation that can be linked to spatially regulated genes expression with a particular focus on phenotypical genes. For example, chromosome 1 was seen to display the largest numbers of gene deregulations and also a concomitant change in nuclear positioning in response to nanotopography. Plotting of deregulated genes in reference to band positioning showed that topographically related changes tend to happen towards the telomeric ends of the chromosomes, where bone related genes are generally clustered. Such an approach offers a better understanding of cell–surface interaction and, critically, provides new insights of how to control stem cell differentiation with future applications in areas including regenerative medicine.

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different lineages including fibroblastic, chondrogenic, myoblastic, adipogenic, and osteoblastic cell types [8–11]. Recent studies have highlighted that MSC function follows form, with alterations in cell adhesion and subsequent cytoskeletal tension modulating lineage commitment [9,12]. There is evidence demonstrating the importance of intracellular tension in MSCs with a high-tension state inducing osteogenic differentiation, whilst a low-tension state inducing adipogenic differentiation [12–14]. Recent advances are indicative of the requirement of an intermediate level of cellular tension for MSC self-renewal [5,15,16].

Interactions between stem cells and the ECM can have indirect or direct effects on cells, otherwise known as mechanotransduction, to elicit changes in gene expression. Indirect mechanotransduction includes the canonical biochemical signalling cascades which result from integrin binding and focal adhesion formation [17]. The second form of mechanotransduction, direct, occurs as a consequence of conformational changes in the cell cytoskeleton, which forms a direct link between the extracellular matrix and the chromosomes via telomeric chromatin/lamin interactions [18,19].

The nucleus itself is supported by the nucleoskeleton consisting of a network of proteins comprising lamin A and C (derived by differential splicing of the same gene) [20], B1 [21] and B2 [22] (products of two genes) in most somatic cell types. The lamina provides structural support to the nucleus, forming part of the link to the cytoskeleton ensuring the correct nuclear and centrosomal organization. The lamina is also associated with DNA replication; lamin B foci are associated with proliferating cell nuclear antigen organization. The lamina is further to the cytoskeleton via the laminae covered with coverslips and incubated for 44 h at 37 °C. The samples were then air dried for 3 days. Cells were used at passages P1–P2 throughout the study.

2.2. Cell extraction and culture

MSCs or skeletal stem cells were enriched from human bone marrow using the STRO1 antibody and magnetic activated cell sorting (MACS) as previously described [2]. MSCs were maintained in basal media (αMEM (PAA)) supplemented with 10% FBS (PAA), 1% v/v) 200 μM l-glutamine (Gibco) and antibiotics (6.74 U/ml Penicillin–Streptomycin, 0.2 μg/ml Fungizone, Gibco) at 37 °C with 5% CO2 in a humidified incubator. MSCs were seeded onto the materials at 1 × 104 cells/ml and allowed to grow for 3 days. Cells were used at passages P1–P2 throughout the study.

2.3. Chromosome territory staining: fluorescence in situ hybridisation (FISH)

MSCs were fixed in 3:1 methanol:acetic acid for 30 min at room temperature and rinsed in 2× SSC (sodium saline citrate; diluted from 2× stock of 3× NaCl, 0.3 M sodium tri-sodium citrate, pH7.4) for 3 h at 37 °C. The appropriate chromosome probe (biotinylated human chromosome 1 paint; Cambio, Cambridge, UK) was brought to 37 °C, vortexed, and pelleted by centrifugation for ~3 s at 11,000 × g. The probe was denatured at 65 °C for 10 min, followed by a 30 min incubation at 37 °C. The samples were rinsed in H2O for 30 s and then dehydrated through a 70%, 90%, 90% (v/v) ethanol series, with a 2 min incubation at each step, followed by a 5 min dehydration step in 100% ethanol. The samples were then air dried for 1 min and incubated in denaturation solution (7:1 formamide: 2× SSC buffer) at 65 °C for 2 min. The samples were quenched using an ice-cold ethanol series as above and air-dried for 1 min. The denatured probe (8–15 μl) was added to each sample, the samples were covered with coverslips and incubated for 44 h at 37 °C in a humidified chamber. Following hybridization, the samples were rinsed in 45 °C pre-warmed 1× SSC buffer for 5 min followed by 2× 5 min washes in stringent wash solution (1:1 formamide: 1× SSC). The probe was detected using the Biotin Printing Kit (Cambio), according to the manufacturer’s protocol. Three replicates of each topography (NSQ50, SQ, FLAT) were used in each experiment.

2.4. Territory analysis

The distances from the nearest edge of the nuclei to the centres of the chromosomal territories and the interterritory distances were measured using ImageJ (version 1.34a; Rasband, W.S., Image J, U.S. National Institutes of Health – http://rsb.info.nih.gov/ij/). Statistics were generated using Prism (Graphpad at www.graphpad.com/prism) the Tukey–Kramer multiple comparisons post-test analysis of variance (ANOVA).

2.5. Affymetrix arrays

MSCs were cultured on the topographies (4 material replicates/biological replica/topography) for 3 days. At this point, the cells were lysed and total RNA was extracted using a Qiagen RNeasy micro kit (Qiagen, UK). Gene expression changes were detected by hybridization of mRNA to Affymetrix HuGene 1.0 ST human arrays according to the manufacturers instructions. Initial bioinformatic analysis was based on rank product. A false discovery rate of 20% was used to upload selected genes changes to the Ingenuity Pathway Analysis (IPA) server to identify canonical signalling pathways, functional pathways and to produce networks. Statistics for functional analysis were carried out by Fischer’s exact test (automatically performed by the software). For the chromosomal band identification a custom script was written to add annotations to the ANOVA results file. The script generated a transcript cluster ID to the chromosome lookup table from the HuGene-1.0-st-v1.lna32 hg19.probeset.csv file obtained from Affymetrix (www.affymetrix.com). Then the results file was parsed and the chromosomal location details from the lookup table, were appended to the corresponding transcript.

2.6. Quantitative real time (q)PCR

MSCs were cultured on topographies for 3 days (4 biological replicates each consisting of 3 replicates each pooled) for each NSQ50, SQ and FLAT) at a density of 1 × 104 cells/ml. Total RNA was extracted using a Quiagen RNeasy micro kit. Real-time qPCR was carried out and analysed as previously described to assess the expression of Runx2, HOPE2S, ALCAM, SOX9 and PPar (Tables 1 and 2). RNA samples were reverse transcribed using the Omniscript First Strand System (Qiagen). Real-time qPCR was carried out using the 7500 Real Time PCR system from Applied Biosystems. GapDH

2. Materials and methods

2.1. Nanopatterning and mastering

The substrates were made in three-step process of electron beam lithography [25] nickel die fabrication and polymer replication using injection moulding. Briefly, the master substrates were fabricated to form an array of 120 nm diameter pits of 100 nm depth and 300 nm pitch in a square (SQ) arrangement with the near square (NSQ50) substrate has a random displacement of ±50 nm, and maintaining an average 300 nm pitch. Nickel dies were made directly from the patterned resist samples and a thin (50 nm) layer of Ni–V was sputter coated on the samples, acting as an electrode in the subsequent electroporation process. The dies were plated to a thickness of approximately 300 μm. The nickel shims were cleaned by stripping the protective polyurethane coating using chloroform in an ultrasound bath for 15 min. An injection mould was used to make polymer replicas in polycarbonate.
served as the house-keeping gene to normalise expression for the genes of interest. In cases where the SYBR Green method was used (Table 1), primer sequences for the genes (GapDH and ALCAM) were validated by dissociation curve/melt curve analysis.

### Table 1: qPCR primer details for SYBR green.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALCAM</td>
<td>ACCTGACCTGACGGCCTT</td>
<td>CCCTGGCTGGTCTT</td>
</tr>
<tr>
<td>GapDH</td>
<td>GTCACTGGTCTGAGCCAAGT</td>
<td>ACCTGGTCTGAGCAAGC</td>
</tr>
</tbody>
</table>

### Western blotting

MSCs were cultured on the topographies for 3 days (4 biological replicas (each consisting of 4 replicas each pooled) for each NSQ50, SQ and FLAT) at a density of 1 x 10^5 cells/mL. MSCs were lysed using protein lysis buffer (20 mM Tris—HCl, pH 7.5, 150 mM NaCl, 1 mM EDTA, 1% v/v Triton X-100) containing phosphatase and protease inhibitors (1% phosphate inhibitor cocktail Roche cat. no. 04906845001, 1% Sigma protease inhibitor cocktail, cat. no. P2714). Proteins were run on a pre-cast NOVEX gradient (4–12%) gel system (Invitrogen) and the samples were then transferred onto a nylon membrane (Immobilon P. Millipore), according to the manufacturer’s protocol. For probing, the blots were incubated in 5% non-fat milk PBS 0.1% (v/v) Tween 20 with the appropriate anti-sera dilution. Antibodies (with dilutions) used were directed to: osteogenic markers Phospho-Runx2 (Abgent, AF5559a) 1:500 total Runx2 (Santa Cruz Biotechnology, sc-10758) 1:500, stem cell marker STRO1 1:500 (R&D systems, MAB1038), adipocytic marker PPary (Santa Cruz Biotechnology, sc-1984), chondrocytic marker Sox9 (Abcam, ab97697), housekeeping gene GapDH (Sigma, G8795); followed by the appropriate 1:4000 goat anti-rabbit, goat anti-mouse or donkey anti-goat IgG HRP-conjugates (Santa Cruz). Detection was performed by enhanced chemiluminescence (Immobilon Western, Millipore).

### 3. Results

#### 3.1. Nuclear organization

Microarray analysis was used to identify potential 'hotspots' for differential gene expression in the genome of MSCs grown on the different nanotopographies and to contribute to the rational selection of specific chromosomes. At the chromosomal level, chromosome 1 (Ch1) displayed the largest number of differentially expressed genes (up and down regulated) in cells cultured on both NSQ50 and SQ (Fig. 1A and B). Overall NSQ50 displayed the most changes (Supplementary Fig. 1) indicating the higher impact this topography has on MSC gene expression and cell growth. In general, both topographies displayed the most regulation changes on predominantly on the larger chromosomes (such as Ch1, Ch2, Ch5, Ch6, Ch12, and ChX) indicating a trend between size and response to nanotopography.

While Ch17 and Ch19, smaller chromosomes, showed deregulations in response to nanotopography, this could, speculatively, be attributed more to biochemical changes over the direct mechanotransduction.

Based on the above observations, Ch1 was investigated further by FISH in order to examine the involvement of chromosomal territory shifts on differential gene expression (Fig. 1C). Distances from the nearest edge of the nucleus to the centre of chromosomal territories were measured for Ch1. Quantification of Ch1 territory position relative to the nuclear periphery was significantly altered (p < 0.01, ANOVA) for cells cultured on the NSQ50 compared to those on the planar control and SQ (p < 0.01, ANOVA) although chromosomal territory position was comparable between SQ topography and Flat control (Fig. 1D). In addition, the Ch1 interterritory distance in cells on the NSQ50 topography was statistically significantly larger (p < 0.05) compared to those on the Flat control and (p < 0.01) when compared to those on the SQ (Fig. 1E).

Further evidence supporting our hypothesis that topographical differences affect the nuclear physiology and organization with concomitant chromosomal shifts are observations of nucleus size differences between cells on the NSQ50 in comparison to cells on the SQ nanotopography and Flat control (Supplementary Fig. 2).

The microarray data was used to identify the chromosomal band positions of differentially expressed genes. Band positions were assayed using Z-score analysis. Z-score measures the probability of the observed numbers of changes, occurring at a particular chromosomal band position with higher scores in either direction indicating that the observed number of gene expression changes are most likely true.

The results from the q-arm data showed that the topography-related changes for both NSQ50 (Fig. 2) and SQ (Fig. 3) were broadly spaced along Ch1 with a trend of activity towards the telomeric (end) region of the chromosome. This was apparent for most of the larger chromosomes (1–15), where analysis showed points of loss of up-regulation and enhanced down-regulation at the telomeric regions (band 20 and above). Interestingly, there were a number of ‘gains’ of gene up-regulation associated with ‘loss’ of gene down-regulation at the centromeric regions (band 10–15) of chromosomes 16 and above.

Looking at the p-arm analysis results for MSCs on NSQ50 (Supplementary Fig. 3), for the largest chromosomes (1–5), analysis showed points of loss of up-regulation (→) and many more substantial gains of down-regulation (→) at predominantly the centromeric regions (bands 11–15) and less at the telomeric regions (band 21 and above). Similar profiles were obtained for chromosomes 6–10. Furthermore, there were a large number of ‘gains’ of gene down-regulation associated with ‘loss’ of gene up-regulation at the centromeric regions of chromosomes 11 and above.

SQ p-arm analysis results, for the largest chromosomes (1–5), showed points of loss of down-regulation (→) and many more substantial gains of up-regulation (→) at the telomeric regions (band 20 and above) (Supplementary Fig. 4). Interestingly, there were a large number of ‘gains’ of gene up-regulation associated with ‘loss’ of gene down-regulation at the centromeric regions (bands 10–15) of chromosomes 6–15. In addition, there were a large number of ‘gains’ of gene down-regulation associated with ‘loss’ of gene up-regulation at the centromeric regions of chromosomes 16 and above.

#### 3.2. Correlating morphology to phenotype

In an effort to correlate morphology to phenotype, protein and gene expression studies were performed. The mean protein expression levels in MSCs grown on NSQ50 (osteogenesis promoting) and SQ (self-renewal supporting) nanotopographies were investigated using Western blotting (Fig. 4A) and gene expression was studied by qPCR (Fig. 4B). The markers used, typically associated with multipotent skeletal or mesenchymal stem cells, were STRO1, HOP26, and ALCAM and differentiation was indicated by tissue specific markers Runx2 (runt related...
transcription factor 2, indicative of initiation of osteogenesis), soft-tissue markers SOX9 (Sry-related high mobility group box 9, a cartilage specific transcription factor) and PPARγ (peroxisome proliferator-activated receptor gamma, an adipocyte-related transcription factor). NSQ50 supported the expression of the osteogenic marker Runx2 in both active (phosphorylated) and total protein forms (Fig. 4A) as well as at the gene transcript level (Fig. 4B) at considerably higher levels than for cells on flat controls and SQ topography (p < 0.05). Examination of soft tissue differentiation markers such as adipogenic marker PPARγ and bone stem cell marker STRO1 showed a reduction at both gene and protein levels of all these markers on the NSQ50 (osteogenic differentiation promoting topography) in comparison to SQ and Flat control (Fig. 4A and B). The chondrogenic marker SOX9 showed reduced gene (p < 0.05) levels in MSCs on NSQ50 with no significant change at protein level noted. For the SQ, self-renewal promoting, nano-topography, statistically significant (p < 0.01) increased protein levels of STRO1 and gene levels of HOP26 and ALCAM (primers

Fig. 1. Nanotopography effects on chromosome 1 (Ch1) territory repositioning within the MSC nucleus. (A) and (B) Microarray was used to select chromosomes that showed either few or many transcript abundance changes. Examples NSQ50 (A) and SQ (B) MSC cultures. Ch1 had the greatest total number of expression changes (mostly down-regulated genes). (C) FISH staining for chromosomal territory positioning. Examples of FISH staining of Ch1 on flat, NSQ50 and SQ in MSC nuclei. Key: Blue – DNA; Green: Ch1 territories. (D) Chromosome territory positioning within the nucleus. For chromosome 1, quantification of territory position relative to the nuclear periphery was significantly altered (p < 0.01) for cells cultured on NSQ50 compared to those on the flat control and SQ. (E) For chromosome 1, quantification of inter-territory position was significantly larger (p < 0.05) for cells cultured on NSQ50 compared to those on the flat control and (p < 0.01) when compared to those on the SQ. Comparison was done by ANOVA *p < 0.05, **p < 0.01, ***p < 0.001, n = 45. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Fig. 2. NSQ50 Z-score analysis of gene activity associated with q-arm band number. The graphs show significant differences from expected changes based on analysis of 1600 iterations of a ‘window’ of genes that were selected using a threshold of 15% for the false discovery rate. Data is shown NSQ50:Flat (up-regulations/down-regulations refer to the NSQ50 topography, relative to the Flat surface). For the largest chromosomes (1–5), analysis showed points of loss of up-regulation (→) and many more substantial gains of down-regulation (←) at the telomeric regions (band 20 and above). Profiles were similar for chromosomes 6–15. Interestingly, there were a large number of ‘gains’ of gene up-regulation associated with ‘loss’ of gene down-regulation at the centromeric regions (band 10–15) of chromosomes 16 and above.
Fig. 3. SQ Z-score analysis of gene activity associated with q arm band number. The graphs show significant differences from expected changes based on analysis of 1600 iterations of a 'window' of genes that were selected using a threshold of 15% for the false discovery rate. Data is shown SQ:Flat (up-regulations/down-regulations refer to the SQ topography, relative to the Flat surface). For the largest chromosomes (1–5), analysis showed points of loss of up-regulation (→) and many more substantial gains of down-regulation (←) at the telomeric regions (band 20 and above). Profiles were similar for chromosomes 6–10. Interestingly, there were a large number of 'gains' of gene down-regulation associated with 'loss' of gene up-regulation at the telomeric regions of chromosomes 11–15. Profiles were very similar for the small chromosomes (16–22).
cannot yet be designed for STR01 in the absence of epitope data so HOP26 and ALCAM genes were used as alternatives) were observed in agreement with Refs. [5,16].

4. Discussion

In the field of regenerative medicine, understanding the stem cell nano/microenvironment is of primary importance. The elucidation of the cell—nanotopographical interaction and its effects on cell morphology and phenotype will provide new insights into the regulation of stem cell differentiation and self-renewal processes. It has previously been shown [1] that a slightly disordered nanotopography NSQ50 promotes osteogenic differentiation and, more recently, we demonstrated that a highly ordered nanotopography, SQ, supports the human skeletal stem cell phenotype [5].

Here, we confirm and extend these results with protein and gene expression analyses to study the differentiation status of MSCs on the two nanotopographies. NSQ50 supported expression of the osteogenic marker Runx2 but not soft tissue marker such PPARγ, SOX9 and STR01. In contrast, SQ (low tension) supported expression of soft tissue markers, especially STR01 but not Runx2. Comparison was done by ANOVA *p < 0.05, **p < 0.01, ***p < 0.001, n = 3. Note: markings above the error bars denote comparison to flat control; comparison between SQ and NSQ50 is denoted by a line.

Fig. 4. Phenotypic changes in response to nanotopography. A). Mean protein expression levels in MSCs grown on NSQ50 and SQ nanotopographies expressed as ratio of the flat control. B) Mean gene expression levels in MSCs grown on NSQ50 and SQ nanotopographies expressed as ratio of the flat control. NSQ50 (high tension) supported the expression of osteogenic marker Runx2 but not soft tissue marker such PPARγ, SOX9 and STR01. In contrast, SQ (low tension) supported expression of soft tissue markers, especially STR01 but not Runx2. Comparison was done by ANOVA *p < 0.05, **p < 0.01, ***p < 0.001, n = 3. Note: markings above the error bars denote comparison to flat control; comparison between SQ and NSQ50 is denoted by a line.

that other studies have implicated nanoscale order and disorder as having large effects on MSC attachment and adhesion based on ability to gather integrins into focal adhesions [27].

Cell—extracellular matrix or cell—surface interactions trigger cascades of signals via membrane proteins at points of adhesion that are transmitted through the cytoskeleton to the nucleus and determine stem cell homing, proliferation and differentiation processes [28]. It is known that as the cytoskeleton is linked to adhesions and also to the nucleoskeleton via LINC (linkers of nucleoskeleton to the cytoskeleton) complexes (possibly facilitated by tensegrity) that alterations in nucleus morphology will follow rearrangements of the cytoskeleton and we have produced preliminary evidence for this in fibroblasts [16,29–31]. It has been demonstrated by others that high tension and cell spreading are essential for the differentiation of MSCs to osteoblasts [9,13]. Here we postulate that the effects may be more than simply biochemical with direct application of tension to the nucleus (based also on nuclear morphology observations in Supplementary Fig. 2) and coupling of the chromosomal telomeres to the nuclear lamina resulting in changes in chromosomal positioning altering DNA accessibility to transcription factors. Such a mechanism of action has also been postulated in response to changes in serum concentration [32].

For the SQ, self-renewal, surface, much smaller (compared to NSQ50) differences to the flat control were seen in chromosomal positioning. Here, we observed expression of soft tissue markers such as SOX9 (mapped on chromosome 7p-arm band 21), PPARγ (12p13), HOP26 (17q11) that are mainly located on smaller
chromosomes at more centromeric positions. We propose smaller chromosomes are less tension-aﬀected as previous studies with ﬁbroblasts also indicate that smaller chromosomes are less responsive to topography. In addition, it is important to note that it is the telomeres that are attached and thus mechanically connected to the nuclear lamina [1,31]. This is perhaps suggestive of more biochemical control rather than direct mechanical control of the genes when situated at lower band numbers. We have previously suggested that small RNAs could be important in MSC self-renewal and this is supportive of a biochemical rather than mechanical control mechanism [5].

However, on our osteogenic, NSQ50 surface, highly signiﬁcant changes in chromosomal positioning and more signiﬁcant telo-
meric changes in regulation were noted. We observed expression of the osteogenic gene Runx2 (mapped on 6p21 — i.e. large chromo-
some, telomeric) but reduced expression of the soft tissue markers. Interestingly, the changes in telomeric gene down-regulations, in
MSCs on the NSQ50 topography and the movement of larger terri-
ories may contribute to osteogenic diﬀerentiation [33].

Literature review provides evidence further supporting that
direct mechanotransduction could be a signiﬁcant part of osteo-
genesis as a number of signiﬁcant osteospeciﬁc genes (osteocalcin at 1q25 — 31, osteopontin at 4q22, osteonectin at 5q31 and alkaline
phosphatase at 2q37) are located at the more telomeric band positions of the larger chromosomes [31,34]. Combined with our
new data, this suggests that the Runx2 osteogenic ‘master’ tran-
scription factor and the genes Runx2 regulates are located in ten-
sionally sensitive areas of the genome. This further demonstrates the tension sensitivity of the osteogenic lineage.

5. Conclusions

In the current study we have demonstrated the potential of
a non-invasive materials approach to understanding mechanisms underlying stem cell growth and diﬀerentiation. Analysis of the architectural and molecular eﬀects of the two diﬀerent nanotopographies on MSC nucleus organisation show modulation of chromosomal positioning and spatially-related gene changes during osteogenesis but not self-renewal. These studies oﬀer new
insight into the diﬀerentiation of MSCs on diﬀerent nano-
topographies and the implications therein for modulation of skeletal function and activity. The wider implications for stem cell fate regulation in other systems (foetal, embryonic, soft tissues) are under investigation in our laboratories.

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Appendix A. Supplementary material

Supplementary material associated with this article can be
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