Nanostructured magnetizable materials that switch cells between life and death

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Abstract

Development of biochips containing living cells for biodetection, drug screening and tissue engineering applications is limited by a lack of reconfigurable material interfaces and actuators. Here we describe a new class of nanostructured magnetizable materials created with a femtosecond laser surface etching technique that function as multiplexed magnetic field gradient concentrators. When combined with magnetic microbeads coated with cell adhesion ligands, these materials form microarrays of ‘virtual’ adhesive islands that can support cell attachment, resist cell traction forces and maintain cell viability. A cell death (apoptosis) response can then be actuated on command by removing the applied magnetic field, thereby causing cell retraction, rounding and detachment. This simple technology may be used to create reconfigurable interfaces that allow users to selectively discard contaminated or exhausted cellular sensor elements, and to replace them with new living cellular components for continued operation in future biomedical microdevices and biodetectors.

Keywords: Magnetic particles; Magnetic gradient concentrator; Culture substrate; Apoptosis; Mechanical force; Cell shape

1. Introduction

The development of microchips containing biological molecules, such as multiplexed arrays of specific DNA sequences, has revolutionized genomics, diagnostics and pathogen detection. In the future, it is likely that living cells will be integrated as system components within biochip-based microdevices used for drug development, medical microdevices and biodetection because of their unique abilities to simultaneously sense and respond to multiple physiological or pathological stimuli. However, biocompatible cell–material interfaces will need to be created to make these technologies a reality.

Microfabrication techniques, such as soft lithography and microcontact printing, have been used to create arrays of adhesive islands that may be used to place cells in predefined positions and maintain them in a functional state [1–3]. These substrates also have been integrated with microfluidic systems that permit cells to be plated on these substrates and nutrient media to be delivered continuously to maintain cell function and to introduce various chemical stimuli [4–6]. But these systems are limited in that they are not reconfigurable. For example, it would be extremely desirable to have future biodetectors containing cells that can be rapidly removed and replaced with new cellular components after exposure to contagious biopathogens in order to reboot the device without requiring addition of expensive enzymes or harsh chemicals that might interfere with the functionality of the system. This type of reconfigurable functionality also might be useful for future drug screening, tissue engineering and biocomputational applications in which living cells are used as primary signal processing elements [7–9].

Several systems have been described that allow users to control the cell adhesive properties of substrates, including thermal responsive polymer surfaces [10], photo-initiated gels [11,12] and electrochemical modulation of
self-assembled monolayers [13,14]. Although these technologies may provide useful methods to rapidly actuate changes in substrate adhesivity and cell behavior, they are not easily reversed within an existing microsystem. Here we describe a microsystem in which cell adhesion, position and viability can be controlled magnetically to meet this challenge.

2. Materials and methods

2.1. Nanofabrication of magnetic gradient concentrators

Arrays of silicon nanospikes were fabricated by femtosecond laser etching, as previously reported [15]. Briefly, silicon chips [flat zone Si(1 1 1)] were masked with 2 μm-thick copper transmission electron microscope grids with hexagonal opening widths of 30 μm. The chips were mounted normal to the laser beam on a three-axis translation stage in a vacuum chamber with a base pressure of less than 10⁻⁶ Pa. Laser pulses (800 nm, 100 fs at 0.3 mJ) were focused to a beam waist of 200 μm on the sample surface. Samples were translated relative to the laser beam vertically at a speed of 200 μm/s to irradiate the entire surface. Laser-etched silicon chips were subsequently coated with a thin (450 Å) layer of magnetizable cobalt using a SHARON TE-4 four-source thermal evaporation system. After irradiation and coating, the resulting structures were imaged with a scanning electron microscope.

Arrays of stainless steel nanospikes were fabricated by femtosecond laser etching of surgical stainless steel blades (#21, Becton Dickinson, Franklin Lakes, NJ) that were mechanically precut into 1 cm² wafers. To create patterned arrays, the surface was polished with a diamond lapping film, and then mounted normal to the laser beam on a three-axis translation stage in a vacuum chamber with a base pressure of less than 10⁻⁶ Pa. Laser pulses (800 nm, 100 fs at 0.15 mJ) were focused on the sample surface, and the sample was translated relative to the laser beam vertically at a speed of 500 μm/s to etch parallel linear valleys with separation of approximately 100 μm (center to center) on the surface of the stainless steel. The sample was translated 90° and linear etching was repeated to generate cross-hatched valleys, leaving an array of square raised areas with sides approximately 50 μm in length.

To create nanospikes on the raised square arrays, wafers were again lightly polished with a diamond lapping film, and then cleaned with acetone followed by a methanol rinse. Wafers were subsequently placed in a glass container, mounted on a three-axis translation stage and filled with distilled water. The surface of the array was irradiated by a 1 kHz train of 100 fs, 800 nm pulses at 400 nm wavelength from a frequency-doubled, amplified titanium:sapphire laser. The laser pulses were focused by a 0.25-m focal-length lens and traveled through 10 mm of water before striking the surface at normal incidence. The focal point is about 10 mm behind the surface and the spatial profile of the laser spot is nearly Gaussian, with a fixed beam waist of 60 μm at the sample surface. The sample was translated relative to the laser beam at a speed of 200 μm/s to create nanospikes over the entire micro-arrayed surface. The dimensions of the resulting structures were determined from scanning electron microscope images as well as analysis with a Dektak 6M profilometer.

Etched stainless steel wafers were coated with Sylgard 184 polydimethylsiloxane (PDMS; Dow Corning, Midland, MI) using a GSP-8 Spincoat device (Specialty Coating Systems/Cookson Electronics, Indianapolis, IN) to prevent surface oxidation and optimize biocompatibility. PDMS base and curing agents were mixed in a 10:1 ratio, degassed by vacuum for 30 min, spin-cast, and cured at least 3 h at 60 °C. Cured PDMS surfaces were made hydrophilic by 8 min exposure to an oxidizing plasma in an UVO-Cleaner 342 (Jelight, Irvine, CA), and wafers were immediately immersed in aqueous Pluronic F-127 solution (1 g/L; BASF, Mount Olive, NJ) for 1 h at room temperature to resist cell adhesion [16]. Wafers were rinsed extensively in phosphate buffered saline prior to incubation with beads and/or cells. The thickness of the PDMS was estimated to be 5 μm based on confocal imaging of coated wafers in which 200 nm green fluorescent beads were added to the PDMS.

2.2. Experimental system

Immortalized human umbilical vein endothelial (HUVE) cells that stably expressed green fluorescent protein (GFP) [17] were cultured in EBM medium (Cambrex) supplemented with EGM-2 MV growth factor supplement (Cambrex) at 37 °C and 5% CO₂. Subconfluent cells were detached by trypsinization, washed once in medium and cultured in EBM containing EGM-2 MV on magnetic substrates. Totsyl-activated superparamagnetic microbeads (4.5 μm diameter; Dynal Biotech, Oslo, Norway) were coated with an RGD-containing peptide (50 μg/ml; PepTite-2000; Integra LifeSciences, Plainsboro, NJ), that mediates cell adhesion through cell surface integrin receptors, as previously described [18,19]. Nanoparticle magnetic concentrators were placed horizontally in a cell culture apparatus that allowed permanent ceramic magnet cylinders (27 MG Oe; Polysciences Inc., Warrington, PA) to be placed directly beneath the concentrators, separated only by thin cover glass (No. 0; Dow Corning). For bead patterning, 100,000 RGD-coated beads were added to each magnetic field concentrator in 10 mm diameter cloning cylinders (Belleco Biotechnology, Vineland, NJ) to avoid dilution of beads in the larger medium reservoir. Beads were readily visualized by fluorescence microscopy due to their auto-fluorescence in the Texas-Red channel. Cells were added to magnetic substrates (2500 cells/substrate) in 10 mm diameter cloning cylinders, and placed in a culture incubator at 37 °C under 5% CO₂ for the indicated times. In some experiments, cells were allowed to bind to the RGD-beads (20 beads/cell) in suspension for 30 min at 37 °C prior to adding them to the magnetic field concentrators. Changes in projected cell areas were measured using computerized image analysis as previously described [1,18]; this was facilitated by visualizing the GFP-labeled HUVE cells using fluorescence microscopy.

The magnetic field gradient concentrators were calibrated by pulling 4.5 μm diameter beads through a glycerol solution of known viscosity (1000 Poise), as previously described [20]. Briefly, bead displacements and velocities were quantitated using IPlab image analysis software and time-lapse brightfield images (<0.1 s intervals) captured with a Nikon Diaphot 300 inverted microscope and a Photometrics CoolSnap HQ CCD digital camera. After recording the beads’ velocities through the fluid, Stokes’ formula for low Reynolds number fluid was used to deduce the forces [Force = 3ρDv, where ρ is the viscosity of the fluid, D is the bead diameter, and v is the velocity of the bead through the fluid]. For force calibration experiments, magnetic field gradient concentrators were placed on edge in glass-bottomed culture dishes, such that beads could be tracked in the X–Y plane of the microscope field as they approached the surface of the magnetic field gradient concentrator. Forces on individual beads were plotted against the distance between the bead and the surface of the magnetic field gradient concentrator.

To measure apoptosis we used a TUNEL in situ cell death detection kit (Roche, Penzberg, Germany) that measures DNA fragmentation as an indicator of programmed cell death. To harvest cells for this assay, magnets were removed from concentrators to release cells and cells were transferred to poly-1-lysine (Sigma) coated 4-chamber slides (Lab-Tek, Naperville, IL). Cells were allowed to adhere to the slides for 30 min and then fixed for 1 h in 4% paraformaldehyde in PBS. Apoptosis-associated DNA strand breakage was identified by following the TUNEL assay kit protocol for adherent cells, and cells were counterstained with DAPI to visualize all nuclei. Positive staining was detected by fluorescence microscopy using a Nikon Diaphot 300 inverted microscope and a Photometrics CoolSnap HQ CCD digital camera, and images were captured and analyzed with IPlab software. At least 100 cells were counted for each condition and results are presented as a percentage of DAPI-stained nuclei that exhibited positive TUNEL staining.

3. Results

The goal of this study was to microfabricate a cell culture environment containing a reconfigurable material interface that can rapidly actuate changes in substrate adhesivity...
and cell behavior. Our design concept was based on past work which showed that anchoragedependent cells, such as endothelial cells, require adhesion to substrates that can resist cell traction forces and produce a minimal degree of cell distortion to maintain cell viability and suppress the cellular suicide program (apoptosis) [1,21]. To distort their shape and avoid entry into the apoptosis pathway, anchoragedependent cells must exert substantial force (>1 nN/μm²) against their individual focal adhesions to the underlying extracellular matrix (ECM) [22–24]. This range of force can be applied to cells using magnetic microparticles coated with ECM ligands [25,26], and forces applied over integrins in this manner have been shown to alter intracellular biochemical activities including translation, signal transduction and gene expression [20,27–29]. This observation raised the possibility of creating a ‘virtual’ adhesive substrate by exerting forces on cell surface integrin-bound magnetic beads using applied magnetic fields that are of sufficient strength to resist cell tractional forces, and thereby maintain cell adhesion and viability; removal of the applied magnetic field would then induce cell retraction, rounding and death (Fig. 1). Importantly, this system could then be reconfigured by adding new beads and cells, thus providing a significant advantage over past materials that, for example, promote substrate release and cell detachment by altering cell temperature [10]. However, to produce pulling forces magnetically, it is necessary to create very high local magnetic field gradients. To do this in a manner that would be useful for future cell-based microdevices where single cell analysis may be required, we needed to generate these in a multiplexed manner with each adhesive magnetic island being on the scale of a single cell.

Regions of magnetic materials with higher curvature exhibit enhanced magnetic field gradients [30]. We have previously shown that irradiation of the surface of a silicon chip with a high power femtosecond laser (800 nm, 100 fs, 0.3 mJ) in the presence of certain gases, such as SF₆, transforms its flat mirror-like surface into a forest of microscopic spikes with very high curvature in the nanometer to low micrometer range [15,31]. Thus, we explored whether these novel laser-etched silicon chips could enhance local magnetic field gradients if coated with a layer of magnetizable material, and if so, whether microarrays of cell-sized islands could be created with this technique.

In our initial studies, arrays of multiple, micrometer-sized islands of high density nanospikes were etched into silicon substrates by masking the chips with a copper transmission electron microscope grid containing 30 μm wide hexagonal openings during laser ablation (Fig. 2a). The substrates were then coated with a thin (450 Å) layer of Cobalt by thermal evaporation to render these substrates magnetizable. When a permanent ceramic magnet (27 MG Oe, Polysciences Inc.) was placed beneath the etched silicon wafer and fluorescent superparamagnetic microbeads (4.5 μm diameter; Dynal) were added to its surface in tissue culture medium, the beads preferentially

Fig. 1. Schematics showing substrates that support cell adhesion and control cell viability. (Left) Anchorage-dependent cells, such as HUVE cells, normally must attach and spread on a rigid ECM-coated culture substrate in order to survive and grow; cells that pull themselves off of a poorly adhesive substrate and round undergo apoptosis. (Right) Schematic of the reconfigurable magnetic control system for switching cells between cell life and death that is described here. Cells are allowed to adhere to ECM-coated microbeads that bind their cell surface adhesion (integrin) receptors. In the presence of a high local magnetic field gradient, magnetic forces exerted on the beads will pull them down onto the surface of a non-adhesive substrate and create a ‘virtual’ cell adhesive island. If the force is great enough to resist cell traction forces, the cells will spread and remain viable. When the magnetic field is removed, the cells will retract the beads, round and switch on a cellular suicide program.

Fig. 2. A multiplexed magnetic field gradient concentrator fabricated by femtosecond laser etching of a silicon wafer. (a) Low power scanning electron micrograph of the cobalt-coated, laser-etched surface of a silicon wafer. Arrays of microspikes contained within hexagonal regions were created using a copper transmission electron microscope grid as a mask during the laser etching process. (b) Fluorescence micrograph showing that fluorescent superparamagnetic microbeads (4.5 μm diameter) are pulled to the hexagonal nanospike regions of the microarray in the presence of an applied magnetic field. (c) High power micrograph of 4.5 μm beads (white arrow) that localize preferentially to the apex of the nanospikes (black arrow). (d) Fluorescence image showing that the magnetic microbeads exhibit a random distribution on the unpatterned regions of these cobalt-coated silicon wafers in the presence of a magnetic field (bar, 50 μm).
attracted to the hexagonal islands containing the nanospikes (Fig. 2b). Higher resolution light microscopic analysis of etched substrates placed on edge and viewed from the side confirmed that the beads were specifically attracted to the apex of the cobalt-coated nanospikes that exhibited the highest curvature (Fig. 2c). In contrast, no patterning of magnetic beads was observed when they were placed on a smooth cobalt-coated silicon chip in the presence (Fig. 2d) or absence (not shown) of an external magnet.

To increase the durability and magnetic conductance of these magnetic concentrators, as well as to eliminate the necessity of coating with cobalt, we explored whether magnetizable stainless steel substrates could be used in place of silicon wafers. Preliminary characterization of laser-irradiated stainless steel (using a surgical scalpel blade) revealed that nanospikes formed slightly below the plane of the steel surface (not shown), whereas on silicon, laser irradiation results in nanospikes that project above the plane of the surface. To present the nanospiked regions above the plane of the steel surface, we etched 2 μm deep valleys in a square (50 × 50 μm) grid pattern by using a computer to precisely control translational movements of planar steel substrates in the path of 800 nm, 100 fs laser pulses in a vacuum (Figs. 3a and b). The remaining raised square surfaces of these etched steel substrates were then irradiated in water with 400 nm, 100 fs laser pulses to produce nanospikes in these elevated regions (Fig. 3c). This resulted in creation of a magnetizable stainless steel surface containing 50 × 50 μm elevated square mesas separated by 50 μm wide valleys at a depth of 2 μm (Fig. 3d). Analysis of high power images of regions within the elevated islands revealed that the steel nanospikes were on average 282 ± 5 nm (S.E.M.) wide and 715 ± 23 nm high, with a center-to-center spacing of 406 ± 78 nm (Fig. 3e).

The steel nanostructured island microarrays were spin-coated with a thin layer (~5 μm) of PDMS to create a flat surface and prevent metal oxidation in the aqueous medium, thereby further increasing biocompatibility. The PDMS surface was subsequently treated with Pluronic F-127 to make the surface non-adhesive to cells, cell-derived ECM proteins, or the magnetic beads. These substrates were placed in modified tissue culture dishes that allowed for permanent ceramic magnets to be placed directly beneath the magnetic concentrators, separated by only a thin (~#0; 80–130 μm thick) glass coverslip that isolates the magnet from culture medium. When magnetic microbeads (4.5 μm diameter; 1500 beads/mm²; Dynal) coated with the RGD cell binding peptide of fibronectin were added in medium in the absence of an applied magnet, they settled down onto the PDMS surface by gravity over a period of minutes but they did not distribute in any discernable pattern (Fig. 4a). In contrast, when the same beads were suspended over the nanostructured stainless steel device in the presence of a single ceramic magnet, virtually all of the beads were preferentially concentrated at sites directly over the square, regularly spaced, nanospiked islands of the microarray (Fig. 4b). Importantly, the beads distributed as a single layer across the entire region of the raised islands containing the nanospikes. With the magnet in place, beads remained in patterned arrays even after gentle medium changes. In contrast, after removing the underlying magnet, the beads were readily dislodged and easily removed by gentle swirling of the medium (not shown).

To quantify the forces exerted on individual beads (and hence, cells) by the stainless steel magnetic field gradient concentrator, we analyzed its ability to pull the magnetic beads through a high viscosity (1000 cP) glycerol solution. Stokes’ law was used to calculate the applied force based on measured changes in bead displacement induced by the applied magnetic field, as previously described [20]. For these experiments, concentrators were placed vertically in glass-bottomed culture dishes so that the beads could be tracked in the X–Y plane of the microscope field as they approached the surface of the stainless steel magnetic field gradient concentrator. These studies confirmed that
individual magnetic microbeads were redirected specifically to the square elevated regions containing the nanospikes once they approached within approximately 20 μm of the nanoetched steel surface (Fig. 5a). Analysis of bead velocities resulted in reproducible force–distance relationships, and revealed that approximately 0.25–2 nN of force was applied to each magnetic bead within about 20 μm from the surface of the magnetic concentrator (Fig. 5b). As these force calibrations were performed with etched nanospike surfaces that were not coated with PDMS, the force generated at approximately 5 μm from the surface of the steel substrate corresponds to the force that would be experienced by the lowest layer of applied magnetic microbeads in the presence of PDMS. This force ranged from approximately 0.9 to 1.3 nN per bead for one magnet or two stacked magnets, respectively (Fig. 5b), which is in the range of the level of traction force living cells apply to individual focal adhesions [22–24].

To explore the utility of the stainless steel magnetic field concentrator as a magnetic cell culture substrate, RGD-coated microbeads were first added to multiple nanostructured square islands with a single ceramic magnet placed beneath the substrate. HUVE cells were plated on these magnetically stabilized microbead arrays at low density (40 cells/mm²) to minimize cell–cell adhesion, and to limit more than one cell from adhering to each island of nanospikes. Cells adhered readily to the magnetically trapped beads that formed the ‘virtual’ adhesive islands, and some exhibited partial spreading (Fig. 6a). These results confirm that magnetic forces applied to the RGD-coated micromagnetic beads are sufficient to resist local cell traction forces (Fig. 6a). Cells that did not come into contact with beads remained non-adherent and were easily removed from the PDMS surface by gentle washing (not shown). In a separate experiment, cells were allowed to bind RGD-coated beads in suspension before the entire mixture was plated on the magnetic field concentrator substrate with a ceramic magnet below. These bead-bound cells also were efficiently pulled to the square nanospiked regions of the microarray (Fig. 6b). In both cases, the bead-bound cells maintained their position over the square islands of the microarray containing the nanospikes for the entire 2-day culture period, as long as the ceramic magnet remained in place during the study.

Computerized morphometric analysis of the projected areas of HUVE cells pre-bound to RGD-beads in suspension and cultured for 24h in the presence of magnetic field gradient concentrators with 0, 1 or 2 stacked ceramic magnets placed beneath the substrate confirmed that cell spreading increased progressively (from 386 to 526 μm²) as the number of underlying ceramic magnets was raised from 0 to 2 (Fig. 7a). In contrast, all cells retracted and detached themselves from the surface of the PDMS-coated substrate when the magnets were removed.
These findings confirm that in the presence of applied magnetic field gradients, these magnetically trapped magnetic beads do, in fact, form ‘virtual’ adhesive substrates that can resist cell traction forces in the absence of covalent linkage to the underlying rigid substrate.

HUVE cells rapidly switch on the cellular suicide program and undergo apoptosis if they are cultured in suspension or on adhesive substrates that cannot effectively resist cell tractional forces and promote cell distortion [1,21]. Therefore, we explored whether removal of the magnetic field gradient and associated cell rounding can be used to switch on the cellular suicide (apoptosis) program in this system. Importantly, analysis of apoptosis using the TUNEL assay to measure DNA fragmentation under the same conditions used to quantitate cell spreading (Fig. 7a) revealed a progressive increase in the percentage of cells undergoing cell death (from 8% to 41%; Fig. 7b) as the number of magnets was lowered from 3 to 0. These results demonstrate that the magnetic field gradient concentrators can effectively resist cell traction forces so that cell distortion can be promoted, and cell viability can be maintained for at least 2 days in culture. They also show that cell death can be actuated simply by removing the applied magnetic field.

4. Discussion

Taken together, these results demonstrate that multiplexed magnetic field gradient concentrators can be created using femtosecond laser etching technology, and that microarrays of ‘virtual’ cell adhesive islands can be created when these materials are combined with microbeads coated with cell adhesion ligands. In the presence of a magnetic field applied with an inexpensive stationary magnet, living cells can be cultured on these microarrays and be maintained in a viable state for days if provided with appropriate nutrients. If desired, the cells adherent to every island, or a subset of islands, can then be abruptly switched into a cell death program, and be physically washed away from the non-adhesive PDMS surface of the magnetic field concentrator without requirement of any enzymes (e.g., trypsin) or harsh chemicals. New beads and living cellular components may then be added through a microfluidic system to ‘reboot’ the system. All of the beads were coated with the same ligand and incubated with the same cell type in the present study. However, by covering these microarrays of virtual adhesive islands with an overlay that contains multiple microfabricated holes in corresponding positions that can be filled individually with different medium or cellular components using automated micro-pipetting techniques, this system can be provided with a combinatorial capability. These more sophisticated multiplexed systems would permit simultaneous analysis of the effects of multiple substrate ligands or soluble stimuli on multiple cell types.

The success of this reconfigurable technology required the development of magnetic field concentrators that enable these surfaces to trap micro- or nanoscale magnetic particles with sufficient force to resist cell generated traction forces. This was accomplished by adapting a femtosecond laser etching technique previously developed to create micro- and nanoscale spikes on flat silicon surfaces [15,31] to create magnetizable substrates that locally concentrate magnetic field gradients due to the high curvature of the spikes. By creating nanospikes on magnetizable stainless steel, we were able to increase magnetic conductance, durability, and biocompatibility of these substrates for studies with living cells, while minimizing cost.

Another important requirement for cellular microarray-based sensors is defined control of cell positioning and maintenance of this selectivity over time. This requires a surface that is resistant to adsorption of molecules and proteins present in the growth medium or secreted by the cells that might support cell adhesion. We were able to...
inhibit protein adsorption by coating the surface of our devices with a uniform smooth layer of PDMS treated with Pluronic detergent, as previously described [16]. This technique proved highly effective for our purposes, as beads and cells were easily removed and replaced on these devices with extremely high fidelity of patterning, even after culturing cells for 2 days on the magnetic devices. In addition to this highly selective non-fouling support for bead and cell localization, the PDMS-coated, laser-etched microarrays concentrated beads in a uniform layer across the entire surface of each raised nanospiked area which likely helps to facilitate cell adhesion and spreading. An important procedural consideration to achieve this uniform distribution is that careful attention must be paid to the final density of beads on the magnetic field concentrators in order to avoid excessive bead tapering on the magnetized nanospiked regions.

Although our experiments demonstrate the utility of these nanostructured surfaces for concentrating microscale magnetic beads, this system also should be compatible with nanoscale magnetic particles. In fact, beads with diameters closer to the size scale of the curvature of the nanospikes may be trapped more effectively by the magnetic gradients generated by each spike. Moreover, reducing the size of ECM-coated beads, in turn, reduces the area available to form cell adhesive sites. Based on studies of force–size relationships of cell adhesions to ECM [22,24], we expect that as bead size is decreased, cell tractional forces transmitted to individual beads will also decrease, thereby lowering the magnetic pulling force required to immobilize individual beads in presence of traction forces exerted by adherent cells.

Other methods for controlling cell positioning and spreading have been described in the context of reconfigurable cell-based processing devices [32]. However, the system described here offers several advantages, including comparative simplicity and flexibility, the ability to create large microarrays of cells that can be analyzed in parallel, and low cost and durability which are critical features for long-term commercial success. The magnetic ‘virtual substrate’ described here also allows for nearly unlimited array configurations, limited only by the resolution of the laser-etching process. Furthermore, because the cell-adhesive molecules are supplied by the removable magnetic beads, the user can easily specify the molecule used to mediate cell attachment for a given application. This same feature may be utilized in conjunction with other technologies such as microfluidics to create reconfigurable microarrays of magnetic microbeads-coated with biological molecules (e.g., DNA, proteins) in devices for genomic or proteomic applications.

A key element of this device is the ability to selectively control cell shape and viability by applying magnetic forces to cell-bound magnetic microparticles. The magnetic field gradients we created were sufficient to support increases in projected cell areas by at least 36%. This degree of cell distortion was large enough to prevent apoptosis, and the level of survival increased when the intensity of the magnetic field was increased, thus confirming that even a small degree of cell distortion can prevent apoptosis in adherent cells, as previously described [1]. This magnetic shape-control capability may provide a way to induce cell death within future biodetectors that have been contaminated by a biological pathogen or toxin. Removal of the magnetic field would allow contaminated cells to be washed away and new magnetic beads and cells delivered to the same multiplexed islands (e.g., using microfluidics) to create a reconfigurable cell-based biochip detector system.

5. Conclusions

We have developed multiplexed magnetic field gradient concentrators that can be used in combination with magnetic microbeads coated with cell adhesion molecules to create ‘virtual culture substrates’. These biocompatible materials can be used to culture multiple cells in precise positions at high density, and to selectively induce cell death on command by removing an external magnet, thereby releasing cells to round up and die. Critical to this approach is the development of a method to fabricate arrays of magnetizable nanospikes that locally concentrate magnetic field gradients at the surface of the material, and thereby provide a way to trap these beads with sufficient force to resist high cell traction forces. Non-specific adhesive interactions with the remaining surface of the stainless steel substrate are inhibited by coating the material with a smooth layer of Pluronic detergent-treated PDMS. This simple system that has minimal power requirements and provides a reconfigurable interface for control of cell position and viability may be useful for future cell-based or biomolecule-based biodetectors, screening technologies, computers and other microsystems devices.

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References