Creation of in vitro disease models using micro-nanoengineering

Allison Bruce

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Creation of *in vitro* Disease Models Using Micro-/Nanoengineering

by

Allison Bruce

Thesis Submitted to the
Benjamin M. Statler College of Engineering and Mineral Resources
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in partial fulfillment of the requirements for the degree of

Master of Science in Engineering

Yong Yang, Ph.D., Chair
Cerasela-Zoica Dinu, Ph.D.
Charter D. Stinespring, Ph.D.

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Abstract

Creation of in vitro Disease Models Using Micro-/Nanoengineering

Allison Bruce

Cells are typically housed in a complex microenvironment (ECM) in vivo, where the cells interact with the extracellular matrix (ECM) and neighboring cells. The ECM provides physical support for cells and affects cell adhesion, migration, proliferation, differentiation, and gene expression. Primary components of the ECM include fibrous proteins which contribute significant nanotopography cues to cells. Due to limitations of two-dimensional (2-D) in vitro studies where cells are cultured on flat, plastic substrates, the conventional cell culture does not mimic the in vivo scenario. This manifests the critical need for biomimetic cell culture platforms which mimic the in vivo microenvironment. During my graduate studies, two in vitro disease models were constructed to investigate cell-substrate and cell-cell interactions.

In Chapter I, nanotopographical effects on cell responses were examined. Focal adhesions and nuclear deformation of normal human lung fibroblasts (NHLFs) were investigated. In general, nanoscale gratings hindered cell spreading and thus reduced cell size while increasing the nuclear volume. On the other hand, nanoscale pillars, depending on the feature size and spacing, might modulate the focal adhesions and nuclear size towards opposite directions. Our observations suggested that it was focal adhesion area instead of focal adhesion size that affected nuclear deformation. Therefore, nanotopography could be optimized to modulate cell adhesion and nuclear volume, which would provide a useful tool to regulate cell phenotypes and functions for end applications.

In Chapter II, the effects of cell-cell and cell-matrix interactions on tumor cell survival within an engineered bone marrow microenvironment for Acute Lymphoblastic Leukemia (ALL) were evaluated. In this study, biologically relevant populations of primary human derived bone marrow stromal cells (BMSCs), osteoblasts and a human ALL cell line representative of an aggressive phenotype were used. Traditional 2-D co-culture, 3-D static culture in collagen, and 3-D cultures with flow were evaluated to determine response to a commonly used chemotherapeutic agent, Ara-C. 3-D co-culture models showed higher survival of tumor cells during Ara-C exposure as well as enhanced protection during chemotherapy stress conferred by microenvironment cells.
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I. Nanotopography Modulated Nuclear Deformation and Focal Adhesions

1. Introduction

1.1 Extracellular Matrix

Regenerative medicine, aiming at the repair or replacement of cells, tissues and organs [1], holds great promise for the treatment of diseases that are untreatable at present, such as various gene and neurological disorders, cardiovascular diseases, as well as hematological malignancies. Realization of this potential remains limited by current challenges associated with the control of cell phenotypes and functions in cell culture [2]. In this respect, the fate of cells is dictated by the in vivo microenvironment where these cells interact with both the extracellular matrix (ECM) as well as with neighboring cells. The ECM serves as a structural support for cells and provides, in concert with the spatio-temporally arranged signaling molecules and external stimuli, topographical and mechanical cues for cell adhesion, spreading, migration, proliferation and differentiation.

The ECM is composed of diverse biomacromolecules including glycosaminoglycans, fibrous proteins such as collagen, elastin, and fibronectin, and nonfibrous proteins such as growth factors and cytokines, with size ranging from several to hundreds of nanometers [3]. The ECM constructed from these biomacromolecules often includes significant topography at the nanoscale level. For example, basement membranes, which are a ubiquitous component of ECM that plays an important role in tissue development and organization, manifest a complex three-dimensional (3-D) texture with sizes in the nanometer range [4-9].

1.2 Nanotopographical Effects on Cellular Responses

A number of recent findings strongly support that cell responses are highly sensitive to topographical cues on a synthetic surface, in particular, at the nanoscale level [10-13]. In addition to inducing pronounced changes to cell morphology, migration, and proliferation, topographical cues could potentially help induce the differentiation of stem cells into certain lineages such as neuron [11,14,15], muscle [16], and bone [12,13].
Cellular responses to external cues begin with formation of focal adhesions via binding and clustering of integrins onto the adhesive ligands present in the substrate. Formation of focal adhesions and actin stress fibers control the adhesion and spreading of cells. As a dynamic process, focal adhesions are constantly assembling and disassembling to enable motility and cellular migration [17]. The shape, size, spacing and depth of nanoscale features have profound influence on cell morphology, spreading and migration. The use of nanotopography can dramatically enlarge the surface area, thus enhancing cell adhesion and proliferation [18,19]; however, the apparent surface that cells can ‘sense’ is determined by the dimensions and geometry (e.g., gratings, pillars, and pits) of nanotopography relative to cell dimensions [12,13,20,21]. For instance, human fibroblasts responded differently to demixed isotropic nanoislands of 13, 35 and 95 nm in height. Fibroblasts displayed highly spread morphology containing well-defined cytoskeleton on the 13-nm islands, but had deformed cell spreading with diffuse actin and less stress fibers when the height of the islands increased [22,23]. Human endothelial cell also showed a highly spread cell morphology on the 13-nm islands [22]. However, on the 95-nm islands, different from fibroblasts, endothelial cells appeared to have larger lamellae, the edges of which appeared to be anchored to the nanoislands and had increased numbers of stress fibers [24].

Anisotropic nanotopography could guide cells to spread and migrate preferentially. Teixeira et al. studied human corneal epithelial cells on silicon (Si) nanogratings with a linewidth ranging from 70 to 1900 nm, with a corresponding pitch of 400 to 4000 nm and two depths of 150 nm and 600 nm. The cells cultured on the smooth control formed focal adhesions and stress fibers with no preferred orientations. On the nanogratings focal adhesions and stress fibers aligned along the nanogratings direction. The focal adhesion widths on any of the nanogratings were smaller than those on the smooth control. The size of focal adhesions increased with the ridge width up to 400 nm and remained constant when the ridge width was beyond 650 nm. On all 600-nm deep grooves, cells aligned along the gratings had higher elongations than cells on the smooth control. Cell alignment resulted from anisotropic cell spreading, as the cells extended and retracted lamellipodia preferentially along the direction of the patterns. They observed that
the average cell areas on all nanogratings were lower than those on the smooth control [20]. The alignment induced by nanotopography might be reduced with an increase in grating width. Up to 90% of the smooth muscle cells (SMCs) would align with nanogratings (350 nm and 500 nm in width) but only 37% of the cells would do so when cultured on 10 µm wide ratings. Greatest alignment was observed on gratings of 800 nm height, and decreased as the height goes down to 300 nm [25]. Nuclear alignment and deformation could also be affected by the anisotropic nanotopography. On polydimethylsiloxane (PDMS) nanogratings (350 nm in width, 700 nm in pitch and 280 nm in depth), 62.0% of nuclei of human mesenchymal stem cells (hMSCs) oriented in a 15° alignment angle compared to a random orientation of the nuclei on the flat control. The nanogratings elongated the nuclei to an elongation aspect ratio (nuclear long axis/short axis) range of 1–5 compared with the range of 1–3 on the flat control. The nanogratings also significantly decreased the average nuclear area to 145.1 ± 4.1 µm² on nanogratings from 194.8 ± 4.8 µm² on the flat control [26].

When the topography reduces the apparent surface area on which cells can sense, the topography restricts the focal adhesions, weakens cellular adhesion, and thus facilitates cell migration [27]. On 350-nm gratings hMSCs exhibited a much smaller size of focal adhesion protein zyxin, 3.2 ± 0.26 µm², compared to 5.3 ± 0.55 µm² in the cells on planar controls. Fluorescence recovery after photobleaching (FRAP) measurements indicated that the turnover of GFP-zyxin proteins was increased on 350-nm gratings. Since the association of zyxin with focal adhesions was force-dependent, smaller zyxin-positive adhesion as well as its higher turnover rate suggested that the traction force in focal adhesions on 350-nm gratings was decreased. These changes led to faster and more directional migration on 350-nm gratings, 15.6 µm/h, compared with 8.3 µm/h observed on planar controls [28].

There is a general notion that proliferation increases with cell spreading. Nanotopography, very often, reduces the surface area cells can sense. Hence, a decrease in proliferation has been observed in cells cultured on nanogratings although the cell alignment, elongation and migration are enhanced [11, 24, 29-33]. For instance, on PDMS nanogratings (350 nm in width, 700 nm in pitch, and 350 nm in depth)
cytoskeleton and nuclei of hMSCs aligned and elongated along the nanogratings. The proliferation of hMSCs on the nanogratings was significantly lower, 26.9 ± 3.1% of BrdU incorporation compared to 35.7 ± 7.6% on flat surfaces [11]. Moreover, Dalby et al. have shown that the near-square nanopits stimulated hMSCs to produce bone mineral in vitro, in the absence of osteogenic supplements at levels similar to the cells cultured with osteogenic media [12]; while highly ordered nanopits permitted prolonged retention of multipotency of hMSCs [34].

1.3 Nuclear Deformation
External cues from cellular microenvironment may initiate cell adhesion, and can be transduced through cytoskeletal filaments and nuclear scaffolds to nucleus, ultimately influencing cell fate and functions [35]. In response to the external cues, the molecular connections between cell adhesion, cytoskeletal filaments and nuclear scaffolds may produce integrated changes in cell and nuclear structure [36]. Cell morphology and cytoskeleton organization influence cells via actin stress fibers [37-39]. Alteration in the actin-mediated tension affects cell spreading, contraction, and migration, thus modulating nuclear shape and plasticity. For instance, the differentiation of stem cells into a chondrocytic phenotype required a rounded cell shape. Direct comparison of cell and nuclear shape of hMSCs indicated that a more rounded nuclear shape was associated with the greatest expression of molecular markers associated with chondrogenesis [40]. Nuclear deformation can lead to conformational adaptation in chromatin structure and organization, which in turn affects transcriptional regulation [41] and then gene expression and protein synthesis [35,42,43], eventually leading to the changes in proliferation, differentiation, or cell death [44,45]. Nuclear deformation strongly depends on the spatial organization and orientation of the actin cytoskeleton. The transmission of mechanical cues from the cytoskeleton to the nucleus may not only depend on cytoskeletal tension, but also on the orientation of the cytoskeletal filaments and the way the filaments are connected to the nucleus [46]. In particular, nuclear volume regulation has been identified as a key mechanism by which cell shape controls the expression of different genes, including those that control the cell cycle [46].
1.4 Focal Adhesion

Nanotopography can rearrange focal adhesion distribution and cytoskeleton assembly, which regulates cell and nuclear shape and polarization. The spatial confinement of focal adhesions, ultimately modulation of cell and nuclear shape, are crucial in the context of mechanotransduction [47]. It has been reported recently that the nuclear volume of human corneal epithelial cells grown on nanogratings increased as the pitch of the nanogratings increased from 400 nm to 2000 nm and there was a significant decrease in the cell modulus at 400 nm compared to the moduli for flat surfaces and for larger pitches of these chemically identical substrates [48].
2. Objective

The objective of this project was to examine nanotopographical effects on cell responses. In order to investigate nanotopography effects on cells, isotropic nanopillars and anistropic nanogratings were designed with dimensions above and below the accepted threshold of 500 nm with varying spacing. Human lung fibroblasts were cultured on this platform and focal adhesions and nuclear deformation were analyzed. Using this model, we planned to correlate focal adhesion size to nuclear deformation to establish optimal topography, thus, providing a tool to regulate cells by modulation of cell adhesions and nuclear volume.
3. Materials and Methods
3.1 Nanofabrication

Nanotopographies including gratings and posts were written in a poly(methylmethacrylate) (PMMA) thin film that was spin-coated onto a Si substrate using electron beam lithography (EBL).

A mixture of polydimethylsiloxane (PDMS) resin and curing agent (Sylgard 184 kit, Dow Corning, MI, USA) in a 10:1.05 w/w ratio was poured onto the EBL mold. After curing for 2 hr at 70°C, the inverse PDMS mold was peeled from the EBL mold.

Multiple PDMS molds were stitched to generate a large area of nanopatterned surface as a substrate so that enough cells could be seeded for subsequent biological analyses as reported previously [39]. Briefly, these PDMS molds with the nanopattern face-down were aligned on a Si wafer. A PDMS prepolymer layer spin-coated on a glass plate was pressed onto the unpatterned side of PDMS molds. The assembly was inverted, placed on a hot plate, and cured at 80°C for 2 hr. The multiple PDMS molds were stitched into a single mold. The stitched mold was then hot-embossed into polystyrene (PS) substrates, which served as a master mold to replicate PDMS working substrates. The PDMS substrates nanotopographies and the flat PDMS as the control were sterilized by using ethanol and then UV exposure, each for 30 min. Prior to cell seeding, the PDMS substrates were incubated with human fibronectin (human, BD Biosciences, 10 μg/ml in PBS) for 15 min.

3.2 Cell Culture

Normal human lung fibroblasts (NHLFs, CRL1490 cells from ATCC, Manassas, VA) were cultured in 2% fetal bovine serum (FBS), 0.1% recombinant human fibroblast growth factor basic (rhFGF-B), 0.1% insulin, and 0.1% gentamicin/amphotecin-B (Lonza). The NHLFs used in the experiments were at passages 3–6.

NHLFs were starved in serum-free medium for 18 hr to induce cell cycle synchronization [53]. Cells were allowed to recover in complete fibroblast medium for 1 hr. The cells were seeded at a density of 2000 cells/cm² and placed in an incubator for 12 hr to allow the cells to adhere.
3.3 Flow Cytometry

Cells were harvested as a single cell suspension in PBS and fixed in cold 70% ethanol for 1 hr on ice. Cells were then washed and resuspended in 0.2% Tween 20 in PBS for 15 min at 37°C. Cells were washed with 2% fetal bovine serum in PBS and resuspended in 1.8% RNase-A in PBS for 15 min at room temperature. Propidium Iodide (50 ug/ml) in PBS was added for 15 min at room temperature. After incubation, the volume of each sample was brought up to 500 ul each and analyzed by flow cytometry.

3.4 Scanning Electron Microscopy (SEM)

The PDMS substrates containing cells were washed with PBS and fixed for 4 hr in 4% paraformaldehyde and 2% glutaraldehyde at ambient. They were then rinsed with distilled water and progressively dehydrated using a Balzers Critical Point CO₂ Dryer, followed by freeze-drying in a graded series of ethanol. Both the PDMS substrates with and without cells grown were sputter-coated with a gold layer of ~10 nm thick using a Denton Vacuum Desk V sputter unit (Denton Vacuum, LLC, Moorestown, NJ, USA). SEM micrographs were obtained from a JEOL JSM-7600F Scanning Electron Microscope.

3.5 Atomic force microscopy (AFM)

AFM characterization was performed using an Asylum MFP-3D Atomic Force Microscope in the tapping mode under ambient conditions. Topography images were recorded simultaneously at the fundamental resonance frequency of the cantilever, with a typical scan rate of 1 Hz and a resolution of 512 samples per line.

3.6 Immunofluorescence Staining

The cells were fixed with 4% paraformaldehyde in PBS for 15 min at room temperature, and permeabilized in a blocking solution, which consisted of 0.03 g/ml bovine serum albumin (BSA, Sigma-Aldrich Co.) and 0.1% goat serum (Sigma-Aldrich Co.) in 0.2% Triton X-100 (Sigma-Aldrich Co.) in PBS for 30 min. Focal adhesions were stained by using anti-paxillin antibody (1:200; Abcam, Cambridge, MA, USA) followed by the Alexa Fluor 555 goat anti-rabbit secondary antibody (1:200; Invitrogen, Carlsbad, CA, USA). F-actin was stained with Alexa Fluor 488 phalloidin (Invitrogen, Carlsbad, CA,
USA), and the cell nuclei were stained and mounted using ProLong Gold Antifade Reagent with DAPI (Life Technologies) and viewed with Zeiss LSM 510 inverted confocal microscope.

For the nuclear volume measurement, z-stack imaging with 0.1 μm intervals between planar images was conducted. To validate the accuracy of z-stack imaging of nuclei, FITC-labelled beads (diameter: 15.4 um ± 0.13 um, BD Bioscience) were used as the standard.

3.7 Image Analysis
Focal adhesion size and nuclear area and volume were analyzed using Imaris. Briefly, creating superficial surfaces overlaying the image; the threshold was set to the TRITC channel and was adjusted to fit the parameters of the confocal image; particles of sizes between 2 and 10 μm$^2$ were extracted using a filter and analyzed.

3.8 Statistical Analysis
Focal adhesion size and nuclear deformation in terms of projected area and volume of NHFLs were presented as mean ± standard error. Approximately 150-200 nuclei images from each of three replicates were collected for each condition. Approximately 1000 focal adhesions for each condition were analyzed from 15-20 cells from each nanotopographies including the flat control. The differences between groups were analyzed by one-way ANOVA and Dunnet’s method, with P<0.05 taken as statistically significant.
4. Results and Discussion
4.1 Design of Nanotopographies

Nanoscale gratings and pillars were of interest because they represented two major nanotopographies. Nanogratings (NGs) would provide anisotropic guidance to cells, while hexagonally arranged nanopillars (NPs) were isotropic. The lateral dimensions of nanotopographies were controlled by EBL writing and the feature height was defined by the thickness of the PMMA thin film. The linewidth of nanogratings were 300, 500 and 1000 nm and the edge-to-edge spacings were 1 and 3 times the linewidth for each design (Fig. 1.1A). The pillars had the diameter of 300, 500, 1000 nm and a center-to-center spacing of 1.3 and 1.9 times the diameter. The corresponding edge-to-edge distances were 90, 150 and 300 nm for the 1X spacing pillars and 270, 450 and 900 nm for 3X spacing (Fig. 1.1B). The height of all nanotopographies was 170 nm as measured by using AFM (Fig. 1.1C). To facilitate the discussion, each nanotopography was named by using its abbreviation (NG or NP) followed by a combination of the feature size (linewidth for nanogratings and diameter for nanopillars) and the spacing. For instance, ‘NG 500-3X’ stood for the nanogratings of 500 nm in linewidth and 1500 nm for spacing, and ‘NP 500-1.9X’ for the nanopillars of 500 nm in diameter and 950 nm for the center-to-center spacing (or 450 nm for the edge-to-edge distance). For each feature size, the spacing was chosen to keep the ratio of the nanostructured surface area over the total area of the corresponding flat surface constant. For example, when the spacing of nanogratings was the same as the linewidth the ratio was 0.50. The increase of the spacing of nanogratings from 1X to 3X the linewidth decreased the ratio to 0.25. Similarly, the increase of the center-to-center spacing from 1.3X to 1.9X the diameter decreased the ratio from 0.50 to 0.25. Hence, the featured surface area could be kept constant so that the cells may be exposed to the same seeding area but at the same time to different topography.
Figure 1.1: A) SEM micrographs of (A) nanogratings, (B) nanoposts, and (C) AFM images of typical nanotopographies. Two-dimensional profile indicated the feature height.
4.2 Spacing-dependent Cell Alignment on Nanogratings

Normal human lung fibroblasts (NHLFs) were serum starved for 18 hr to give rise to a larger than 92% population of cells possessing a synchronized nuclear cycle, G1 phase, substantially reducing the variation in starting nuclear volume. After the cell adheres onto the substrate, increasing intracellular tension may mature early stage focal complexes into focal adhesions, which can reach several micrometers in length. It was reported that focal adhesions of MC3T3-E1 preosteoblasts grew in length from 4 hr to 12 hr culture [47]. We were interested in the early-stage cell-nanotopography interactions and hence investigated the cell responses at 12 hr. Shown in Fig. 1.2, NHLFs displayed multipolar morphology on the nanopillars, similar to those on the control flat surfaces; while the cells might orient along the nanogratings direction.

![Figure 1.2: Nanotopography-dependent cell spreading.](image)

The nuclei were stained with DAPI in blue, the actin filaments were stained with phalloidin in green, and focal adhesions were stained with paxillin in red.
It has been observed that nanogratings could provide contact guidance for cells to align along the nanogratings direction [11,20,25,26,28-32]. Surprisingly, NHFLs did not align on all nanogratings studied. As shown in Fig. 1.3A, the cells did not display obvious alignment preference when the nanogratings had the same spacing as the linewidth. On the other hand, the cells preferentially oriented along the nanogratings direction when the spacing was 3X the linewidth for all three linewidths studied. We further examined the cell spreading using SEM (Fig. 1.3B). The cells seemed to stay on NG 300-1X, but could reach the bottom of the grooves for the other five nanogratings. When the spacing was 1X linewidth, the filopodia could reach out perpendicular to the grating direction (NG 500-1X and NG 1000-1X), while the filopodia of the cells on NG 300-3X and NG 500-3X preferred to extend along the grating direction. In our nanotopography design, the feature height was 170 nm. The cells could overcome the grating barrier and reach the neighboring gratings when the spacing was narrow. When the spacing was three times the linewidth, it might be easier for the cell to follow contact guidance and to orient along the gratings. When the spacing was further increased, i.e. 3000 nm for NG 1000-3X, the lamellipodia could reach the groove bottom and the filopodia could extend perpendicular to the grating direction. Both confocal and SEM observations suggested that although nanogratings might provide contact guidance the human fibroblasts could bridge the gaps and did not show preferential orientation when the spacing was narrow. When the spacing was significantly increased, however, it may be easier for the cells to elongate along the ridge. Therefore, the nanograting-induced cell orientation is spacing dependent.
Figure 1.3: Cell spreading on nanogratings. A) Confocal and B) SEM images of NHLFs grown on nanogratings. The yellow arrows point to the nanograting direction. Scale bars = 1 µm.
4.3 Nanogratings Modulated Focal Adhesions and Nuclear Size

As the first step for the adhesion-cytoskeleton-nucleus mechanotransduction pathway, focal adhesions of cells in response to nanotopographies were examined. We analyzed confocal images for focal adhesion protein paxillin in the cells on the nanogratings and the flat surface, and compared structures between 2 and 10 μm², the typical size for mature focal adhesions [50]. The focal adhesion structures in the high-magnification confocal images (Fig. 1.4A) were extracted using Imaris (Fig. 1.4B) and the focal adhesions were analyzed.

![Figure 1.4: Image analysis using Imaris. The confocal images of focal adhesion protein paxillin structures on (A) nanogratings were extracted using Imaris after applying threshold and shown in (B).](image)

Nanotopography can increase the total surface area, depending on the shape and size of the features. We defined the ratio of total surface area versus the corresponding flat surface to reflect the extent of nanotopography-induced increase in the surface area. Fig. 1.5 showed that when the ratio was low (all nanogratings except NG 300-1X) there was no obvious difference in the average focal adhesion size between the nanogratings and the flat control, 4.6 ± 0.1 μm². However, the average focal adhesion size of cells on NG 300-1X was significantly smaller than the flat control. The spacing of NG 300-1X was 300 nm while other nanogratings had spacing larger than 500 nm, indicating that there was a critical spacing of between 300 and 500 nm. This observation agreed well with the previous report that the cells could fill 800-nm gratings completely but not reach of bottom of 400-nm gratings [51]. The results suggested that
nanogratings may affect the focal adhesion size, depending on not only the surface area increase but also the spacing.

![Graph showing focal adhesion size comparison]

**Figure 1.5:** Nanogratings-modulated focal adhesion size. The straight horizontal line indicated the average focal adhesion size of NHLFs on the flat control.

Of particular interest was the nuclear deformation, which would result in changes of gene and protein expression [42,43] and was strongly associated with cell proliferation and differentiation [45,46]. To validate the accuracy of the confocal z-stack measurement of nuclear volume, we measured the FITC-spheres with a diameter of $15.4 \pm 0.13 \, \mu m^2$ which were embedded in collagen I gel. The measured sphere volume was $1893.8 \pm 71 \, \mu m^3$, which agreed well with the calculated volume of $1912.3 \pm 56 \, \mu m^3$ (measurement error < 1%). The nuclei confocal images were reconstructed in Imaris and nuclei area and volume were analyzed (Figure 1.6). Compared to the nuclear area and volume on the flat surface, the cells displayed larger nuclear area (Fig. 1.7A) and volume (Fig. 1.7B) on all nanogratings except on NG 300-3X. McKee et al. measured the nuclear deformation and nuclear elastic modulus of human corneal
epithelial cells grown on equally spaced nanogratings with a pitch from 400 nm to 2000 nm and a constant depth of 300 nm. They found that the nuclear volume increased as the pitch of nanogratings increased from 400 nm to 2000 nm. The 400-nm nanogratings resulted in the smallest volume and other nanogratings had higher nuclear volume than the flat surface. The elastic modulus of aligned cells was nanograting-dependent, but not simply increased with increasing pitch size [48]. This differences were believed to be resulted from several issues. McKee et al. did not eliminate the variation by synchronizing the cell cycle. They only measured six cells, which might not represent the average response from the whole cell population. Different cell types studied might be another factor. Nanogratings, very often, reduced the surface area cells can sense, which can result in a decrease cell spreading and proliferation although the cell alignment, elongation and migration are enhanced [11,24,29-33]. In the present study, NHLFs spread less on nanogratings but had increasing nuclear volume for all but one pattern.

**Figure 1.6:** Typical projected nuclear area and reconstructed nuclear volume of cells on nanogratings.
Figure 1.7: Nuclear deformation on nanotopographies. (A) Nuclear area and (B) nuclear volume comparison of cells grown on nanogratings. The straight horizontal lines indicated the average nuclear area and volume of NHLFs on the flat control.
4.4 Cellular Responses to Nanoposts

Different from nanogratings, NHFLs spread in all directions on nanopillars. Except for NP 300-1.3X where the PDMS pillars were densely packed and collapsed into bigger aggregates because of dimensional instability, the cells could confine on the pillars and the filopodia extension on the top of the pillars for both small (300 nm) and large (1000 nm) pillars (Fig. 1.8). Focal adhesion protein paxillin displayed random orientation on the isotropic pillars (Fig. 9). Image analysis of focal adhesions showed that the average focal adhesion size was smaller than that on the flat control (Fig. 1.10). In particular, the focal adhesion size on NP 500-1.9X was significantly smaller than the flat control.

Figure 1.8: SEM micrographs of human fibroblasts grown on (A) nanopillars 300-1.9X and (B) nanopillars 1000-1.9X. The boxed regions were enlarged to show the filopodia extension.
**Figure 1.9:** Image analysis using Imaris. The confocal images of focal adhesion protein paxillin structures on (A) nanogratings were extracted using Imaris after applying threshold and shown in (B).

**Figure 1.10:** Nanopillars-modulated average focal adhesion size.
Interestingly, the cells on NP 500-1.9X displayed a larger nuclear volume than that on the flat surface (Fig. 1.11). In NP 500-1.9X, the edge-to-edge distance of 450 nm was large enough so that the cells would completely confine to the space and thus the nanopillar-induced total surface area increase enhanced cell spreading. When the spacing was further increased to 900 nm of NP 1000-1.9X, the increase in surface area and cell spreading was not significant. Therefore, the nuclear volume increased in NP 500-1.9X. When the edge-to-edge distance was smaller than 300 nm (NP 1000-1.3X, NP 300-1.9X and NO 500-1.3X) the cells might not see obvious difference between the nanopillars and the flat surface and the nuclear volume was almost the same.

Correlation between focal adhesions and nuclear volume was also analyzed. There was no significant difference in the average focal adhesion size and nuclear volume of the cells on nanogratings (except NG 300-1X) and the flat surface. However, NP 500-1.9X enhanced cell spreading, and consequently increased the nuclear volume. Roca-Cusachs et al. suggested that cell spreading promoted DNA synthesis by inducing nuclear swelling, thereby induced an increase in proliferation, actin polymerization (F-actin content), and stress fiber formation, whereas elongation dramatically affected entanglement and bending of actin filaments but had little or no effect on proliferation, formation of stress fibers, and F-actin content [46].

It was noted that some nanotopography had smaller focal adhesions size but bigger nuclear volume. For example, the cells on NP 500-1.9X had significantly smaller average focal adhesion size than the flat control but the correspondent nuclear volume was significantly larger than that on the flat surface. We postulated that the quantity of total focal adhesions instead of the average focal adhesion size determined cell spreading and thus nuclear volume. Jean et al. measured nuclear deformation in response to alterations of cell shape as cells detach from a flat surface. They found that the nuclear deformation appeared to be in direct and immediate response to alterations of the cell adhesion area [52]. As a matter of fact, it was observed that the cells were easier to detach from nanogratings than the flat surface during the cell fixation process.
Figure 1.11: Nanopillar modulated deformation of (A) nuclear area and (B) nuclear volume of fibroblasts.

Nuclei Area ($\mu m^2$)

- **A**:
  - 1000-1.9X: 639 ± 15 μm²
  - 500-1.9X: 700 μm²
  - 1000-1.3X: 700 μm²
  - 300-1.9X: 700 μm²
  - 500-1.3X: 700 μm²

Nuclei Volume ($\mu m^3$)

- **B**:
  - 1000-1.9X: 1218 ± 35 μm³
  - 500-1.9X: 1240 μm³
  - 1000-1.3X: 1240 μm³
  - 300-1.9X: 1240 μm³
  - 500-1.3X: 1240 μm³

* Total Surface Area/Flat Surface
5. Conclusion

Focal adhesions and nuclear deformation of NHFLs in response to a variety of nanotopographies have been investigated. It was found that the orientation of the focal adhesions and nuclei along the nanogratings depended on not only the linewidth but also the spacing. Nanogratings, in general, hindered cell spreading and thus reduced the cell size while increasing the nuclear size. On the other hand, nanopillars, depending on the feature size and spacing, might modulate the focal adhesions and nuclear towards opposite directions. Our observations also suggested that it was focal adhesion area instead of the focal adhesion size that determined the cell spreading and nuclear deformation. Taken together, nanotopography could be optimized to modulate the cell adhesions and nuclear volume, which provide the useful tool to regulate cells for the end applications.
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II. Three-dimensional Microfluidic Co-Culture Model of the Bone Marrow Microenvironment for the Study of Acute Lymphoblastic Leukemia

1. Introduction

1.1 Physiological Environment

The bone marrow microenvironment is the primary site of normal hematopoietic cell development as well as the anatomical location in which diverse types of leukemia initiate and progress. It is notably the most frequent site of leukemic relapse, as well as a site of metastasis for many solid tumors including breast, lung, and prostate cancer. Held in common to all tumor cells that either originate or migrate to this site is the propensity to be refractory to treatment [1] thus positioning them to contribute to relapse of disease. As such, it is important to attempt to model this site appropriately to investigate tumor cell survival in this context and to develop drug screens that consider its complexity.

With that goal in mind the current model includes cell types that contribute to the heterogeneity found in this unique space with a focus on an aggressive form of acute lymphoblastic leukemia (ALL) that constitutively expresses the Bcr-Abl fusion protein as a representative model. Bone marrow stromal cells (BMSC) and osteoblasts comprise just two of the well characterized populations that are essential to the developmental support of both normal and leukemic cells in postnatal bone marrow [2,3]. These two populations provide secreted cytokines and chemokines as well as support of cell signaling that is prompted through physical interaction of hematopoietic cell surface receptors with adhesion molecules including, but not limited to Vascular Cell Adhesion Molecule-1 (VCAM-1) and fibronectin. The heterogeneity of the supportive microenvironment is significant in terms of cellular constituents but models must also consider extracellular matrix (ECM). Various collagens comprise a significant component of the ECM [4-7] with Collagen I being particularly abundant in the marrow space [8,9]. Of additional influence on hematopoietic cell development is the elasticity of the matrix, which has profound effects on tumorogenesis [10-13]. The interstitial fluid flow in bone, being extremely slow (between 0.1 and 4.0 μm/s [14]), plays an important role in nutrient transport and establishment of the microenvironment [15,16]. The interstitial flow has been reported to regulate tumor cell growth,
differentiation, migration and metastasis [17-21], and to promote angiogenesis and tumorigenic activity of stromal cells [22].

1.2 Current Tumor Models

Several models to investigate the influence of the bone marrow microenvironment have been developed that are informative, but continue to rely on combinations of relevant cell types grown in two dimensions (2-D). With recognition of the importance of architecture to the unique anatomy of the bone marrow effort is warranted to improve on these models to attempt to move closer to biological relevance. The 3-D models have been shown to restore cellular morphologies and phenotypes characteristic of in vivo tumor development [23-26]. Simply switching culture dimensionality from 2-D to 3-D drastically affects cell morphology [27], proliferation [28], differentiation [29], gene and protein expression [24,30-33], and metabolism [34]. Reflecting the impact of dimensionality, GB1 glioma cells were shown to elongate and flatten in 2-D culture, destroying the typical pseudo-spherical morphology and fillapodial characteristics, but closely resembles the original in vivo phenotype in 3-D culture [35]. Just as cancer cell gene expression patterns can differ, chemotherapy drugs display distinct sensitivities in 2-D versus 3-D [24,36]. Two dimensional glioblastoma models were more sensitive to the chemotherapy agent TMZ than 3-D models or the clinical population [35]. Additionally, treatment of MDA-231 3-D cultures, but not MDA-231 monolayers, with the DNA-damaging therapeutic cisplatinum resulted in up-regulation of TGFβ1 – a response that is predictive of the response in vivo [37]. Three dimensional models also allow investigation of cell-cell interactions [38]. For example, Gao et al. observed the distance dependent effects of soluble factors secreted by L929 fibroblast cells on tumor cell cycle progression in multilayered hydrogels containing living cells [39]. The 3-D matrix model has been further incorporated into microfluidic platforms, thus enabling systematic investigation of both physiological and pathological phenomena in vitro [22,40-47]. In carefully controlled experimental approaches, cancer cell migration [48,49], tumor-stromal cell interactions [50], and tumor-endothelial cell interactions [41,51] have been studied in 3-D microfluidics. For example, breast cancer cells (MDA-MB-231) preferentially migrated along the streamline in a 3-D microfluidic platform, but the direction depended on cell density, CCR7
receptor activity and interstitial flow velocity [49]. Additionally, extravasation and migration distance of breast cancer cells were significantly enhanced in a vascularized osteo-cell conditioned microenvironment with human BMSC and endothelial cells [52]. These findings collectively emphasize the value of developing 3-D microfluidic models to interrogate biological questions, including those related to the bone marrow microenvironment, as the focus of the current work.
2. Objective

The objective of this project was to investigate the influence of the bone marrow microenvironment on the chemoresistance of aggressive leukemic cells for Acute Lymphoblastic Leukemia (ALL). In our study, we co-cultured three biologically relevant populations of bone marrow niche cell types: bone marrow stromal cells, human osteoblasts, and an aggressive leukemic cell line. The study compared traditional 2-D static tumor alone and co-culture models to 3-D static and 3-D dynamic models. By creating this model, we set out to establish an *in vitro* model which addresses *in vivo* characteristics, such as, dimensionality, cell-cell interactions, and interstitial fluid flow. By incorporating these crucial features, we hoped to create a model which better mimics the *in vivo* scenario, therefore, contributing to cancer progression and treatment analysis.
3. Materials and Methods

3.1 Fabrication of a Microfluidic Platform

The microchannels of 500 \( \mu \)m width, 75 \( \mu \)m height and 2 cm length were generated via conventional photolithography [53]. The micropattern created with AutoCAD was printed on a transparency film that served as the photomask and SU-8 2050 (MicroChem Corp., Newton, MA, USA) was used as the substrate.

Polydimethylsiloxane (PDMS) resin and curing agent (Sylgard 184 kit, Dow Corning, MI, USA) was mixed at a 10:1.05 w/w ratio and poured over the SU-8 mold. After curing at 70\( ^\circ \)C for 2 hr, the inverse PDMS mold was peeled from the SU-8 mold.

The microfluidic platforms were assembled in two manners, conventional oxygen plasma assembly and novel microtransfer assembly as described previously [54]. Oxygen plasma provided a permanent assembly while microtransfer assembly was reversible. Briefly, the PDMS mold with microchannels and the glass coverslip were exposed to oxygen plasma at 300 mTorr (40 Pa), 50 watt for 15 s in a March PX-250 Plasma Asher from Nordson Co. (Westlake, OH, USA). The oxidized PDMS microchannels and glass coverslip were brought into contact and then placed in an oven at 125\( ^\circ \)C for 15 min to complete the assembly. During microtransfer assembly, 5 wt.% PDMS prepolymer in hexane solution was spin-coated onto a Si wafer to form a thin film, which served as an adhesive layer to assemble PDMS microchannels onto a glass coverslip. The coverslip was pretreated with oxygen plasma as described above. The assembly was post-cured at 80\( ^\circ \)C for 2 hr. The microtransfered assembly allowed to peel the PDMS mold off from the coverslip after cell studies and thus the cell-encapsulated hydrogel matrix could be retrieved for analysis.

Prior to cellular studies, the platform was sterilized by injecting and retaining 70% ethanol within the microchannels for 15 min followed by UV exposure for 15 min.

3.2 Cell Culture

Human bone marrow stromal cell (BMSC) cultures were initiated from human bone marrow from consenting donors, with approval by the West Virginia University Institutional Review Board. All
primary BMSC cultures were initiated from donors with no previous chemotherapy exposure. BMSC were maintained in α-modification of Eagle's medium (α-MEM; Gibco, Grand Island, NY, USA) supplemented with 10% fetal bovine serum (Hyclone, Logan, UT, USA), 1% L-glutamine (Gibco), 1% penicillin/streptomycin (Sigma, St. Louis, MO, USA), and 0.1% 2-β− mercaptoethanol (Sigma). Human osteoblasts (HOB) were purchased from Promocell (Heidelberg, Germany) and maintained in osteoblast growth media. The human Philadelphia chromosome positive (Ph+) B lineage ALL cell line SUP-B15 (CRL-1929) was obtained from ATCC and cultured in Iscove's DMEM (Mediatech, Manassas, VA, USA) supplemented with 10% fetal bovine serum, 2 mM L-glutamine, 0.05 μM 2-β−mercaptoethanol and 1% penicillin/streptomycin.

3.3 Tumor Models

Two groups of culture conditions were compared that included SUP-B15 leukemic cells alone or SUP-B15 cells co-cultured with HOB or BMSC. The total cell density was maintained at 20x10⁶ cells/mL. In co-culture experiments, SUP-B15 cells were mixed at a 10:2:2 ratio with HOB and BMSC. The cells were resuspended in 100 μL complete culture medium, gently mixed with 10% 10X PBS, 2% NaOH, and 88% Collagen I (rat-tail, BD Biosciences, Franklin Lakes, NJ, USA), and then injected into the microfluidic platform previously described. The platform was incubated at 37°C for 1 hr to allow collagen I to gelate before the culture media was continuously pumped through the platform at a flow rate of 800 μL per day using a Harvard Apparatus PhD Ultra Pump (Harvard Apparatus, Holliston, MA, USA) to sustain cell culture for a predefined period of time. For comparison, both tumor cells alone (SUP-B15 only) and co-culture with BMSC and HOB were studied in 2-D and 3-D configurations under the static condition. In the 2-D static model, cells were seeded in 6-well tissue culture polystyrene (TCPS) plates. In the 3-D static model, the mixture of cells (SUP-B15 + BMSC + HOB) with collagen I was plated in 24-well culture plates and allowed to gelate.

To demonstrate cell-cell interactions in 3-D matrices, SUP-B15 leukemic cells, BMSC and HOB were fluorescently labeled by using Cell Tracker Green CMFDA, Red CMTPX, and far red shown in blue, respectively. The Cell Trackers were obtained from Life Technologies, Grand Island, NY, USA.
3.4 Chemotherapeutic Reagents and Treatment

Antimetabolite chemotherapeutic drug, Ara-C (Sigma) was used to evaluate therapeutic effects on tumor cells. Ara-C was stored at -80 °C at 10 mg/mL, and diluted in the base media used for leukemic cell culture immediately before use. After 72 h incubation, tumor cells cultured in 2-D and 3-D under static conditions were treated with 1 μM Ara-C for 48 hr to best mimic an ALL patient’s drug serum concentration during treatment, while 3-D dynamic platforms were exposed to continuous flow of 1 μM Ara-C in complete culture media for 48 hr. Viability was evaluated by both flow cytometry and immunofluorescence image analysis.

3.5 Flow Cytometry

Collection of tumor cells from 3-D matrices was accomplished by incubation of cultures in type I collagenase (Life Technologies). The collagenase was used to digest collagen I in the 3-D static model according to the manufacture protocol. Apoptotic cells were detected by staining with Annexin V-Alexa 555 assay (Life Technologies) and gated against cells stained with CellTracker™ Green CMFDA. Data were acquired using a FACSFortessa flow cytometer with Cell Quest Pro software (BD Biosciences, Franklin Lakes, NJ, USA). Experiments were repeated in triplicate.

3.6 Immunofluorescence Staining

Annexin V Staining. Apoptotic cells were detected by staining with Annexin V-Alexa 555 (1:50) assay. Three-dimensional static and dynamic cell-matrix samples were washed with PBS and then stained with Annexin V overnight at 4°C. Samples were washed, fixed with 1% paraformaldehyde (PFA), and imaged on the same day. After immunofluorescence staining, cells were imaged using a Zeiss LSM 510 confocal microscope.

3.7 Confocal Image Analysis Using Imaris

Immunofluorescence images were analyzed using Imaris software (Bitplane, Windsor, CT, USA). The surface function was used to reconstruct objects in the FITC (green) and TRITC (red) channels. First, the threshold was adjusted to a level where the superimposed confocal image matched that of the created objects. The objects created were then filtered based on size to remove unwanted noise and to evaluate
only tumor cell viability. Remaining objects were counted based on color. Red Annexin V-positive cells were counted as not viable, while green Annexin V-negative cells were counted as viable.

3.8 Statistical Analysis

Viability percentages were expressed as mean ± standard error from three independent experiments. Immunostained images of no less than 10,000 tumor cells/sample for the 2-D model and 200 tumor cells/sample for the 3-D models were evaluated. The differences between groups were analyzed by one-way ANOVA and Tukey’s method for multiple comparisons. p < 0.10 was taken as statistically significant for tumor alone and co-culture comparison and p<0.05 was taken as statistically significant for 2-D and 3-D comparison.
4. Results and Discussion

4.1 Essential Elements in Tumor Models

The current study sought to develop a 3-D model with utility for investigation of tumor cell biology in the context of the bone marrow microenvironment as the site of initiation of leukemic disease and as the frequent site of metastatic malignancies characterized by therapeutic resistance. Inherent to this design were considerations of interstitial flow, dimensionality and cell-cell interactions (Figure 2.1) as critical parameters that impact on cell morphology and signaling that impacts on critical downstream pathways. BMSC and HOB were included as representative, but not exclusive, elements of the bone marrow microenvironment that have been well documented to influence both normal hematopoietic cell development as well as impacting on tumor cell quiescence, proliferation, and survival in this unique anatomical niche [8,9]. Collagen I was chosen for inclusion based on its predominance in the bone marrow, however, it was recognized that the extracellular matrix required to generate a truly biologically representative model would be more complex and include several types of collagens, hyaluronate, and fibronectin as well as other structural components [8,55,56].

![Figure 2.1: Schematic illustration of tumor models. The three factors tested in this study include dimensionality, cell-cell interactions and interstitial flow.](image)
4.2 Microfluidic Design and Fabrication

A typical 3-D microfluidic platform was shown in Fig. 2.2. Tumor cells alone or co-culture with BMSC and HOB mixed with Collagen I were injected through the “Cells In” inlet into four microchannels, which were 500 µm in width, 75 µm in height and 2 cm in length. After Collagen I gelated at 37°C incubator for 1 hr, the culture medium was flowed through a “Media In” inlet into the four microchannels for a predefined period of time (Fig. 2.2B). Near the inlet and outlet, perfusion channels comprising of four rows of 100 µm X 100 µm columns with a spacing of 80 µm were fabricated to help retain the cells within the main microchannels (Fig. 2.2C). The velocity profile within the microchannels deviated from Poiseuille flow because the compliant collagen hydrogel, the various pore size and cell-induced matrix remodeling. important [22]. By tracking the movement of injected TRITC-labelled beads, the velocity was determined to be 2.3 µm/s, which was within the physiological range.

Figure 2.2: A typical 3-D microfluidic co-culture model. (A) The microfluidic platform consists of 4 microchannels, each 500 µm in width, 75 µm in height and 2 cm in length. The divergent angles are 30°. Near the inlets and outlet there are perfusion channels. The enlarged boxed area illustrates multiple cell types embedded in a 3-D matrix. (B) Optical image of the microfluidic platform. (C) SEM images of top view (upper panel) and side view (lower panel) of the perfusion channels.
4.3 Co-Culture Models

Confocal and Imaris generated reconstruction images (Figure 2.3) show that in both 2-D and 3-D static as well as in 3-D dynamic experimental circumstances all cell populations can be readily discerned with 3-D reconstruction demonstrating direct cell-cell interactions with HOB and BMSC. Cross sections (CS) (Figure 2.3A) demonstrate the compaction of the 2-D static model compared to the 3-D circumstance in which the dimensionality can be appreciated. The cell-cell interactions are further emphasized in Fig. 2.3B upon rotation of the 3-D reconstruction images around the X and Y axis indicating tumor cell interaction with structural cells (BMSC and HOB).
Figure 2.3: Representative tumor models. (A) Confocal and Imaris reconstruction images of the tumor models at low magnification (20X), high magnification (40X) and high magnification cross sections (CS). The cell-cell interactions in z-direction in the boxed region of CS images are enlarged in Row 4. The heights for 2-D and 3D CS images are 10 µm and 30 µm, respectively. (B) 3-D reconstruction images rotated via x- and y-axis revealing cell-cell interactions.
4.4 Drug Sensitivity across Platforms

It has been reported that both stromal cells and osteoblasts enhance leukemic cell survival during chemotherapy exposure with previous experiments typically completed in traditional 2-D cultures [57,58]. The relevance of this observation is that surviving tumor cells in the context of leukemia directly correlate with the likelihood of relapse of disease [59] and as such more biologically relevant models to interrogate the protective effect of critical elements of the microenvironment are warranted. In the current study, we determined that this trend of microenvironment protection of tumor cells grown with access to BMSC or HOB was also observed in both 3-D static and 3-D dynamic (microfluidic) models. Flow cytometry and confocal microscopy were used to evaluate viability of tumor cells in the 2-D and 3-D models (Fig. 2.4). Regardless of whether the 3-D configuration was static or had a steady media flow during culture, it was associated with significantly higher survival of tumor cells during Ara-C exposure as well as enhanced protection during chemotherapy stress conferred by microenvironment cells (Fig. 2.5, 2.6).
Figure 2.4: (A) Typical flow cytometry analysis for untreated 2-D tumor model (left) and Ara-C treated 3-D co-culture model (right) using collagenase to digest collagen matrix. (B) Typical immunofluorescence image analysis of untreated (left) and treated (right) 3-D dynamic tumor models.
Figure 2.5. Comparison in chemoresistance of tumor cells to Ara-C among a variety of tumor models. (A) The effects of dimensionality, co-culture and interstitial flow on the viability of tumor cells. Bracket shows significant (p<0.1) difference between treated tumor and treated co-culture.

Figure 2.6: The effects of 3-D culture on tumor cell viability under the co-culture condition. Brackets reveal significant (p<0.05) difference in 3-D and 2-D models. All 3-D models showed significant difference when compared to 2-D models.
The observed chemoresistance difference in 3-D culture can be attributed to the enhanced cell-matrix interactions. Over the past decade, a growing number of reports have shown that there are different signaling mechanisms in 3-D microenvironments compared to 2-D substrates [60,61]. Moreover, tumor cells cultured in three dimensions have been shown to be less sensitive to apoptosis induced by radio-chemo treatments or by death receptor ligation compared to cells cultured in two dimensions [62]. Cytotoxic effects of chemotherapeutic agents have also been shown to be significantly reduced in cells cultured in three dimensions [24,63,64]. These data suggest that investigation of leukemic cell protection by bone marrow microenvironment cues in traditional 2-D culture models can provide meaningful information in which trends are confirmed in 3-D culture, but that more accurate measures of the magnitude of effect may require further development of 3-D models. No readouts were observed that were not consistent between 2-D and 3-D based experiments, and as such, practical downstream evaluation of cells may also drive the experimental design choice. Potentially experiments in which large numbers of tumor cells are required for recovery at the cessation of cytotoxic exposure may be best aligned with 2-D approaches, with 3-D lending itself best to single cell analyses or polymerase chain reaction (PCR) based applications. We have documented that leukemic cells can be recovered via collagenase digestion and cell sorting with high quality RNA that is appropriate for RT-PCR being easily recovered. This will broaden the realm of experiments that will be possible related to gene expression studies in this microenvironment model.
5. Conclusion

Relapse of leukemic disease in high risk categories of ALL remains a significant clinical challenge, with pre-clinical models providing platforms for investigation of pathways that influence tumor cell quiescence, the regulation of tumor stem cell phenotype, and modulation of apoptotic pathways as just a few of the elements that converge on response to therapy. In combination with appropriate in vivo models, it remains essential to optimize in vitro models to more accurately mimic the complexity of the bone marrow microenvironment to provide new tools for investigation of the mechanisms that underpin survival of residual disease and relapse. The microfluidics model in the current study represents one step in this effort with relevance to both leukemic disease as well as metastasis of solid tumors to the bone marrow where they also benefit from the protective niche during therapy.

The current study showed decreased chemotherapeutic drug sensitivity of leukemic cells when co-cultured with BMSC and HOB and exposed to flow. The engineered 3-D microfluidic co-culture model allowed precise control over the mechanical properties of matrix and fluidic shear stress and enabled systematic investigation of effects of cell-cell and cell-matrix interactions on cancer initiation and progression, thus improving our limited comprehension of the role of microenvironmental signals in cancer biology.
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