

Assaying stem cell mechanobiology on microfabricated elastomeric substrates with geometrically modulated rigidity

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Published online 27 January 2011; doi:10.1038/nprot.2010.189

We describe the use of a microfabricated cell culture substrate, consisting of a uniform array of closely spaced, vertical, elastomeric microposts, to study the effects of substrate rigidity on cell function. Elastomeric micropost substrates are micromolded from silicon masters comprised of microposts of different heights to yield substrates of different rigidities. The tips of the elastomeric microposts are functionalized with extracellular matrix through microcontact printing to promote cell adhesion. These substrates, therefore, present the same topographical cues to adherent cells while varying substrate rigidity only through manipulation of micropost height. This protocol describes how to fabricate the silicon micropost array masters (~2 weeks to complete) and elastomeric substrates (3 d), as well as how to perform cell culture experiments (1–14 d), immunofluorescence imaging (2 d), traction force analysis (2 d) and stem cell differentiation assays (1 d) on these substrates in order to examine the effect of substrate rigidity on stem cell morphology, traction force generation, focal adhesion organization and differentiation.

INTRODUCTION

Mounting evidence suggests that physical signals in the cellular microenvironment, particularly matrix rigidity, can mediate stem cell differentiation¹⁻⁸. In early studies, mouse mammary epithelial cells were observed to increase differentiation when grown on soft collagen gels, as opposed to tissue culture plastic⁹. Tubulogenesis of human umbilical vein endothelial cells (HUVECs) was also shown to depend on the underlying substrate rigidity¹⁰. Recent studies have directly shown that matrix rigidity may regulate stem cell differentiation¹¹⁻¹⁶. Human mesenchymal stem cells (hMSCs) grown on polyacrylamide (PAA) gels alter their properties in relation to substrate rigidity^{11,12}. This landmark work further demonstrated that substrate rigidity defines lineage commitment of hMSCs¹². Similarly, substrate rigidity has recently been shown to regulate neural and skeletal muscle stem cell function, including differentiation^{13,15}. Another recent study also suggests that hMSCs are quiescent when grown on PAA gels, with properties similar to bone marrow¹⁷. These cells are arrested in their progression through the cell cycle, but can be induced to re-enter the cell cycle and differentiate when presented with a more rigid substrate.

It remains elusive how mechanical signals in cell microenvironments are transduced into biochemical and cellular functional responses, a process known as mechanotransduction. Mechanotransduction is believed to depend, in part, on myosin-based cytoskeletal (CSK) tension and the integrin-based transcellular focal adhesions (FAs) that physically tether the CSK to the extracellular matrix (ECM)^{6,18-26}. Indeed, the CSK structure and tension depend on matrix rigidity²⁷⁻³². Fibroblasts grown on stabilized collagen gels generate contractile traction forces within the gels, form stress fibers and assemble fibronectin into fibrils; however, fibroblasts cultured on freely floating gels do not demonstrate these behaviors³². In another study, magnetic beads coated with RGD, a peptide sequence that binds integrins, were deposited on cells and magnetically twisted to apply a shear stress to the cell surface. The resultant observation was that cell stiffness increases

with bead deformation^{30,31}. A recent study using optical tweezers further suggests that cells sense ECM rigidity at individual adhesion sites, and that CSK tension at these adhesion contacts responds proportionally to ECM rigidity²⁷. Together, these studies support the involvement of CSK tension in the mechanotransduction process.

The CSK tension-mediated mechanotransduction process may involve integrin-based FA stress signaling^{25,26,33-36}. Integrins physically tether the CSK to the ECM, and further cluster to activate biochemical signaling networks by nucleating FA signaling proteins³⁷⁻⁴². Because FAs provide both the mechanical linkage between the CSK and the ECM and a scaffold for intracellular signaling, it is thought that FAs provide a conduit to sense mechanical stimuli and transduce them into biochemical responses important for the regulation of stem cell differentiation^{5,7}. Indeed, both externally applied and intracellular CSK forces at integrins have been shown to alter FA assembly and downstream FA signaling^{27,43-46}. Thus, it is plausible that ECM mechanics regulate hMSC differentiation by increasing CSK tension, which is then transduced into biochemical signals through increased FA stress and modified FA signaling.

Accordingly, numerous methods have been developed to examine how forces, both sensed and exerted at FAs, regulate biochemical responses and ultimately, cell function. In particular, deformable substrates with precisely engineered mechanical properties have been used extensively. The first substrates of this kind consisted of ultrathin silicone films, which are compliant to the extent that adherent cells are able to induce wrinkles within the film when they contract⁴⁷. However, it is inherently difficult to quantify the traction forces exerted by cells from the wrinkling patterns. Consequently, more advanced techniques to quantify traction forces have been engineered, such as gelatin and PAA gel-based traction force microscopy (TFM), microfabricated horizontal cantilevers and elastomeric substrates⁴⁸⁻⁵³.

Among these approaches, PAA gel-based TFM and microfabricated elastomeric micropost arrays are the two most widely adopted techniques for measuring traction forces. In TFM, fluorescent beads are embedded near the surface of a ligand-functionalized PAA gel that has been pre-stressed against a rigid surface^{49,50}. Cells cultured on this substrate exert traction stresses that deform the gel principally in the plane of the surface. These deformations are generally orders of magnitude smaller than the thickness of the gel and can be tracked by observing the displacements of the embedded beads. Consequently, by treating the PAA gel as an incompressible, linearly elastic material of semi-infinite thickness, the traction field $F(r)$ and displacement field $u(r)$ are related by the Fredholm integral equation of the first kind:

$$u_i(r) = \int dr' G_{ij}(r-r') F_j(r') \quad (1)$$

where $G(r-r')$ is the tensorial Green's function representing the displacement at r caused by the application of a point force at r' . Given $u(r)$, which is measured from the bead displacements, equation (1) must be inverted to solve for $F(r)$. This is a computationally intensive, ill-posed problem. Moreover, to achieve stable unique solutions, regularization schemes must be implemented, such as restricting traction forces to specific sites of adhesion and imposing constraints on the deformation field of the cell^{50,54,55}. Despite these limitations, TFM has been continually refined with the development of faster computational algorithms such as Fourier-transform traction cytometry and improved methods for computing the displacement field from beads^{54,56,57}.

Elastomeric micropost arrays represent a drastically different approach to measure traction forces^{53,58}. Here, a substrate consisting of arrays of uniformly spaced, vertical, elastomeric posts is fabricated with photolithography and replica molding with the silicone elastomer polydimethylsiloxane (PDMS). After ECM proteins are microcontact printed across the tips of these posts, cells are able to adhere, spread out and exert contractile forces that deflect underlying posts. Each post, therefore, functions as a cantilever. For tip deflections that are small compared with the height of the posts, the posts can be conveniently modeled as linearly elastic beams subjected to pure bending. The force F applied at the tip and the resultant deflection x are described by

$$F = \left(\frac{3EI}{L^3} \right) x \quad (2)$$

where E is the elastic modulus, I is the area moment of inertia and L is the height of the post. The term contained within the parentheses, referred to as the spring constant, is therefore a measure of the stiffness of the post. Compared with TFM, traction forces are relatively simple to compute with the micropost arrays. The first generation of micropost arrays had relatively wide post-to-post spacing that constrained cell spreading and movement⁵³. However, this concern has been mitigated with the development of more closely spaced micropost arrays⁵⁸⁻⁶⁰.

In addition to traction force measurement, deformable substrates are also used to modulate substrate stiffness and, thereby, affect the traction stresses generated by adherent cells. Gels composed of natural ECMs such as collagen-I, fibrin, Matrigel or synthetic materials such as PAA and PEG have been created with defined stiffness^{28,61}. Gels derived from natural ECMs more closely mimic the *in vivo*-like environment for cells, as they present adhesive ligands in native conformations and may sequester other components such as growth factors⁶². However, the bulk mechanical properties

of these gels are difficult to control. Changes in stiffness cannot be decoupled from other parameters such as ligand density and fiber thickness⁶¹. Moreover, gels composed of filamentous, semi-flexible biopolymers such as collagen and fibrin exhibit nonlinear elasticity in that they stiffen when subjected to low strains^{63,64}. In contrast, gels derived from synthetic materials are chemically inert and must be functionalized with adhesive peptides and proteins using linker chemistry. The advantage is that synthetic gels have well-defined bulk mechanical properties. PAA gels, for example, are linearly elastic over a broad range of strains and have been formulated to exhibit elastic moduli ranging from 2 Pa to 55 kPa^{28,49}. Yet, synthetic and natural gels alike are not immune to molecular-scale changes in porosity, wettability, hydration, polymer chain mobility and binding properties of immobilized adhesive ligands that accompany changes in bulk stiffness^{65,66}. A recent study using synthetic gels for human pluripotent stem cells indicates that these molecular-scale changes can have profound effects on stem cell function⁶⁷.

Similar to synthetic and natural gels, elastomeric micropost arrays can also be used to control substrate stiffness. However, as we describe in this protocol, the stiffness of micropost arrays can be controlled independently without affecting bulk or nanoscale mechanics, and adhesive ligand topography. This is achieved by exploiting the fact that the spring constant of a micropost is inversely proportional to the height of the post to the third power. By fixing the micropost cross-sectional area and changing only the height, micropost arrays of varying stiffness can be fabricated^{58-60,68,69}. The bulk and nanoscale properties of the PDMS remain unchanged as does the amount of ECM functionalized onto the tips of the microposts.

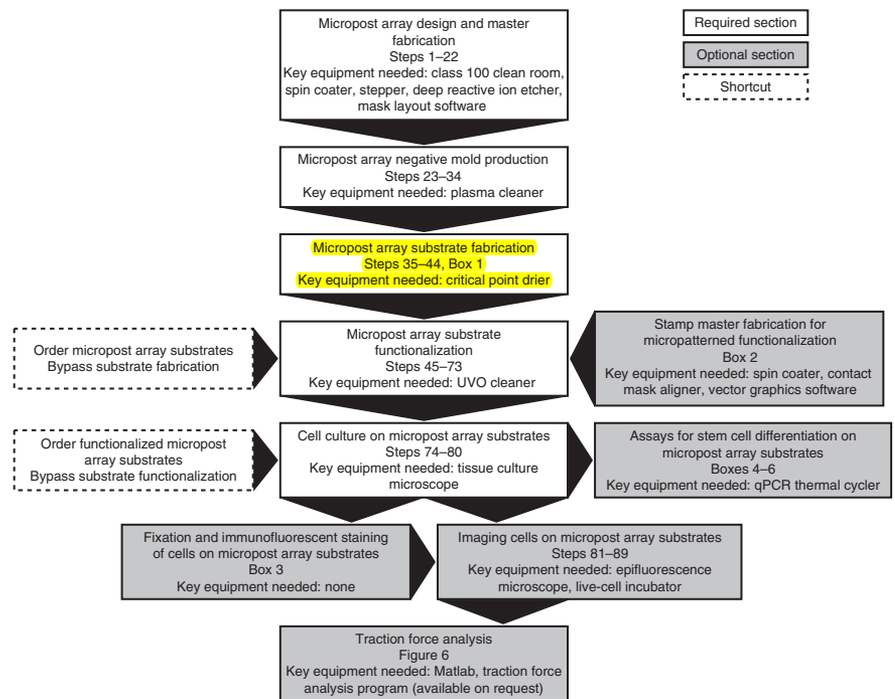
To demonstrate the utility of the elastomeric micropost array as an effective means for both traction force measurement and substrate rigidity modulation, we describe in detail our protocol for studying of the effect of micropost rigidity on stem cell function, including differentiation. This protocol, which is presented in a modular manner, will cover the steps required to generate a micropost array substrate with minimum functionality (Fig. 1), as well as a number of optional applications, focusing on traction force imaging and analysis and stem cell differentiation assays. As originally described in a *Nature Methods* publication by Fu *et al.*⁶⁰, the protocol indicates where the user can bypass steps that require special equipment, by requesting micropost array substrates from our group (<http://www.seas.upenn.edu/~chenlab/micropostform.html>). Furthermore, to inform the reader as to the suitability of this protocol for his/her particular system of study, we also discuss the applications of this method as well as the advantages and disadvantages of this method compared with similar methods; we also provide technical insights for adapting the protocol for a broad range of applications.

Applications of micropost array substrates

Deformable substrates have traditionally been used to study single cell-substrate interactions such as cell locomotion, traction force generation, FA dynamics and CSK mechanics^{50,52,53,70,71}. However, the elastomeric micropost arrays have also been used to study a broader range of biological questions, illustrating the versatility of this tool^{58,72-82}. One notable application has been in the study of multicellular systems. Using an array of cylindrical microposts similar to the ones described in this protocol, du Roure *et al.*⁵⁸ measured the forces generated by migrating sheets of epithelial cells. Saez *et al.*⁷⁶ modified the microposts to have an oval cross-section



Figure 1 | Flow diagram of the different sections of the protocol. The required and optional sections are color-coded as white and gray, respectively, with arrows indicating the order in which the sections are performed. **Note that some sections require key equipment, such as a plasma cleaner, critical point drier or UVO cleaner.** If these machines are not available, limited quantities of either non-functionalized or functionalized micropost array substrates can be ordered through our online service (<http://www.seas.upenn.edu/~chenlab/micropostform.html>). These shortcuts are indicated by boxes with dashed borders on the flow chart.



and, therefore, exhibit stiffer properties along their major axis than along the minor one. On micropost arrays of anisotropic stiffness, epithelial sheets were observed to grow and migrate along the direction of greatest rigidity⁷⁶. In studies in which unfettered migration is not desired, multicellular constructs have been constrained on the microposts with patterned micro-contact printing. Nelson *et al.*⁷² used this approach to grow a monolayer of endothelial cells on the microposts in the shape of an asymmetric annulus and observed that regions of high tractional stress, such as along the convex, outer boundary of the annular monolayer, correlated with regions of greater proliferation. Similarly, Ruiz and Chen⁷³ found that hMSCs patterned in various multicellular geometries preferentially expressed osteogenic markers in regions of greater stress and adipogenic markers in regions with lower stress.

Other studies have used micropost arrays to measure the forces transmitted through cell-cell contacts. The micropost array substrate differs from continuous, deformable substrates in that forces exerted on each post normally are not transmitted to neighboring posts through the substrate. Liu *et al.*⁷⁴ exploited this feature to examine the role of cell-cell tugging force on adherens junction growth. To increase the occurrence of two cells forming a single contact, endothelial cells were seeded on bowtie-shaped micropatterns on the microposts, such that only one cell could occupy each half of a bowtie. Paired cells exist in a state of quasi-static equilibrium in which the net traction force sums to zero. As such, intercellular tugging force can be calculated from the vector sum of the traction forces exerted by each cell. Using this approach, Liu *et al.* observed a positive correlation between junction size and tugging force. Ganz *et al.*⁷⁹ examined adherens junction-mediated forces in a different manner by immobilizing an N-cadherin-Fc chimera, which mimics the adherens junction protein N-cadherin, onto the microposts. They showed that C2 myogenic cells grown on these substrates exerted forces on the cadherin contacts that were similar in magnitude to those observed at FAs.

A third broad application of the microposts has been in the development of models of biological processes. Two similar but distinct approaches have been developed to study the mechanics of leukocyte transmigration through an endothelial monolayer. Rabadzey *et al.*⁷⁷ cultured an unpatterned monolayer of endothelial cells on microposts, subjected these cells to laminar shear flow in a

parallel plate flow chamber and observed the effects of neutrophil adhesion and transmigration on traction forces generated within the monolayer. Liu *et al.*⁷⁸ patterned ‘mini’ monolayers of five to ten endothelial cells and titrated the monocyte concentration to isolate the effects of individual monocytes on each monolayer. Both studies observed that firm adhesion and transmigration of leukocytes triggered increases in endothelial traction force, which was greatest at the site of leukocyte attachment. In yet another unique implementation, micropost arrays have been used to examine the mechanics of platelet-mediated clotting or thrombus formation. Single platelets are too small to culture on micron-scale microposts. Instead, Liang *et al.*⁸⁰ formed microthrombi, composed of many platelets, on top of the microposts; they were able to obtain dynamic measurements of the clotting force as a function of thrombin activity.

Aside from its applications for passively reporting traction force, the micropost array substrate has been integrated with actuation technologies as a means to both apply forces to cells and to measure responses of traction forces. Studies have shown that application of tangential forces at integrin-mediated adhesions leads to reinforcement of the CSK-ECM linkages^{27,43}. Motivated by this finding, Sniadecki *et al.*⁷⁵ engineered micropost array substrates in which magnetic nanowires were embedded in a sparse number of microposts. These select microposts can then be actuated under a uniform magnetic field to impart nanoNewton forces to individual adhesions of adherent cells. The surrounding passive microposts are able to report changes in global traction force induced by local force application. Interestingly, Sniadecki *et al.* observed enhanced FA assembly only at the site of force application and found that local forces could induce long-range relaxation of traction forces.

The most insightful studies with micropost array substrates have taken advantage of the ability to carry out paired analyses of the traction forces, with a functional output such as proliferation or adherens junction assembly. Fu *et al.*⁶⁰ have taken this approach

a step further by showing that the early contractile state of single hMSCs that have been exposed to differentiation media can predict the later onset of their differentiation. In that study, individual cells were constrained to micropatterns on the micropost arrays and monitored daily for changes in their traction forces. After 1 week, the cells were stained for differentiation markers. Subsequently, Bayesian classifier analysis was used to determine whether the traction forces measured on different days could predict the differentiation outcome for each cell.

Advantages and disadvantages of micropost array substrates

The existing applications of micropost array substrates highlight a number of advantages and disadvantages for adapting this tool for new applications. One positive attribute of the micropost arrays is that they have been used to study diverse types of cells, such as epithelial cells, endothelial cells, fibroblasts, leukocytes and mesenchymal stem cells, in well-defined adhesive and mechanical contexts. In addition to mesenchymal stem cells, it is likely that other mechanosensitive stem cell types, such as hematopoietic, skeletal muscle and embryonic stem cells, can also be studied on the micropost arrays^{15,83–86}. Moreover, the ability to microcontact-print defined patterns of different types of proteins on the micropost arrays provides control over the adhesive topography that is difficult to achieve on gel-based substrates. Another advantage of the micropost arrays is that they are very amenable to paired analyses of traction forces and also to a second functional output that is not directly linked to traction force. Whereas cells cultured on PAA gels need to be detached in order to measure their traction forces, cells cultured on micropost arrays do not, and they can be fixed and stained for cellular components such as differentiation markers or CSK and adhesion proteins.

Despite all of their advantages, the micropost array substrates cannot completely replace other approaches for measuring traction forces or modulating substrate rigidity. One potential limitation is generating ultracompliant micropost arrays that are equivalent to the softest PAA gels. Neural differentiation, for example, has been observed on PAA gels, with elastic moduli ranging from 0.1 to 1 kPa¹². In comparison, the softest microposts that have been used to study stem cell differentiation have an equivalent stiffness of 1.5 kPa⁶⁰. It is possible to generate ultra-compliant microposts, either by fabricating taller microposts or using a lower ratio of cross-linking agent to PDMS prepolymer to cast substrates. However, there are significant technical issues such as preventing very soft microposts from sticking to each other during substrate fabrication and functionalization. As such, PAA gels are currently better suited than micropost arrays for culturing cells on ultra-compliant substrates.

The discrete topography of the microposts can be either an advantage or disadvantage, depending on the application. In the studies of cell-cell tugging force, localized force actuation and transmigration, the discrete topography is beneficial in that it either prevented transmission of forces through the substrate or provided space for cells to move vertically^{74,75,77,78}. However, if the goal is to understand how cells mechanically interact with each other through the substrate or respond to global substrate deformations, then continuous deformable substrates may be more suitable than micropost arrays^{87,88}. The discrete topography of the micropost array raises another potential issue in that altering post density and adhesive area can influence the distribution of cellular adhesions, which

can possibly affect cellular behavior^{59,68}. Therefore, it is paramount to use the knowledge of how the cell types of interest adhere and spread in tissue culture when designing the geometric parameters of a micropost array.

Experimental design

Special equipment needs for fabricating micropost array masters.

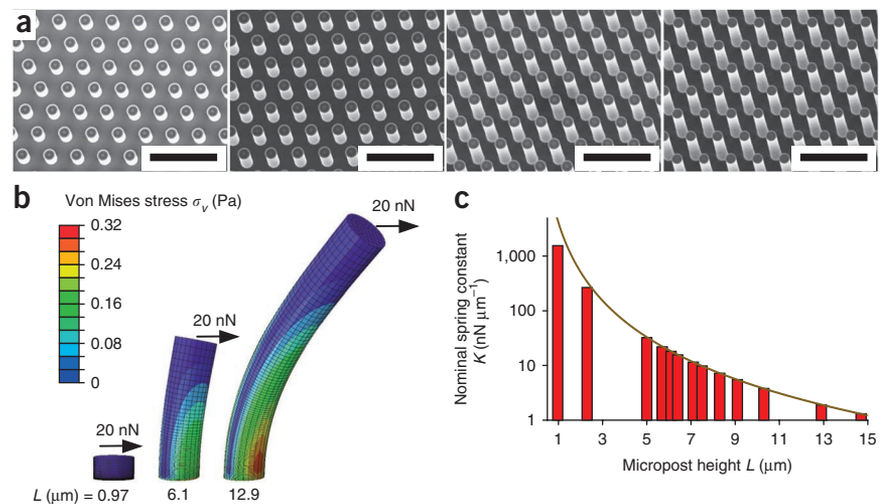
This protocol describes the fabrication of silicon micropost array masters with a micropost diameter of $\sim 2 \mu\text{m}$ and center-to-center spacing of $4 \mu\text{m}$. To fabricate features with these dimensions, one must first have access to an advanced microfabrication facility such as the Massachusetts Institute of Technology (MIT) Microsystems Technology Laboratory or Cornell Nanofabrication Facility. There are two special pieces of equipment at these facilities that are required to achieve high-density micropost arrays. The first is a lithographic stepper to pattern the photoresist that will define the post diameter and spacing in the micropost array. The stepper patterns photoresist by projecting UV light through a reticle photomask and reduction lens onto the wafer. The reduction lens focuses the collimated light on a region of the wafer smaller than the reticle, such that the features on the reticle are scaled down. This not only allows the stepper to pattern features in the photoresist as small as 500 nm but is also economically advantageous, as photomasks with increasingly smaller features become prohibitively expensive. The second piece of equipment is a deep reactive ion etcher (DRIE) that can etch silicon microstructures with very high aspect ratios. It accomplishes this feat by iteratively passivating and etching the wafer, also known as the Bosch process⁸⁹. During the passivation step, the wafer is uniformly coated with C_4F_8 , a chemically inert Teflon-like polymer. This polymer is then removed by a nearly isotropic plasma etch containing SF_6 ions. Rapid cycling between the two steps minimizes lateral etching, resulting in a very directional etch. Previously, the Bosch process has been used to fabricate arrays of holes in silicon that can be used to cast PDMS microposts⁵⁸. Here, we use the same process to fabricate the inverse structure consisting of arrays of silicon microposts (Fig. 2a). This subtle but significant difference confers technical advantages in replica micromolding of the PDMS micropost substrates, which will be elaborated upon below.

If access to an advanced microfabrication facility is not practical, micropost array masters with larger micropost diameters and spacings can be fabricated with more basic microfabrication equipment. A detailed protocol for such approaches is described elsewhere⁹⁰. Briefly, a master is generated by spin-coating a layer of SU-8 photoresist, from ~ 7 to $12 \mu\text{m}$ thick, on a silicon wafer, and then exposing the photoresist to UV light through a photomask on a contact mask aligner. Development of the unexposed photoresist will then leave behind an array of SU-8 photoresist microposts that can be used for replica-molding of the PDMS micropost array substrates. This approach obviates the need for a stepper and DRIE machine. However, the theoretical resolution R of the contact photolithography is

$$R = \frac{3}{2} \sqrt{\frac{\lambda z}{2}} \quad (3)$$

where λ is the wavelength of light and z is the photoresist thickness⁹¹. SU-8 maximally absorbs UV light with a wavelength of

Figure 2 | Characterization of micropost array masters and substrates. (a) Scanning electron micrographs of silicon micropost array masters of four different heights. From left to right, the heights of the microposts are 2.3, 5, 8.3 and 12.9 μm . Scale bars, 10 μm . (b) FEM simulations of the deflection of PDMS microposts in response to an applied force at the tip. (c) The nominal spring constant (K), as computed from FEM analysis (bars) and equation (2) (curve), is plotted for PDMS microposts of different heights L . Reprinted with permission from reference 60.



365 nm. Thus, to fabricate microposts with a diameter of 2 μm , the thickness of SU-8 photoresist cannot exceed 9.7 μm . Such posts are shorter than several of the taller silicon micropost array masters that we have generated. Moreover, the theoretical resolution is rarely achieved because of defects such as uneven photoresist coating and mask damage.

Design considerations for fabrication of reticle photomask and silicon micropost arrays. The critical parameters to consider when designing the reticle photomask for patterning the micropost array master are the center-to-center spacing and diameter of the microposts. These two parameters are interdependent and should be optimized for the cell types of interest. For example, closely spaced arrays with a center-to-center spacing of 2 μm are well-suited to measure forces in small cells such as epithelial cells⁵⁸. An even smaller spacing in the submicron range may be necessary for smaller cellular bodies such as platelets⁸⁰. We have found that, below a center-to-center spacing of 4 μm , there is no noticeable difference in the spread area or morphology of intermediate-to-large-sized cells such as mesenchymal stem cells, fibroblasts and endothelial cells when compared with cells on continuous substrates^{59,60}. However, spacings ranging from 6 to 9 μm adversely affect the ability of cells to spread and migrate on micropost arrays as they would on continuous substrates⁵⁹. Once the post spacing has been chosen, a suitable post diameter must be used such that adjacent posts do not stick to each other. We and others have observed that using a post diameter that is equal to half the center-to-center spacing provides excellent spatial resolution while mitigating excessive post collisions and stiction^{59,60,92}. Posts with a diameter larger than half the center-to-center spacing can still be used to measure traction forces. However, the sensitivity of the measurement will be lower because of the smaller distances over which adjacent posts can deflect before they collide. Decreasing the post diameter too much may also produce problems with measurement sensitivity. We found that for post diameters ranging from 0.67 to 0.83 μm , cells exerted constant strain energies on micropost arrays of different post densities, such that post deflections were almost indistinguishable from noise at the highest post densities³⁹. Thus, for practical purposes, it is recommended that a post diameter of at least 1 μm be used, which will yield an adhesive area per post that is at the lower bound of the FA areas reported in the literature¹⁸. Taking these variables into consideration, the reticle photomask that we use in this protocol is designed to pattern micropost arrays with a center-to-center spacing of 4 μm and a micropost diameter of 2 μm .

In addition to optimizing the micropost array geometry, a number of other issues must be addressed in the design of the reticle photomask. First, to facilitate microcontact printing of fibronectin onto the micropost arrays, large flat structures should be placed around the arrays to serve as weight-bearing structures that prevent the stamp from collapsing the microposts. In our design, we divided the micropost array into four 2 mm \times 2 mm quadrants, separated by rectangular flat structures. Second, the lithographic stepper used in this protocol is equipped with an $\times 5$ reduction lens. Therefore, the scale of the array of micropost circles in our reticle photomask is five times larger than it would be on the photoresist. This will be different for a stepper with a different reduction lens. In addition, the micropost masters described in this protocol are comprised of cylindrical pillars, as opposed to cylindrical pits that have also been used elsewhere^{58,76}. There are two main advantages to using cylindrical pillars. First, negative replica molds can be cast from pillar-based silicon masters in batches and then used repeatedly to generate large numbers of PDMS micropost array substrates. This minimizes the frequency of casting from the silicon masters, which are difficult to replace if damaged during handling. Second, PDMS micropost substrates are cast on a rigid backing such as a glass cover slip or slide. If pit-based masters are used, the glass cover slip would have to be pressed against the rigid silicon master during curing. Peeling a rigid substrate away from a rigid master is difficult and can cause one or both devices to break. As positive-tone photoresist is used to pattern the pillar arrays in this protocol, the reticle photomask should be negative, with the circle patterns opaque and the surrounding area transparent. The reticle photomask is designed with mask layout software such as L-Edit Pro or AutoCAD, and must be outsourced to a commercial mask-writing company for production.

During the actual fabrication of the silicon micropost array masters, photolithographic patterning of the micropost array and subsequent DRIE etching of the silicon are the key steps that must be optimized. This process will likely involve frequent inspection of the silicon masters at each stage of the process flow with an optical microscope and a scanning electron microscope. For the photolithography step, UV exposure conditions must be optimized with test exposures to ensure that the patterned photoresist features reproduce the reticle mask features with high

fidelity. Moreover, the photoresist should be at least 1 μm thick to provide a sufficient protective layer against the DRIE plasma, which etches the photoresist ~ 100 times more slowly than silicon. For the DRIE step, standard recipes have been developed that use an inductively coupled plasma source to create plasmas with high electron density, low pressure and low energy. Although these plasmas can result in high etch rates and directional etching, some lateral etching still occurs, which may lead to microposts with smaller diameters than expected. As such, the aspect ratio of micropost height to diameter is limited by the DRIE process to be less than ~ 20 . In our masters, the expected micropost diameter was 2 μm , but the actual micropost diameter was 1.83 μm . Although this was within our tolerances, if the discrepancy between the expected and actual micropost diameter is too large, then different parameters in the photolithography and DRIE steps will have to be optimized.

Characterization of mechanical properties of micropost array substrates. Before using elastomeric micropost array substrates for experiments, the mechanical properties of the microposts should be rigorously characterized. Equation (2) provides a good approximation of the spring constants of microposts for small post deflections. However, for larger deflections, which are normally observed with microposts that have aspect ratios greater than 5, the linear approximation of equation (2) is not valid. Moreover, for short microposts with aspect ratios less than 1.5, equation (2) is also not an accurate approximation because of a significant shearing component in the post deformation⁹³. One way to validate the spring constant approximations from equation (2) is to use a micromanipulator to push a calibrated glass micropipette tip against the tip of a micropost^{53,94}. Because the spring constant of the micropipette tip is known and the deflections of the micropipette tip and micropost can be measured, the spring constant of the micropost can be calculated. This method has previously been used to determine the spring constant of microposts with a diameter and height of 3 and 10 μm , respectively^{53,94}. However, for microposts with smaller diameters and microposts with large aspect ratios, calibration with micropipettes becomes increasingly difficult for a couple of reasons. First, the glass micropipettes generally have tip diameters $> 0.5 \mu\text{m}$ and would be difficult to position near the tips of smaller microposts. Second, tall microposts may be hard to calibrate, as they will not present a sufficient reactive force to deflect the much stiffer micropipette tip.

Therefore, to more accurately determine the spring constant of the PDMS micropost, we have used a finite element modeling (FEM) package such as ABAQUS (Simulia, Dassault Systèmes) (Fig. 2b,c). We modeled the micropost as a neo-Hookean hyperelastic cylinder composed of hexahedral mesh elements. The diameter and range of heights for the micropost were measured from scanning electron microscopy (SEM) and surface profilometry. PDMS is known to change its mechanical properties over time, depending on the curing times and temperatures that are used⁹⁵. Therefore, to determine the Young's modulus of the micropost, we cured strips of PDMS at 65 and 110 $^{\circ}\text{C}$ for different lengths of time and then stretched the strips on a tension tester such as the 5848 MicroTester (Instron). These tests subsequently showed that the Young's modulus stabilizes at a value of $2.5 \pm 0.5 \text{ MPa}$ for PDMS that has been cured at 110 $^{\circ}\text{C}$ for 20 h. To simulate bending of the micropost, we specified fixed boundary conditions for the

base of the micropost and applied a range of traction shear loads across the top of the post. Post displacement was measured at the center node on the top surface, and then plotted against the applied force. The nominal spring constant K can then be determined by computing the slope of the force-displacement curve as the displacement approaches zero. Our FEM simulations assume that the microposts are fixed against a rigid substrate, when in fact they are attached to a thin elastic PDMS layer. As such, K will have to be adjusted to account for substrate warping. Correction factors for this adjustment have recently been determined for a wide range of micropost aspect ratios⁹³.

Fabrication considerations for a micropatterned stamp master.

To micropattern islands of fibronectin on the micropost array substrate, a microstructured PDMS stamp consisting of raised and recessed regions is used. This stamp is cast against a micropatterned photoresist-coated wafer that is itself produced using conventional photolithography⁹⁶. Compared with the micropost array masters, a micropatterned stamp wafer is much simpler to fabricate. The spread areas of single cells generally range between 100 and 10,000 μm^2 . As such, the smallest islands of fibronectin that normally would be micropatterned have a minimum feature dimension of 10 μm . Photoresist features of this size are readily patterned with a contact mask aligner. Moreover, photolithography can be performed with a transparency photomask that can be designed with vector graphics software such as Adobe Illustrator and obtained through low-cost, high-resolution photoplotting services. The main challenge with micropatterning is to make sure that the microstructured stamp does not collapse under its own weight and, thereby, transfer fibronectin beyond the micropatterns. Microstructured stamps can collapse if the raised regions buckle or if the recessed regions collapse, also known as roof collapse. Generally, buckling is not a concern as the aspect ratio of the raised regions will be < 0.5 (ref. 96). Theoretical and experimental studies on stamp deformation have suggested that roof collapse can be avoided if the ratio of the height of the raised regions to the width of the recessed regions is at least 0.3 (ref. 97). To illustrate an example, if the raised regions are 100 $\mu\text{m} \times 100 \mu\text{m}$ squares, the height of the raised regions should be $< 50 \mu\text{m}$. If the height is 30 μm , adjacent raised squares must be spaced no more than 100 μm apart. If the height is 15 μm , adjacent squares must be no more than 50 μm apart. Therefore, the spacing of raised features should dictate the viscosity of the photoresist used to pattern the stamp master. More viscous photoresists such as SU-8 2025 are suitable for stamps with more widely spaced patterns, whereas thinner resists such as SU-8 2010 are ideal for more closely packed patterns.

Alternative strategies for substrate functionalization. Fluorescent labeling of microposts for traction force analysis can be tailored for different studies. Specifically, other molecules with similar structure to DiI (1,1'-dioleil-3,3',3'-tetramethylindocarbocyanine methane sulfonate) but different fluorescent spectra can be used to label the microposts. The dye DiD, e.g., fluoresces in the far-red, or Cy5, channel. This is particularly useful if the Cy3 channel, which is used for DiI, must be used for another purpose such as fluorescently labeling cellular structures. Moreover, labeling with DiD is also helpful if the Cy2 channel will be used to image an experimentally important signal, such as a GFP-fused



protein. The reason for this is that DiI can be weakly detected in the Cy2 channel, known as bleed-through. For both DiI and DiD, it is important to optimize the labeling concentration to ensure sufficient signal-to-noise ratio while minimizing any undesirable bleed-through. Alternatively, lipophilic dyes can be omitted entirely by using fluorophore-conjugated proteins. Fluorescently labeled fibronectin can be mixed with non-labeled fibronectin and microcontact-printed to label only the tips of the microposts^{58,76}. Another post labeling method is to adsorb Alexa Fluor 594-conjugated BSA to the shafts and base of the microposts⁸². The main reason to use this strategy is if the cell type of interest takes up DiI in significant amounts, thereby presenting severe artifacts for traction force analysis. If this strategy is used, a concentration of 20 $\mu\text{g ml}^{-1}$ BSA-Alexa Fluor 594 is recommended as a starting point. Further optimization of the labeling concentration is necessary, as BSA-Alexa Fluor 594 may hinder the adsorption of F127 Pluronics and thereby allow cells to crawl down the posts.

Fibronectin is not the only type of protein that can be printed on the microposts. Other ECM proteins such as collagen and vitronectin can be used as well. Adsorption conditions and protein concentrations will have to be optimized empirically for different proteins. For example, collagen must be adsorbed to a stamp in a 1% (vol/vol) acetic acid solution to ensure that the collagen fibrils do not precipitate. Another important consideration is that some proteins may not be printable, as doing so would destroy their biological activity. For these situations, a potential solution would be to covalently link the protein of interest to a linker protein such as an antibody fragment or biotin. The protein could then be 'printed' on the microposts by first printing a binding partner of the linker protein, such as protein G or avidin, and then immersing the substrate in a solution with the protein of interest.

Microcontact printing of adhesive proteins and subsequent treatment with Pluronic to block nonspecific protein adsorption to the shafts of the microposts ensures that cells are only able to adhere to the tips of the microposts and not crawl in between. A UVO (ultraviolet-ozone) cleaner is crucial for this process, as it can

temporarily hydrophilize a PDMS micropost substrate. Adhesive proteins that are adsorbed onto a PDMS stamp can then be transferred onto a relatively hydrophilic micropost substrate^{98,99}. Moreover, the surface of UVO-treated PDMS is reversibly modified, unlike the surface of plasma-treated PDMS, which is glass-like and much more hydrophilic¹⁰⁰. The mild hydrophilicity of UVO-treated PDMS, therefore, permits Pluronic, which has a hydrophobic polymer backbone, to adsorb to the surface. If a UVO cleaner is not available and preventing cell invasion into the micropost array is not critical, fibronectin may be adsorbed over the entire surface by immersing the substrate in the fibronectin solution^{58,69}.

Cell seeding strategies on the micropost array substrates. Seeding cells on the micropost array substrates is an empirical process in which the main parameters that need to be optimized are seeding density and seeding time. These parameters are dependent on the cell type. Different cell types adhere to fibronectin with differential efficiency. For example, hMSCs and HUVECs attach within 10–15 min of seeding, whereas epithelial cells can take up to 1 h. Different cell types also have different final spread areas. hMSCs are two to three times larger than HUVECs, and both cell types are significantly larger than epithelial cells or immune cells. Therefore, if a significant number of isolated cells on the substrate is desired, hMSCs should be seeded at a lower density of $\sim 1,500$ cells per cm^2 , whereas HUVECs should be seeded at around 5,000–10,000 cells per cm^2 . Note that these seeding densities are significantly higher than the desired density on the substrate. The rationale is that it is better to seed at a high density and wash off excess floaters after a short incubation time than it is to seed at a low density and allow the majority of cells to slowly attach without washing. This minimizes the time that adherent cells spend in suspension. For micropatterned micropost array substrates, these parameters may be different, as the cells will require more incubation time to land on a smaller adhesive surface. In our experience, seeding density can be left unchanged while seeding time is extended to 3 h (or even overnight) to obtain a good percentage of cells adherent to micropatterns on the micropost arrays.

MATERIALS

REAGENTS

- Hydrogen peroxide (30% (vol/vol); Sigma-Aldrich, cat. no. 216763) **! CAUTION** It is an oxidizing agent and is corrosive. Wear goggles, gloves and laboratory coat when handling.
- Sulfuric acid (95% (vol/vol); Sigma-Aldrich, cat. no. 320501) **! CAUTION** It is corrosive. Wear goggles, gloves and laboratory coat when handling.
- Deionized water
- Hexamethyldisilazane (HMDS; Sigma-Aldrich, cat. no. H4875) **! CAUTION** It is flammable. Avoid prolonged exposure or inhalation. Wear goggles, gloves and protective clothing and handle in a properly ventilated chemical hood.
- Megaposit SPR 700-1.0 photoresist (MicroChem) **! CAUTION** It is flammable. Avoid prolonged exposure or inhalation. Wear goggles, gloves and protective clothing and handle in a properly ventilated chemical hood.
- MF CD 26 developer solution (MicroChem) **! CAUTION** It is corrosive and is an irritant. Wear goggles, gloves and protective clothing when handling.
- Shipley AZ4620 photoresist (MicroChem) **! CAUTION** It is flammable. Avoid prolonged exposure or inhalation. Wear goggles, gloves and protective clothing and handle in a properly ventilated chemical hood.
- Acetone (Fisher Scientific, cat. no. A18-4) **! CAUTION** It is an irritant and inflammable.

- Isopropanol (Fisher Scientific, cat. no. A416-4)
- Araldite 2012 epoxy (Freeman Supply, cat. no. 056452)
- SU-8 2002/2010/2025 photoresist (MicroChem) **! CAUTION** It is inflammable. Avoid prolonged exposure or inhalation. Wear goggles, gloves and protective clothing and handle in a properly ventilated chemical hood.
- Norland 68 optical adhesive (Norland Products, cat. no. NOA 68)
- (Tridecafluoro-1,1,2,2-tetrahydrooctyl)-1-trichlorosilane (United Chemical Technologies, part no. T2492) **! CAUTION** It is corrosive and toxic. Wear gloves, goggles and handle in a properly ventilated chemical hood.
- Sylgard 184 polydimethylsiloxane (PDMS, Dow Corning; K.R. Anderson, cat. no. 2065622)
- Ethanol (200 proof; Fisher Scientific, cat. no. 04-355-222)
- Propylene glycol monomethyl ether acetate (Sigma-Aldrich, cat. no. 484431) **! CAUTION** It is inflammable. Avoid prolonged exposure or inhalation. Wear goggles, gloves and protective clothing and handle in a properly ventilated chemical hood.
- Fibronectin (BD Biosciences, cat. no. 356008)
- DiI (1,1'-dioleil-3,3',3'-tetramethylindocarbocyanine methane sulfonate) (Invitrogen, cat. no. D-3886)
- F127 Pluronic (Sigma-Aldrich, cat. no. P2443)



PROTOCOL

- PBS (Invitrogen, cat. no. 10010-023)
- 10× PBS (Invitrogen, cat. no. 70011-044)
- Mammalian cells (e.g., human bone marrow-derived mesenchymal stem cells; Lonza, cat. no. PT-2501)

Cell culture medium (for hMSCs)

- Dulbecco's modified Eagle's medium (low glucose; Invitrogen, cat. no. 11885-092)
- Fetal bovine serum (qualified; Invitrogen, cat. no. 10437-028)
- L-Glutamine (Invitrogen, cat. no. 25030-081)
- Trypsin (0.05% (wt/vol))-EDTA (2 mM) (Invitrogen, cat. no. 25300-054)
- Paraformaldehyde (16% (vol/vol); Electron Microscopy Sciences, cat. no. 15710) **! CAUTION** It is an irritant to eye and skin. Wear goggles and gloves when handling.
- Triton X-100 (Sigma-Aldrich, cat. no. X100)
- Goat serum (Invitrogen, cat. no. 16210072)
- BSA (Sigma-Aldrich, cat. no. A4503)
- Primary antibodies (Anti-vinculin, Sigma-Aldrich, cat. no. V9131; anti-FAK(pY397), BD Biosciences, cat. no. 611722)
- Secondary antibodies (Alexa Fluor 647 goat anti-mouse IgG (H + L), Invitrogen, cat. no. A-21235)
- DAPI (Invitrogen, cat. no. D3571)
- Alexa Fluor 488-conjugated Phalloidin (Invitrogen, cat. no. A12379)
- Fluoromount G (Electron Microscopy Sciences, cat. no. 17984-25)
- Nuclease-free water (Ambion, cat. no. AM9938)
- Qiagen RNeasy Micro Kit (Qiagen, cat. no. 74004)
- 2-Mercaptoethanol (Mallinckrodt Baker, cat. no. 4049-00) **! CAUTION** It is an irritant and inflammable. Wear goggles, gloves and handle in a properly ventilated chemical hood.
- dNTP mixture (dATP, dTTP, dCTP, dGTP, 10 mM each; Invitrogen, cat. no. 18427013)
- Oligo dT₁₂₋₁₈ primer (Invitrogen, cat. no. 18418012)
- RNaseOUT ribonuclease inhibitor (Invitrogen, cat. no. 10777019)
- MMLV reverse transcriptase (Invitrogen, cat. no. AM2043)
- qPCR Taqman probes (Alkaline phosphatase (ALP), Hs00758162_m1; collagen type I, Hs 00164004; GAPDH, Hs 99999905_m1; FrzB, Hs00173503_m1; IBSP, Hs00173720; lipoprotein lipase, Hs00173425_m1; PPARG1/2, Hs00234592_m1; Runx2, Hs00231692_m1; Applied Biosystems)
- TaqMan universal PCRmaster mixture (Applied Biosystems, cat. no. 4304437)
- Oil Red O (Sigma-Aldrich; cat. no. O0625) **! CAUTION** Avoid inhalation of dust and eye contact.
- Formalin (10% (vol/vol); Sigma-Aldrich, cat. no. HT501128)
- Fast blue RR salt (Sigma-Aldrich; cat. no. FBS-25)
- Naphthol AS-MX phosphate alkaline solution (Sigma-Aldrich; cat. no. 85-5)
- Citrate concentrated solution (Sigma-Aldrich; cat. no. 85-4C)

EQUIPMENT

- L-Edit Pro (Tanner EDA) layout/verification software for reticle photomask design. Alternatively, AutoCAD (Autodesk) can be used. Adobe Illustrator (Adobe) should be used for transparency photomasks.
- High-resolution chrome mask (Advance Reproductions or Microtronics)
- Silicon wafers (N-type, 1-0-0 orientation, Ph-doped; Silicon Quest International). Generally, steppers handle 4-inch or 6-inch wafers, whereas contact mask aligners handle 3-inch or 4-inch wafers.
- Wet processing station (with Piranha solution tank and wafer dump rinser; Semifab, Model WPS-400 & 800)
- Wafer spin dryer (Marteq Process Solutions)
- Two contact hot plates
- Vacuum desiccators (Cole-Parmer, cat. no. EW-06514-30 or equivalent)
- Automated photoresist coat and develop system (SSI 150 Dual Track Spinner System, Semiconductor Systems)
- Projection stepper (Nikon NSR-2005i9, Nikon Precision)
- Metallurgical microscope (ME600 or equivalent, Nikon Instruments)
- Surface profilometer (KLA-Tencor-Prometrix P10, KLA-Tencor)
- Deep reactive ion etcher (ICP Deep Trench Etching Systems, Surface Technology Systems)
- Scanning electron microscope (SEM, Zeiss SUPRA 40 High Throughput FESEM, Carl Zeiss)
- Wafer die saw (Disco Abrasive System DAD-2H/6T, Disco Abrasive Systems)
- Tweezers (Electron Microscopy Sciences): wafer tweezers, flat narrow tweezers and fine-tipped curved tweezers are recommended.
- Pyrex crystallizing dishes (Sigma-Aldrich)

- Compressed N₂ gas (Airgas)
- Microscope glass slides (Fisher Scientific)
- UV curing chamber (Electro-Cure 500, Electro-Lite Corporation)
- Glass Pasteur pipettes (Fisher Scientific, cat. no. 136786A)
- Plastic transfer pipettes (Fisher Scientific; cat. no. 137117M)
- Top-loading balance (Denver Instrument, model no. TR-402 or equivalent)
- Aluminum weighing dish (43-mm diameter; Heathrow Scientific, cat. no. HS14521A)
- Two laboratory ovens (Isotemp or equivalent, Fisher Scientific)
- Razor blades
- Plasma cleaner (Plasma Prep II, SPI Supplies)
- Analytical balance (Mettler Toledo, model no. AG245 or equivalent)
- Borosilicate cover glass (No. 1 thickness, various dimensions, Fisher Scientific)
- Glass plate (McMaster-Carr)
- Ultrasonic pen cleaner
- Bone-dry liquid CO₂ (with siphon in tank, Airgas)
- **Critical point drier (PVT-3D, Tousimis)**
- Spin coater (WS-400-NPP-Lite or equivalent, Laurell Technologies Corporation)
- Transparency mask (CAD/Art Services)
- Contact mask aligner (Karl Suss MJB3 or equivalent, Suss MicroTec)
- UV filter glass (U-360, Hoya Corporation)
- TexWipes (ITW Texwipe, cat. no. TX312)
- Aluminum weighing dish (110 mm diameter; Fisher Scientific, cat. no. 08-732-108)
- MatTek Petri dishes (35 mm, with 20 mm holes; MatTek Corporation, part no. P35-20-C-NON)
- Mini Alpha polyester swabs (ITW Texwipe; cat. no. TX754B)
- Reusable Attofluor live-cell chamber (Invitrogen, cat. no. A7816)
- UVO cleaner (Jelight, model no. 342)
- Tissue culture supplies (e.g., micropipette tips, serological pipettes, tissue culture flasks, conical tubes, Fisher Scientific)
- Petri dishes (100 or 150 mm; Fisher Scientific)
- Hemocytometer
- Biological hood
- Tissue culture incubator
- Benchtop centrifuge (Thermo Scientific CL2 model, Thermo Scientific, or equivalent)
- Compressed CO₂ gas (Airgas)
- Tissue culture microscope (TMS or equivalent, Nikon Instruments)
- Advanced fluorescence microscope (Zeiss Axiovert 200M, Zeiss; Nikon Eclipse Ti, Nikon; or equivalent)
- Cage or stagetop live-cell incubator
- Parafilm (Fisher Scientific)
- Microcentrifuge (Eppendorf 5418 model, Eppendorf or equivalent)
- Optically clear adhesive seal sheets (ThermoScientific, cat. no. AB-1170)
- PCR plates (96-well; BioExpress, cat. no. T-3085-1)
- Thermocycler (MasterCycler Gradient, Eppendorf)
- Real-time PCR System (ABI 7300, Applied Biosystems)
- ABAQUS (Simulia, Dassault Systèmes)
- 5848 MicroTester (Instron)

REAGENT SETUP

Piranha solution Combine one part 30% (vol/vol) hydrogen peroxide and three parts 95% (vol/vol) sulfuric acid in a Pyrex crystallizing dish or beaker. The solution should be prepared immediately before use.

Ethanol (70% (vol/vol)) Mix 35 ml ethanol with 15 ml deionized water. The solution can be stored indefinitely at room temperature (20 °C).

DiI stock solution (50 µg ml⁻¹) Dissolve 25 mg of DiI paste in 500 ml ethanol. Filter-sterilize the solution to remove large DiI crystals. The solution should be protected from light and can be stored indefinitely at 4 °C.

F127 Pluronic stock solution (2% (wt/vol)) Dissolve 2 g F127 Pluronic flakes in 100 ml PBS. Filter-sterilize the solution. The solution can be stored indefinitely at room temperature.

Fibronectin stock solution Dissolve lyophilized fibronectin in 1 ml of sterile, deionized water and prepare aliquots of 50–100 µl. Aliquots of solubilized fibronectin are stable for 2 weeks at –20 °C, as per the manufacturer's guidelines.

Paraformaldehyde fixation solution (3.7% (vol/vol)) Mix 30 ml deionized water, 4.5 ml 10× PBS and 10 ml 16% (vol/vol) paraformaldehyde. The solution can be stored at –20 °C for up to 1 year.

Triton X-100 permeabilization solution (0.1% (vol/vol)) Mix 100 μ l Triton X-100 and 100 ml of 1 \times PBS. The solution can be stored indefinitely at room temperature.

Goat serum solution (33% (vol/vol)) Mix 16.67 ml of goat serum and 33.33 ml of 1 \times PBS. The solution can be stored at 4 °C for up to 3 months.

BSA solution (1% (wt/vol)) Dissolve 500 mg bovine serum albumin in 50 ml of 1 \times PBS. The solution can be stored at 4 °C for up to 3 months.

Oil Red O stock solution Dissolve 300 mg of Oil Red O in 100 ml isopropanol

and mix by stirring. The solution can be stored for up to 3 months at room temperature.

Isopropanol (60% (vol/vol)) Mix 30 ml isopropanol with 20 ml deionized water. The solution can be stored indefinitely at room temperature.

Cell culture medium for hMSCs Combine 500 ml Dulbecco's modified Eagle's medium (low glucose), 50 ml FBS and 5 ml L-glutamine. The medium can be stored at 4 °C for up to 6 months.

PROCEDURE

Micropost array design and master fabrication ● **TIMING 2 weeks**

1| Design reticle photomask using AutoDesk AutoCAD or Tanner EDA L-Edit Pro. See Experimental design for suggestions on designing a photomask.

2| Obtain photomask from a company specializing in high-resolution photomasks. We use Advance Reproductions or Microtronics for chrome masks. Turnaround time is ~1 week.

■ **PAUSE POINT** The photomask can be stored in a clean and dry environment at room temperature indefinitely.

3| Immerse a 6-inch silicon wafer in Piranha solution for 10 min to remove organic residue, dump-rinse the wafer in deionized water for 10 min, and then spin-dry the wafer with a spin-rinse dryer.

! **CAUTION** The Piranha solution is potentially explosive and must be kept away from organic solvents and materials. Proper goggles, aprons and gloves must be worn during handling.

▲ **CRITICAL STEP** The operational conditions described for micropost array master fabrication strongly depend on many process parameters (e.g., temperature, gas flows, pressure and radio frequency power) and the specific equipment used. The parameters given in Steps 3–13 of the procedure serve only as guidance and should be optimized empirically for each fabrication process flow.

4| Dehydrate the wafer at 200 °C for 10 min on a contact hot plate, allow the wafer to cool down to room temperature and treat the wafer with HMDS vapor in a vacuum desiccator for 10 min.

5| Position the wafer on the chuck of the spin coater and dispense 5 ml SPR700-1.0 photoresist onto the center of the wafer. Spin the wafer at 500 r.p.m. for 8 s to spread the photoresist across the entire wafer and then quickly ramp up to 4,000 r.p.m. for 30 s to create a thin, uniform photoresist layer.

▲ **CRITICAL STEP** The wafer should be coated with photoresist within 60 min of completing the HMDS coating step.

? **TROUBLESHOOTING**

6| Soft-bake the wafer at 95 °C for 60 s on a contact hot plate to evaporate the solvent in the photoresist and densify the photoresist film.

7| Insert the reticle photomask (obtained from Step 2) and wafer into the Nikon projection stepper. Expose the wafer through the reticle photomask, with an exposure dose of ~170 mJ cm⁻². Remove the wafer from the stepper and return the photomask to storage.

8| After exposure, bake the wafer at 115 °C for 60 s on a contact hot plate to drive diffusion of photocatalyzed acid and enhance the spatial resolution of the photoresist pattern.

9| Develop the wafer with MF CD 26 developer solvent for 30 s, rinse the wafer with deionized water for 2 min and spin-dry the wafer.

10| Inspect the wafer with a metallurgical microscope to confirm that the photoresist pattern matches the desired patterns on the photomask. If not, strip away the photoresist with Piranha solution, and restart the fabrication process from Step 3.

▲ **CRITICAL STEP** Wafer inspection after photoresist development is necessary for monitoring (i) whether the correct mask has been used; (ii) the quality of the photoresist film (i.e., uniformity of thickness, absence of particulates and streaks) is satisfactory; and (iii) whether the feature dimensions are within the specified tolerances.

? **TROUBLESHOOTING**

PROTOCOL

- 11| Bake the wafer at 130 °C for 60 s on a contact hot plate to harden the photoresist.
- 12| Determine the thickness of the patterned photoresist film with the surface profilometer.
■ **PAUSE POINT** The patterned wafer can be stored in a clean and dry environment at room temperature for at least 1 week.
- 13| Etch the wafer with the DRIE machine through the exposed silicon surrounding the patterned photoresist using C_4F_8 and SF_6 plasma. The Si-DRIE-etch recipe shown below yields an etch rate of $\sim 1.47 \mu\text{m min}^{-1}$.

	Etch mode	Passivation mode
Process time	6 s	4.5 s
Overrun	0.5 s	0 s
Platen generator power	80 W	60 W
Coil generator power	600 W	600 W
Gas	SF_6 (70 sccm)	C_4F_8 (35 sccm)
Etch rate	$1.47 \mu\text{m min}^{-1}$	NA

? TROUBLESHOOTING

- 14| Inspect the wafer in the SEM to measure the post diameter and height, and to further ensure that silicon microposts with straight sidewalls are obtained (**Fig. 2a**).
- 15| Use the surface profilometer to measure the silicon etch depth and determine the uniformity of the silicon etch profile across the entire wafer.
■ **PAUSE POINT** The wafer can be stored in a clean and dry environment at room temperature indefinitely.
- 16| Dehydrate the wafer at 200 °C for 10 min on a contact hot plate, allow the wafer to cool down to room temperature and treat the wafer with HMDS vapor in a vacuum desiccator for 10 min.
- 17| Position the wafer on the center of the spin coater and dispense 10 ml of AZ4620 photoresist onto the center of the wafer. AZ4620 is a thick photoresist used to coat and protect the etched silicon structures from debris produced by the wafer die saw in Step 19. Spin the wafer at 500 r.p.m. for 30 s to spread a thick layer of photoresist across the entire wafer.
▲ **CRITICAL STEP** The wafer should be coated with photoresist within 60 min of completing the HMDS coating step.
- 18| Bake the wafer at 130 °C for 60 s on a contact hot plate to harden the photoresist.
■ **PAUSE POINT** The wafer coated with photoresist can be stored in a clean and dry environment at room temperature indefinitely.
- 19| Use the wafer die saw to cut the wafer into individual micropost array master devices. The final silicon micropost array master device has dimensions of $\sim 21 \text{ mm} \times 21 \text{ mm}$.
■ **PAUSE POINT** The silicon micropost array master coated with photoresist can be stored in a clean and dry environment at room temperature indefinitely.
- 20| Immerse and sonicate the silicon master device in acetone for 10 min to completely dissolve the photoresist, rinse the silicon master device in isopropanol for 2 min and blow-dry the silicon master device with a nitrogen gun.
■ **PAUSE POINT** The silicon micropost array master can be stored in a clean and dry environment at room temperature indefinitely.
- 21| To reinforce the master for repeated castings, cut a glass slide to $\sim 25 \text{ mm} \times 25 \text{ mm}$ and mount the master onto the glass slide with Araldite epoxy. Seal the edges of the master with SU-8 photoresist or Norland 68 optical adhesive and cure in a UV chamber.
- 22| To fluorosilanize the silicon micropost array master device, place the silicon device inside a vacuum desiccator alongside a glass slide. Dispense two or three drops of (tridecafluoro-1,1,2,2-tetrahydrooctyl)-1-trichlorosilane with a Pasteur pipette

onto the slide. Place the Pasteur pipette inside the chamber, replace the chamber cover and apply vacuum for 4 h.

! CAUTION Add the silane to the desiccator inside a properly ventilated chemical hood and connect the desiccator to a vacuum line in the fume hood if possible.

■ PAUSE POINT After silanization, the silicon micropost array master can be stored in a clean and dry environment at room temperature indefinitely.

Micropost array-negative mold production ● TIMING 2.5 h

23| Prepare a 10:1 mixture of uncured PDMS by dispensing 60 g of PDMS base prepolymer and 6 g of curing agent into a plastic cup. These amounts are sufficient for four castings of a micropost array master in small aluminum weighing dishes (as described in Step 25), but can be adjusted if more or fewer negative molds are desired. Mix the PDMS components for 3 min.

24| Degas the mixed, uncured PDMS in a vacuum desiccator for 1 h.

▲ CRITICAL STEP Use separate desiccators for silanization and degassing PDMS to ensure that there is no contamination of the PDMS with residual silane left inside the desiccator.

25| Place a silicon micropost array master in a small aluminum weighing dish and pour the degassed PDMS into the dish to a height of ~0.75 cm. Wait for 5–10 min for any bubbles introduced during pouring to rise to the surface and dissipate. Do not degas the dish of PDMS inside a vacuum desiccator.

▲ CRITICAL STEP Cast negative molds within 3 h of degassing the PDMS, as the PDMS will cure slowly at room temperature and become increasingly viscous.

26| Cure the dish of PDMS at 110 °C for 10–15 min in a laboratory oven. Remove the dish from the oven and let it cool to room temperature.

27| Carefully tear away the aluminum weighing dish without flexing the cured PDMS or master. The silicon master will not break easily if mounted onto a glass slide backing.

28| Cut away any PDMS undercutting the master with a razor blade.

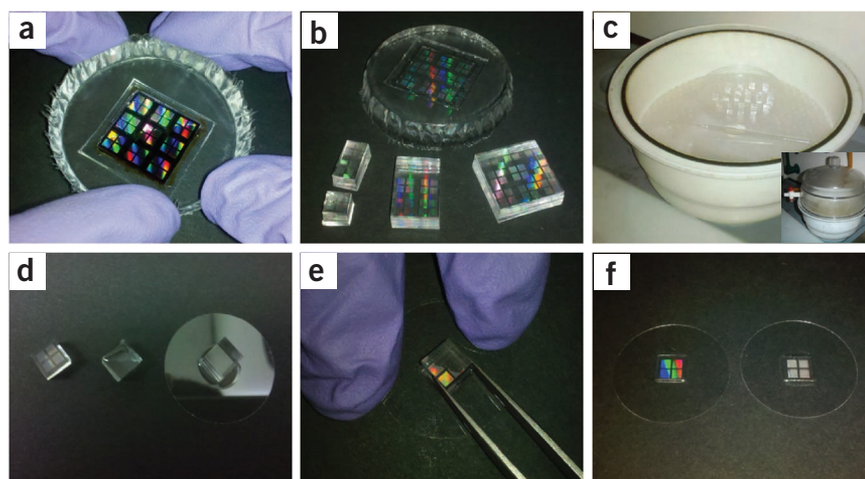
29| Place the bulk PDMS and master against a flat surface, firmly hold down the PDMS at one corner of the master and peel the PDMS at a slow constant rate from the diagonal corner to yield the negative mold (**Fig. 3a**).

30| If desired, repeat Steps 25–29 to cast additional negative molds of the master.

31| Cut away excess PDMS surrounding the micromolded region of the negative molds with a razor blade to ensure that the top surface of the mold is completely flat.

32| If desired, cut the negative molds into smaller molds so that the micropost array substrates can be cast on different-sized cover slips (**Fig. 3b**).

Figure 3 | Replica molding of a micropost array master. **(a,b)** A PDMS negative mold is peeled from the silicon master **(a)** and can be cut to different sizes **(b)** to cast arrays of different area. **(c)** The negative molds are then fluorosilanized in a vacuum desiccator. **(d)** Silanized negative molds are coated with a thin layer of uncured PDMS (middle mold), sandwiched against a cover slip (right mold) and cured. **(e)** A cured substrate is released by clamping the substrate while using tweezers to peel the mold. **(f)** Substrate quality can be quickly determined by looking at how the array diffracts light. A flawless substrate (left) diffracts light into many colors while a flawed substrate (right) has opaque regions.



PROTOCOL

33 | To activate the surface of the negative molds with plasma, place the molds face-up inside the chamber of a plasma cleaner (e.g., SPI Plasma Prep II) using a suitable carrier such as a 100 mm Petri dish. Evacuate the chamber until the pressure reaches 500–1,000 mTorr and then activate the plasma source.

34 | To fluorosilanize the negative molds, place molds inside a vacuum desiccator alongside a glass slide. Dispense two to three drops of (tridecafluoro-1,1,2,2-tetrahydrooctyl)-1-trichlorosilane with a Pasteur pipette onto the slide. Place the Pasteur pipette inside the chamber, replace the chamber cover and evacuate the chamber overnight (**Fig. 3c**).

■ **PAUSE POINT** Negative molds can be left in the desiccator indefinitely or can be removed after 1 d of silanization and stored in a clean, dry environment indefinitely.

Micropost array substrate fabrication ● **TIMING 22–24 h**

35 | Prepare a 10:1 mixture of uncured PDMS by dispensing 10 g of PDMS base prepolymer and 1 g of curing agent into a plastic cup. Vigorously mix the PDMS components for 5 min.

▲ **CRITICAL STEP** This particular PDMS mixture (as opposed to the one used for negative molds) must be exactly 10:1 in order to reproducibly cast substrates with the same elastic modulus. Weigh the PDMS prepolymer and curing agent with an analytical balance (e.g., Mettler Toledo AG245). The longer mixing time of 5 min is a precaution to ensure that the PDMS is well mixed.

36 | Degas the mixed, uncured PDMS in a vacuum desiccator for 1 h.

37 | Dispense a small amount of degassed PDMS onto the top surface of each negative mold using a plastic transfer pipette. Spread out the PDMS over the top surface by either pressing two molds together like a sandwich or gently blowing the PDMS with a stream of N₂ (**Fig. 3d**).

38 | Place the PDMS-covered molds in a vacuum desiccator for 15 min to remove any air bubbles that may be trapped between the uncured PDMS and molds.

39 | Clean cover slips in preparation for casting against the molds. Blow-dry the cover slips with a stream of N₂ and plasma clean for 2 min at 100 mA. Operate the plasma cleaner as described in Step 33. If plasma cleaner is unavailable, cover slips can be cleaned with detergent or Piranha solution.

40 | Lay out the clean cover slips on a clean glass plate and remove PDMS-covered molds from the vacuum desiccator.

41 | With a pair of tweezers, flip each mold onto a cover slip, sandwiching the uncured PDMS between the mold and the glass. Apply firm, uniform pressure to squeeze out the excess PDMS and ensure that the resultant casting is as uniformly level as possible (**Fig. 3d**).

42 | Cure the molds and cover slips at 110 °C for 20 h in a laboratory oven. Remove the substrates and molds from the oven and let them to cool to room temperature.

▲ **CRITICAL STEP** PDMS is fully cured after 20 h at 110 °C and will not cure further when stored at room temperature. However it will continue to cure if left at 110 °C for longer periods of time. For example, PDMS cured for 48 h will be more cross-linked, and therefore stiffer, than PDMS cured for 20 h. Therefore, it is recommended that the curing time be kept consistent in repeat castings.

43 | Peel off each negative mold from its cover slip to yield the micropost substrate. To minimize the likelihood of breaking the glass, use two fingers to clamp down the cover slip against a flat surface while the other hand grips the mold with tweezers. Peel the mold at a slow, constant rate from one corner toward the other until the substrate is released (**Fig. 3e**).

? **TROUBLESHOOTING**

44 | Inspect each substrate by tilting it back and forth under a light (**Fig. 3f**). If a substrate appears opaque, the micropost array is collapsed. If the substrate diffracts light into different colors, the majority of microposts are likely upright. Substrates that are macroscopically flawless should be examined further with an optical microscope to check whether there are collapsed or missing microposts scattered within the array. Missing posts, which leave behind stumps, appear hollow when viewed under an optical microscope with ambient illumination. The short stumps are also invisible when the substrate is immersed in liquid, which is another method to distinguish them from intact microposts. Transfer flawless substrates to a storage container (e.g., Petri dish). Flawed substrates can be returned to flawless state by sonicating the substrates in

BOX 1 | SUPERCRITICAL DRYING OF COLLAPSED MICROPOST SUBSTRATES

● TIMING 1 H

Operating instructions for critical point drying is machine dependent. The following instructions are for generic operation.

1. Cool the drying chamber to below 0 °C by opening the cool meter valve that connects the machine to a liquid CO₂ tank equipped with a siphon.
2. Fill the chamber with ethanol and transfer the substrates to the chamber. Substrates may be stacked as long as they do not stick to each other. As such, the PDMS squeezed out during casting that surrounds the array region is a suitable spacer.
3. Close the chamber and open the Fill meter valve to allow liquid CO₂ to enter the chamber. The presence of Schlieren lines indicates that liquid CO₂ and ethanol are mixing.
4. When the chamber is full, open the Purge meter valve to allow the chamber contents to exit. The fill and purge rates are not exact, but the position of the Purge meter valve should always be lower (i.e., more closed), than the position of the Fill meter valve. A mixture of ethanol and CO₂ will exit the tubing connected to the purge outlet.
5. Wait for a sufficient amount of time until the Schlieren lines disappear and only dry ice is observed exiting the purge tube. This indicates that all the ethanol has been replaced by liquid CO₂.
6. Close all valves and heat up the chamber until the pressure and temperature exceed the critical point of CO₂ (1,037 psi and 31 °C). Above this point, liquid CO₂ is converted to gaseous CO₂.
7. Open the purge valve to vent the CO₂ from the chamber.
8. Remove the substrates from the critical point drier and inspect as described in Step 44 of the PROCEDURE.

ethanol for 1–2 min to make the microposts stand upright and then supercritically drying the substrates in a critical point drier (see **Box 1**).

■ **PAUSE POINT** Flawless substrates can be stored in a dry environment indefinitely. Flawed substrates that are stored in ethanol do not have to be supercritically dried immediately.

Micropost array substrate functionalization ● TIMING 4 h

45| Cast micropatterned or flat slabs of PDMS to use as stamps. If micropatterned stamps are desired, see **Box 2** for steps to make a micropatterned silicon wafer before performing any steps in this section. Otherwise, prepare sufficient 30:1 or 10:1 uncured PDMS in a plastic cup to fill a 100-mm or 150-mm Petri dish to a height of 0.75 cm. The 30:1 PDMS is softer than 10:1 and is recommended for flat stamps. The 10:1 PDMS is recommended for micropatterned stamps, as the increased stiffness reduces the likelihood that the stamp will collapse under its own weight. Mix the PDMS components for 3 min.

46| Degas the mixed, uncured PDMS in a vacuum desiccator for 1 h.

47| For a flat slab, pour the degassed PDMS into a 150-mm Petri dish. For a micropatterned slab, pour PDMS over a micropatterned silicon wafer placed inside a Petri dish or aluminum weighing dish of suitable size. Wait for 5–10 min for the bubbles introduced during pouring to rise to the surface and dissipate.

48| Cure the PDMS at 65 °C for at least 2 h or at 110 °C for at least 15 min (if using an aluminum weighing dish). Remove the dish of PDMS from the oven and allow it to cool for 5 min. Proceed to Step 52 if you are not casting a micropatterned PDMS slab.

■ **PAUSE POINT** Curing time for the slab of PDMS is not critical; accordingly, the dish can be left in the oven for a couple of days.

49| If you are casting a PDMS slab against a micropatterned wafer, carefully separate the PDMS and wafer from the dish without flexing the silicon wafer.

50| Cut away any PDMS that has undercut the wafer with a razor blade.

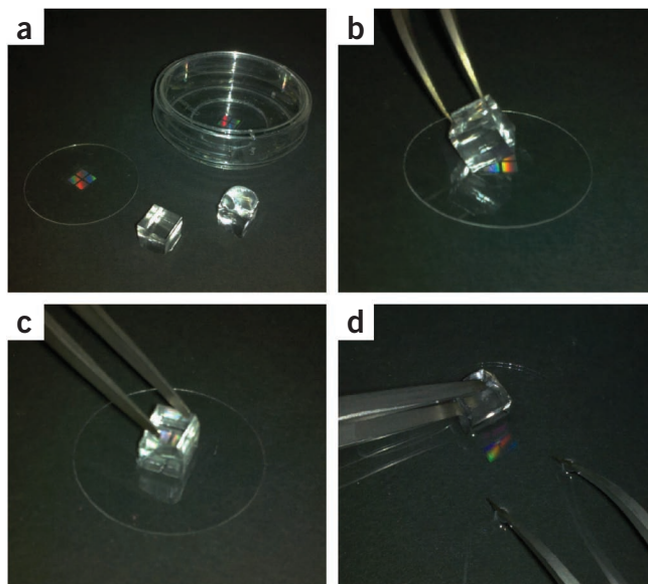
51| Place the bulk PDMS and wafer against a flat surface, firmly hold down the PDMS at one end of the wafer and peel the micropatterned PDMS slab at a slow constant rate from the other end.

52| Use a razor blade to cut the slab of PDMS into stamps that fit the area of the micropost arrays to be functionalized (**Fig. 4a**).

53| Notch the bottom corner of each stamp to indicate the orientation of the stamping surface.

PROTOCOL

Figure 4 | Microcontact printing on micropost array substrates. (a) PDMS stamps are cut to the size of the micropost array and then coated with fibronectin solution (right stamp). Micropost array substrates can be left unmounted (left substrate) or mounted (right substrate) onto a Petri dish. (b) A dried, coated stamp is aligned and laid on top of the micropost array. (c) Gentle pressure is applied with tweezers to ensure good contact between the stamp and the micropost tips. (d) The substrate and stamp are placed in ethanol before the stamp is peeled, to minimize any damage that peeling forces may impart.



54 | Prepare flawless micropost substrates (from Step 44) for functionalization. Cut away any excess PDMS surrounding the array with a razor blade (**Fig. 4a**). If substrates are to be used for live-cell imaging at high magnification, proceed to Step 55. Otherwise, skip to Step 56.

55 | For live-cell imaging with high-magnification, low-working-distance objectives (**Fig. 4a**), micropost substrates can be mounted onto a Petri dish with a 20-mm hole cut out (e.g., MatTek dish) using UV-curable adhesive (option A) or PDMS (option B); alternatively, they can be mounted onto a reusable stainless steel dish (option C):

(A) Mount substrate to a MatTek Petri dish with a cut-out hole using UV-curable adhesive

- (i) Use an Alpha polyester swab to spread a thin, narrow layer of Norland 68 optical adhesive on the back of the Petri dish, around the perimeter of the cut-out hole.
- (ii) Lay down the micropost substrate over the hole.
- (iii) Expose the entire assembly in a UV curing chamber to cure the adhesive.

(B) Mount substrate to a MatTek Petri dish with a cut-out hole using PDMS

- (i) Use an Alpha polyester swab to spread a thin, narrow layer of uncured 10:1 PDMS on the back of the Petri dish, around the perimeter of the cut-out hole.
- (ii) Lay down the micropost substrate over the hole.
- (iii) Cure the entire assembly at 65 °C for 1 h in laboratory oven.

(C) Mount substrate onto a reusable stainless steel dish

- (i) Insert substrate in a reusable stainless steel dish (e.g., Attofluor).

■ PAUSE POINT Steps 49–55 are not time sensitive and can be performed anytime before the rest of this section. PDMS stamps and trimmed/assembled substrates can be stored in a clean and dry environment indefinitely until the day of use.

56 | Sonicate stamps (from Step 53) in ethanol for 5 min to clean off any particulates.

57 | Dry the stamps in a sterile environment with a stream of N₂ and then transfer them to a new Petri dish.

58 | Prepare a sufficient working volume of fibronectin solution, diluted to a final concentration of 50 µg ml⁻¹ in sterile, deionized water, to coat the stamps.

▲ CRITICAL STEP Do not mix the fibronectin solution with a small-bore micropipette tip, which will shear the protein. Instead, use a large-bore micropipette tip or flick the tube to mix.

59 | Dispense droplets of fibronectin solution onto the top surface of each stamp. Sweep over the droplet with a micropipette tip to spread the solution along the edges and corners of the stamp (**Fig. 4a**).

60 | Let fibronectin adsorb onto the stamp for 1 h.

61 | Wash away excess fibronectin solution by flooding the Petri dish with deionized water until the stamps are submerged.

62 | Submerge the stamps in a second Petri dish filled with deionized water to wash again.

63 | Dry the stamps with a stream of N₂ and transfer stamps to a new Petri dish.

64 | Expose the micropost substrates to UV ozone for 7 min in a UVO cleaner to temporarily hydrophilize the surface.

BOX 2 | STAMP MASTER FABRICATION FOR MICROPATTERNED FUNCTIONALIZATION ● TIMING 2 WEEKS

▲ **CRITICAL** Many process parameters in this section, such as photoresist spin-coating speeds, baking times and UV exposure duration are machine specific and dependent on the experimental design. The parameters given are for guidance only and should be optimized empirically for each type of application.

▲ **CRITICAL** Steps 3–8 should be performed in a clean room (class 10,000 or equivalent). Equipment and reagents should be handled inside the clean environment for these steps. Silicon wafers should not be exposed to an unclean environment until after Step 8.

1. Design photomask using vector graphics software such as Adobe Illustrator. See Experimental design for suggestions on designing a photomask.
2. Obtain transparency photomask from a printing company specializing in high-resolution printing. We use CAD/Art Services for this service. Turnaround time is ~1 week.
3. Dehydrate a new silicon wafer at 200 °C for 10 min on a contact hot plate, remove the wafer from the hot plate and let it cool to room temperature.
4. If a plasma cleaner is available, use it to clean the wafer for 2 min at 100 mA to remove surface contaminants; place wafer inside the chamber of a plasma cleaner and operate, as described in Step 33 of the PROCEDURE.
5. Position the wafer on the chuck of the spin coater and dispense 5 ml of SU-8 2000 series photoresist on the center of the wafer (see Experimental design for suggestions on choosing the right photoresist). Spin the wafer at 500 r.p.m. (acceleration 84 r.p.m. per s) for 10 s to spread out the photoresist and then ramp up to 1,500 r.p.m. (acceleration 336 r.p.m. per s) for 30 s to produce a uniform photoresist layer.

? TROUBLESHOOTING

6. Soft-bake the wafer at 65 °C for 1 min and then at 95 °C for 2 min on two contact hot plates to evaporate the solvent and densify the photoresist film. Remove the wafer from the hot plate and let it cool to room temperature.

? TROUBLESHOOTING

7. Expose the wafer to 365 nm UV light through the photomask with a contact mask aligner. The exposure dose is 130 mJ cm⁻², although this must be optimized based on the experimental design.

8. After exposure, bake the wafer at 65 °C for 1 min followed by at 95 °C for 2 min on two contact hot plates to cross-link exposed regions of photoresist. Let the wafer cool slowly to room temperature on the hot plate.

9. Develop the photoresist by submerging and agitating the wafer in two successive dishes of propylene glycol monomethyl ether acetate (PGMEA) for 3 min and 10 s.

10. Rinse away residual PGMEA by submerging and agitating the wafer in two successive dishes of isopropanol for 10 s each. Blow-dry the wafer with a stream of N₂.

11. Inspect the wafer on a metallurgical microscope for pattern fidelity of photoresist.

? TROUBLESHOOTING

12. Hard-bake the wafer on a contact hot plate at 175 °C for at least 15 min. Let the wafer cool to room temperature.

? TROUBLESHOOTING

13. Fluorosilane the wafer in the same manner described in Step 22 of the PROCEDURE.

65| Lay fibronectin-coated stamps on top of the substrates and apply gentle pressure to transfer fibronectin to the tips of the microposts (**Fig. 4b,c**). Let the stamp remain in contact for at least 15 s.

▲ **CRITICAL STEP** Microcontact-print the fibronectin within 30 min of exposing the substrates to UV ozone. Otherwise the substrates will lose hydrophilicity and will have to be re-exposed to UV ozone for 7 min.

66| Peel away the stamps. For taller, softer microposts, submerge the substrates in ethanol before peeling away the stamps (**Fig. 4d**).

? TROUBLESHOOTING

67| If they are not already submerged in ethanol, submerge the functionalized substrates in pure ethanol, followed by 70% (vol/vol) ethanol and three successive washes with deionized water to dilute out the ethanol. There are two methods of washing, depending on whether a substrate is mounted onto a live-cell viewing dish (option A) or is unmounted (option B).

(A) Washing dish-mounted substrates

- (i) Add 2 ml of ethanol to the dish, swirl the dish and then aspirate.
- (ii) Add 2 ml of 70% (vol/vol) ethanol, swirl the dish and then aspirate.
- (iii) Add 2 ml of deionized water, swirl the dish and then aspirate. Repeat this two more times and leave the substrates covered with water.

PROTOCOL

(B) Washing unmounted substrates

- (i) Transfer the substrates to a Petri dish containing ethanol.
- (ii) Transfer the substrates to a Petri dish containing 70% (vol/vol) ethanol and swirl the dish.
- (iii) Transfer the substrates to a Petri dish containing deionized water and swirl the dish. Repeat this step twice with another two Petri dishes of deionized water.

▲ **CRITICAL STEP** Pure ethanol, which has a low surface tension, is used to wet the micropost array substrates without collapsing the microposts. Ethanol (70% (vol/vol)) serves to sterilize the substrates before cell culture. Minimize exposure of the substrates to air during successive washes in water to prevent the microposts from collapsing. For dish-mounted substrates, do not aspirate the liquid completely; leave the array region covered with a puddle of liquid. For unmounted substrates, use tweezers to transfer substrates quickly between Petri dishes and then forcibly submerge them in aqueous liquids (as PDMS is very hydrophobic). Use this technique for all subsequent liquid handling steps in the protocol (Steps 69, 70, 72, 73, 75 and 79).

68| If the microposts are to be fluorescently labeled, prepare a working solution of 5 $\mu\text{g ml}^{-1}$ DiI by diluting 50 $\mu\text{g ml}^{-1}$ DiI stock solution in deionized water. Otherwise, proceed to Step 71.

69| Submerge substrates in DiI solution and incubate for 1 h, protected from light, using option A for dish-mounted substrates or option B for unmounted substrates:

(A) Dish-mounted substrates

- (i) Aspirate deionized water and add 2 ml DiI solution.

(B) Unmounted substrates

- (i) Transfer the substrates to a Petri dish filled with DiI solution.

70| Wash away the excess DiI solution with deionized water, using option A for dish-mounted substrates or option B for unmounted substrates:

(A) Dish-mounted substrates

- (i) Aspirate the DiI solution in the dish.
- (ii) Add 2 ml of deionized water, swirl the dish and then aspirate. Repeat two more times and leave the substrates covered with water.

(B) Unmounted substrates

- (i) Transfer the substrates to a Petri dish containing deionized water and swirl the dish. Repeat this step two more times.

71| To block un-stamped regions of the substrates with F127 Pluronic, prepare a working solution of 0.2% (wt/vol) Pluronic by diluting the 2% (wt/vol) Pluronic stock solution in PBS.

72| Submerge substrates in Pluronic solution and incubate for 30–60 min, protected from light (if labeled with DiI), using option A for dish-mounted substrates or option B for unmounted substrates:

(A) Dish-mounted substrates

- (i) Aspirate the deionized water and add 1–2 ml Pluronic solution.

(B) Unmounted substrates

- (i) Transfer the substrates to a Petri dish filled with Pluronic solution.

73| Wash away the excess Pluronic solution with deionized water and PBS, using option A for dish-mounted substrates or option B for unmounted substrates:

(A) Dish-mounted substrates

- (i) Aspirate the Pluronic solution in the dish.
- (ii) Add 2 ml of deionized water, swirl the dish and then aspirate. Repeat this step once.
- (iii) Add 2 ml of PBS.

(B) Unmounted substrates

- (i) Transfer the substrates to a Petri dish containing deionized water and swirl the dish. Repeat this step once.
- (ii) Transfer the substrates to a Petri dish containing PBS.

■ **PAUSE POINT** At this point, the substrates can be used immediately for an experiment or stored at 4 °C for up to 1 week. For storage, seal the container holding the substrates with Parafilm. If the substrates are fluorescently labeled, wrap the container with aluminum foil.

Cell culture on micropost array substrates ● TIMING 1–24 h

74| Warm cell culture medium and trypsin to 37 °C before seeding cells on micropost substrates.

75| Replace PBS covering substrates with cell culture medium, using option A for dish-mounted substrates or option B for unmounted substrates:

(A) Dish-mounted substrates

(i) Aspirate PBS and add 2 ml of medium.

(B) Unmounted substrates

(i) Transfer the substrates to a Petri dish filled with medium.

76| Re-suspend adherent cells by rinsing the cells with PBS twice and then adding 2 ml of trypsin-EDTA into the T-75 culture flask. Incubate the cells at 37 °C/5% CO₂ for 5 min, tap the side of the flask to detach the cells, and then add 4 ml of cell culture medium to neutralize trypsin. Pipette the cell suspension into a 15-ml conical tube and centrifuge at 300g for 5 min to pellet the cells. Aspirate the medium, and resuspend the cell pellet in 5 ml of cell culture. Detailed tissue culture procedures have also been published elsewhere¹⁰¹.

77| To count resuspended cells, pipette 20 µl of cell suspension into the notch of a hemocytometer. Observe the hemocytometer under a tissue culture microscope and count the number of cells within the nine large grids on the hemocytometer. Divide the cell count by the number of grids inspected and multiply by 10,000 to determine the number of cells per ml of suspension.

78| Replate cells onto the substrates by adding the cells to a final density of 1 to 5 × 10³ cells per cm² of culture area, including the area in the culture vessel surrounding the micropost arrays. For a substrate in a 35-mm Petri dish, 1 to 5 × 10⁴ cells is an appropriate number of cells for seeding. Place the substrates in an incubator to allow cells to attach.

▲ CRITICAL STEP The density of cells on the micropost array substrates must be empirically optimized, as dictated by the goals of the experiment. Parameters to optimize include initial cell seeding density (Step 78), cell seeding time (Step 79) and incubation time for allowing cells to spread before experimentation (Step 80). These are dependent on cell type and whether micropatterns are used. See Experimental design for general seeding strategies.

? TROUBLESHOOTING

79| After allowing the cells to attach for an empirically optimized time, remove floating cells by repeatedly washing the substrates, using option A for dish-mounted substrates or option B for unmounted substrates:

(A) Dish-mounted substrates

(i) Aspirate the liquid in the dish. Do not aspirate directly above the area where cells are attached.

(ii) Add 2 ml PBS, swirl the dish to disperse floating cells that have pooled and aspirate. Repeat this step (usually three to seven times) until the majority of floating cells have been removed.

(iii) Aspirate the PBS and replace with fresh medium.

(B) Unmounted substrates

(i) Transfer the substrates to a Petri dish filled with PBS and gently swirl the dish to wash away floating cells that have pooled near the surface of the substrate. Repeat this step one or two times.

(ii) Transfer the substrates onto a Petri dish with fresh medium.

? TROUBLESHOOTING

80| Allow substrates seeded with cells to incubate at 37 °C until ready to perform experiment. Cells can be fully spread on the micropost arrays in as little as 2 h. For long-term experiments, cells have been cultured on the micropost arrays for up to 2 weeks and could possibly remain viable for even longer periods. For traction force analysis, cells can be imaged live or fixed. If cells will be imaged live, proceed to Step 81. If cells will be fixed and stained for cellular structures, proceed to **Box 3**, which describes the protocol for fixation and staining, and then proceed to Step 81. For stem cell differentiation analysis, there are three separate assays described in this protocol that can be carried out at this point in the procedure instead of proceeding to Step 81. **Box 4** describes the quantification of stem cell differentiation markers by quantitative PCR (qPCR); **Box 5** describes the assay for adipogenic differentiation by Oil Red O staining; and **Box 6** describes the assay for osteogenic differentiation by ALP staining.

? TROUBLESHOOTING

BOX 3 | FIXATION AND IMMUNOFLOUORESCENCE STAINING OF CELLS ON MICROPOST ARRAY SUBSTRATES ● TIMING 7–8 H

This fixation protocol can be optimized depending on the experimental design. The standard method is to fix the cells and then permeabilize. In certain cases, such as staining for focal adhesions, a Triton extraction before fixation may yield better signal by removing more cytoplasmic background.

1. If a dish-mounted substrate is to be fixed and stained, separate the substrate from the dish using option A for substrate mounted with Norland 68 optical adhesive, option B for substrate mounted with PDMS or option C for substrate inserted in a reusable dish:

(A) Substrate mounted with Norland 68 optical adhesive

(i) Cut out the substrate with a diamond-tipped pen.

(B) Substrate mounted with PDMS

(i) Pry away the substrate with a razor blade. The Petri dish is reusable.

(C) Substrate inserted in a reusable dish

(i) Remove the substrate from the dish.

▲ **CRITICAL STEP** Make sure the cultured surface remains wet.

2. Fix cells by immersing substrates in 3.7% (vol/vol) paraformaldehyde for 20 min.

▲ **CRITICAL STEP** Protect the substrates from light if labeled with DiI.

3. Wash the substrates in three successive Petri dishes of PBS.

4. Permeabilize the cells by immersing the substrates in 0.1% (vol/vol) Triton X-100 solution for 10 min.

▲ **CRITICAL STEP** Protect the substrates from light if labeled with DiI.

5. Block the substrates against nonspecific adsorption of antibodies by immersing them in 2 ml of 33% (vol/vol) goat serum or 1% (wt/vol) BSA solution for 1 h.

▲ **CRITICAL STEP** For blocking, it is recommended that normal serum from the host species of the secondary antibodies be used, which is most commonly goat. BSA can be used in other cases. Protect the substrates from light if labeled with DiI.

6. Prepare an appropriate dilution of primary antibodies in 33% (vol/vol) goat serum or 0.1% (vol/vol) Triton X-100 solution. Typical dilutions range from 1:100 to 1:500. Circular (25 mm) and 22-mm square cover slips require 150–200 µl each, whereas 18-mm circular cover slips require 75–100 µl of antibody solution.

7. Dispense droplets of primary antibody solution on a sheet of Parafilm for each substrate to be stained.

8. Place each substrate face-down on a droplet and use tweezers to gently rock the substrate up and down a few times to evenly distribute the solution. Let the substrates incubate with the primary antibody for 1 h.

▲ **CRITICAL STEP** Protect the substrates from light if labeled with DiI.

9. Wash the substrates with PBS three times to remove the unbound primary antibody. Each PBS wash should last for 3–5 min.

10. Prepare an appropriate dilution of secondary antibodies and other fluorescent stains (e.g., DAPI, fluorophore-conjugated phalloidin) in 33% (vol/vol) goat serum or 0.1% (vol/vol) Triton X-100 solution. Typical dilutions range from 1:100 to 1:2,000. The same volumes used for the primary antibody solution apply for the secondary antibody solution.

11. Dispense droplets of secondary antibody solution on a sheet of Parafilm for each substrate to be stained.

12. Place each substrate face-down on a droplet and gently rock up and down a few times. Let the substrates incubate with the secondary antibody for 1 h. Protect from light.

13. Wash the substrates with PBS three times to remove the unbound secondary antibody. Each PBS wash should last for 3–5 min.

14. Transfer each stained substrate face-up to a microscope slide, dispense 1–2 drops of Fluoromount G and cover with a clean cover slip.

15. Allow Fluoromount G to cure for ~4 h before imaging. Proceed to Step 81 of the PROCEDURE for imaging the fixed and stained cells.

▲ **CRITICAL STEP** The highest quality images are obtained within 24 h of mounting. After 24 h, Fluoromount G is fully cured and its index of refraction will match that of borosilicate glass. As a result, the microposts will be very hard to see under phase illumination and must be located with fluorescence.

Imaging cells on micropost array substrates ● TIMING variable

81| If performing live-cell imaging, set up the microscope and equilibrate the temperature and CO₂ in the live-cell incubator.

82| Place a dish-mounted substrate or a slide-mounted fixed substrate on the microscope stage. Add objective immersion oil if necessary.

83| Adjust the focus to bring the microposts into view.

84| Scan the micropost array for cells to image (**Fig. 5a–c**). This is easiest if live cells are labeled with GFP or if fixed cells are stained for cellular structures. In that event, use appropriate fluorescence excitation to detect the labeled cells. If cells are not fluorescent, but the microposts are labeled with DiI, it is more convenient to use fluorescence rather than phase illumination to identify cells. Contractile cells can be identified by looking for well-deflected microposts that can be

BOX 4 | QUANTIFICATION OF GENE EXPRESSION IN STEM CELLS ON MICROPOST ARRAY SUBSTRATES ● TIMING 3 H

▲ **CRITICAL** The general protocol described here is as described in reference 60. Users can of course switch to their familiar protocols for RNA extraction as well as for qPCR. Detailed instructions are provided separately by the kits/qPCR machines that are used for the protocols.

▲ **CRITICAL** For this assay, cells should be cultured on an array area of 2.25 cm² at a density of 5,000 cells per cm² to obtain enough RNA for quantification. Let the cells attach and fully spread overnight. Substrates should be unmounted and immersed in a multi-well or Petri dish format.

1. Treat cells on the micropost array substrates with different chemicals; in this case, treat the cells with 2 ml of osteogenic medium, adipogenic medium or mixed medium (osteogenic:adipogenic = 1:1) for 14 d. Untreated cells left in growth media are used as a control.

2. After treatment, aspirate medium and rinse the cells with PBS twice. Keep cells in PBS.

3. Under a microscope, scrape away cells that are not on the micropost array part of the substrate using a small-bore micropipette tip.

4. Aspirate PBS and discard the scraped cells.

5. To extract RNA from cells attached to the micropost array substrate using the Qiagen RNeasy Micro kit or an equivalent extraction kit, first add 10 µl of 2-mercaptoethanol per 1 ml Buffer RLT (included in the Qiagen kit) and mix by vortexing.

6. Disrupt the cells by adding 350 µl Buffer RLT directly on the micropost array substrate.

7. Harvest lysate with a cell lifter. Pipette the lysate directly into a RNase-free tube. Pass the lysate at least five times through a 20-gauge needle fitted to an RNase-free syringe.

■ **PAUSE POINT** Samples can be preserved in a -80 °C deep freezer at this point.

8. Add 1 volume (usually 350 µl) of 70% ethanol to the homogenized lysate, and mix well by pipetting. Do not centrifuge.

9. Transfer the sample, including any precipitate that may have formed, to an RNeasy MinElute Spin Column in a 2-ml collection tube. Close the tube gently, and centrifuge for 15 s at ≥8,000g (≥10,000 r.p.m.). Discard the flow-through.

10. Add 350 µl Buffer RW1 to the RNeasy MinElute Spin Column. Close the tube gently and centrifuge for 15 s at ≥8,000g (≥10,000 r.p.m.) to wash the column. Discard the flow-through.

11. Add 10 µl DNase I stock solution to 70 µl Buffer RDD (included in the Qiagen kit). Mix by gently inverting the tube.

12. Pipette the DNase I incubation mix (80 µl) directly onto the RNeasy MinElute silica gel membrane, and place on the benchtop at room temperature for 15 min.

13. Add 350 µl Buffer RW1 (included in the Qiagen kit) into the RNeasy MinElute Spin Column, and centrifuge for 15 s at ≥8,000g. Discard the flow-through and collection tube.

14. Transfer the RNeasy MinElute Spin Column into a new 2-ml collection tube. Pipette 500 µl Buffer RPE (included in the Qiagen kit) onto the RNeasy MinElute Spin Column. Close the tube gently and centrifuge for 15 s at ≥8,000g (≥10,000 r.p.m.) to wash the column. Discard the flow-through.

15. Add 500 µl of 80% (vol/vol) ethanol to the RNeasy MinElute Spin Column. Close the tube gently and centrifuge for 2 min at ≥8,000g (≥10,000 r.p.m.) to dry the silica gel membrane. Discard the flow-through and collection tube.

16. Transfer the RNeasy MinElute Spin Column into a new 2-ml collection tube. Open the cap of the spin column, and centrifuge in a microcentrifuge at maximum speed for 5 min. Discard the flow-through and collection tube.

17. To elute, transfer the spin column to a new RNase-free 1.5-ml collection tube. Pipette 14 µl RNase-free water directly onto the center of the silica gel membrane. Close the tube gently and centrifuge for 1 min at maximum speed to elute.

18. Dissolve 1 µl RNA in 99 µl nuclease-free water and quantify RNA concentration using a spectrophotometer. The RNA concentration is calculated as follows: $(OD_{260} \text{ value} \times 40 \times \text{dilution factor}) / 1,000 = x \mu\text{g } \mu\text{l}^{-1}$

19. Take 0.5 µg of RNA and mix with 1 µl Oligo dT₁₂₋₁₈ (500 µg ml⁻¹), 1 µl 10 mM dNTP mixture and sterile water to a final volume of 12 µl.

20. Heat the mixture to 65 °C for 5 min and quickly chill on ice. Collect the contents of the tube by brief centrifugation and add 4 µl of 5× first-strand buffer, 2 µl of 0.1 M DTT and 1 µl RNaseOUT Recombinant Ribonuclease Inhibitor (40 U µl⁻¹).

21. Mix the contents of the tube gently and incubate at 37 °C for 2 min.

22. Add 1 µl (200 units) of MMLV reverse transcriptase, and mix by pipetting gently up and down. Incubate for 50 min at 37 °C. Then inactivate the reaction by heating at 70 °C for 15 min.

23. Add sterile water to a final volume of 100 µl.

■ **PAUSE POINT** Samples can be stored at -20 °C for up to 1 year at this point.

24. For qPCR, pipette 5 µl of each sample to one well of a 96-well PCR plate. Triplicate each sample.

▲ **CRITICAL STEP** Make sure that each sample is loaded at the same volume. Change the pipette tip after loading onto each well.

25. Make PCR master mixture by mixing ten parts of 2× universal PCR mixture, one part of TaqMan gene expression assay and four parts of nuclease-free water. Add 15 µl to each well in the 96-well plate. Seal the 96-well plate with optically clear adhesive seal sheets.

26. Load 96-well PCR plate in qPCR thermal cycler (e.g., ABI 7300) and perform the experiment according to the protocol provided by the vendor.

BOX 5 | OIL RED O STAINING FOR ADIPOGENIC DIFFERENTIATION OF STEM CELLS ON MICROPOST ARRAY SUBSTRATES ● TIMING 2 H

1. Treat cells on the micropost array substrates by adding 2 ml adipogenic medium for 14 d. Change medium twice a week.
2. After treatment, rinse cells with PBS twice and then fix cells with 10% (vol/vol) formalin for 30 min.
3. During fixation, prepare Oil Red O working solution by mixing three parts of Oil Red O stock solution with two parts of deionized water. For example, for a 10-ml working solution, mix 6 ml of Oil Red O stock solution with 4 ml of water. Allow solution to sit at room temperature for 10 min and then filter it through glass fiber filter paper.
▲ CRITICAL STEP The Oil Red O working solution must be freshly prepared and is stable for only 2 h at room temperature. Filtering through glass fiber filter paper removes undissolved Oil Red O particles, which may jeopardize cell counting.
4. After fixation, aspirate the fixing solution and rinse cells with PBS three times to wash out the formalin.
5. Aspirate the PBS and add 2 ml of 60% (vol/vol) isopropanol to cover each substrate. Incubate for 5 min.
6. Aspirate 60% (vol/vol) isopropanol and add 2 ml of Oil Red O working solution to each substrate. Incubate at room temperature for 10 min.
▲ CRITICAL STEP The 60% (vol/vol) isopropanol pretreatment in Step 5 eliminates the background of Oil Red O binding. Do not rinse the cells with PBS in between 60% (vol/vol) isopropanol treatment and Oil Red O staining.
7. Aspirate the Oil Red O working solution and add 2 ml of 60% (vol/vol) isopropanol to rinse away excess Oil Red O. Then rinse the cells with PBS three times.
8. Stain nuclei with DAPI at a dilution of 1:2,000 in 2 ml PBS. Incubate substrates at room temperature for 30 min and protect from light.
9. Count cells under microscope. The percentage of Oil Red O-positive stained cells is calculated as follows: (number of Oil Red O-positive cells/number of DAPI-stained nuclei) × 100% (see **Fig. 8**).

BOX 6 | ALKALINE PHOSPHATASE STAINING FOR OSTEOGENIC DIFFERENTIATION OF STEM CELLS ON MICROPOST ARRAY SUBSTRATES ● TIMING 2 H

- ▲ CRITICAL STEP** Given the fact that the number of cells on the micropost array substrate may be low for this assay, the quantification assay of alkaline phosphatase (ALP) activity may not be feasible. Instead, it is better to count the number of ALP activity-positive cells directly and then normalize with the total number of cells stained by DAPI in the same field.
1. Treat cells on the micropost array substrates with osteogenic induction medium for 14 d. Change the medium twice a week.
 2. After treatment, rinse cells with PBS three times.
 3. Prepare citrate working solution by diluting 2 ml of concentrated citrate into 98 ml of deionized water. The citrate working solution can be kept at 4 °C for further experiments.
 4. Prepare the fixation solution by mixing two parts of citrate working solution with three parts of acetone. For example, to prepare 10 ml of fixation solution, mix 4 ml of citrate working solution with 6 ml of acetone.
▲ CRITICAL STEP Fresh fixation solution must be prepared immediately before use.
 5. Aspirate PBS and add 2 ml of fixation solution to each substrate and fix cells at room temperature for 30 s.
 6. Aspirate fixation solution, rinse cells with PBS twice and keep cells in PBS.
 7. Prepare substrate solution as follows: carefully cut a fast blue RR salt capsule (keep under –20 °C), dissolve the fast blue RR salt powder in 48 ml H₂O and mix by vortex until all the fast blue salt has dissolved. The color of the solution will become yellow. Add 2 ml of naphthol AS-MX phosphate alkaline solution into the fast blue RR solution and mix by inverting the tube five times.
▲ CRITICAL STEP Substrate solution must be prepared immediately before use. Once naphthol AS-MX phosphate alkaline solution is added to the fast blue solution, the solution must be used within 30 min.
 8. Aspirate PBS from cells and add 2 ml of substrate solution to each substrate. Incubate at room temperature for 30 min and protect from light.
 9. Aspirate the substrate solution and rinse cells with PBS once.
 10. Stain nuclei with DAPI at a dilution of 1:2,000 in 2 ml PBS. Incubate substrates at room temperature for 30 min and protect from light.
 11. Rinse cells with PBS twice. The cell body of ALP activity-positive cells will become blue after staining.
 12. Count cells under microscope. The percentage of ALP activity-positive cells is calculated as follows: (number of ALP (+) cells/number of DAPI-stained nuclei) × 100% (see **Fig. 8**).

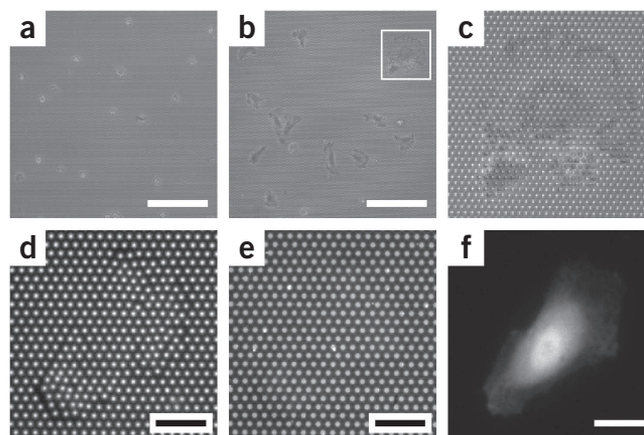
connected to form a closed perimeter. Weaker cells can be located by adjusting the fine focus up and down over the length of the microposts to find posts that have slight deflections.

? TROUBLESHOOTING

85| For a target cell, use fluorescence illumination and adjust the fine focus to the tips of the microposts. Acquire an image of the tips of the microposts, known as the ‘top image’ (**Fig. 5d**).



Figure 5 | Basic imaging of cells on micropost array substrates. (a) A low-magnification, bright-field image of round, but adherent, cells after only 30 min of spreading on the micropost array. Scale bar, 150 μm . (b,c) A low-magnification, bright-field image of cells after overnight incubation on the micropost array. Scale bar, 150 μm . The inset is magnified in c to show a spread cell. (d) A high-magnification, fluorescent image of the tips of the microposts under a cell. (e,f) The base positions of the same microposts are shown in e and the GFP-expressing cell is shown in f. These three images were acquired with an $\times 40$ oil-immersion objective. Scale bar, 20 μm .



86 | Optionally, an image of the undeflected positions of the microposts can be acquired for traction force measurement. Adjust the fine focus to $\sim 1 \mu\text{m}$ above the base of the microposts. Acquire an image, known as the ‘base image’ (Fig. 5e).

87 | Optionally, if the target cell is fluorescently labeled, whether live or fixed, an image of the cell can also be acquired, referred to as the ‘cell image’ (Fig. 5f).

88 | Repeat Steps 84–87 for as many cells as desired. For live-cell imaging, individual cells can also be repeatedly imaged over time to generate time-lapse movies. Alternatively, if the microscope is equipped with a motorized xy stage, multiple cells can be imaged in rapid sequence.

89 | To analyze traction forces from the acquired images, see Figure 6 for the general algorithm. Detailed instructions for a custom-written MATLAB program are available on request.

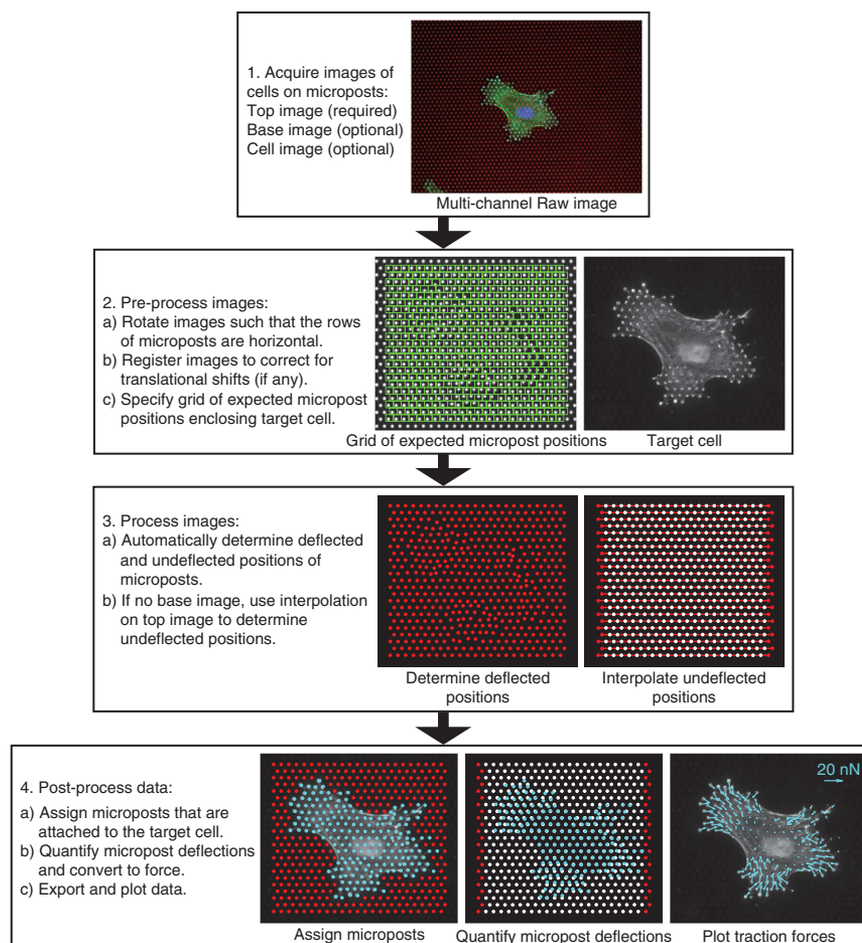


Figure 6 | General algorithm for analyzing traction forces from the micropost array substrate. Key steps for analyzing a representative cell are illustrated. Detailed instructions for a custom-written MATLAB program can be supplied on request.

? TROUBLESHOOTING

Troubleshooting advice can be found in **Table 1**.

TABLE 1 | Troubleshooting table.

Step	Problem	Possible reason	Solution
5, 10, Box 2 (step 5)	Spin-coated photoresist is not uniform and contains traces of particulates	Amount of photoresist is insufficient	Dispense enough photoresist onto the center of the wafer before spin coating
		Air bubbles or particulates deposited during dispensing	Pour directly from the stock bottle or a particulate-free container. Wipe the bottle opening with a lint-free laboratory wipe after each pour to prevent particulate build up
		Wafer surface is not clean	Re-clean the wafer with Piranha solution and dry with a stream of N ₂ . If the wafer cannot be completely cleaned, use a new wafer
10, Box 2 (step 11)	Photoresist pattern on the wafer does not reproduce the photomask features with good fidelity	The photomask is contaminated	Gently clean the photomask with solvents and then blow dry with N ₂
		Iridescent material along the border of the photoresist-silicon interface is observed	Re-develop the wafer
		The features in the photoresist do not match the dimensions of the photomask	Optimize the exposure dose
		For SU-8 photoresist, features are wider at the top than at the bottom of the photoresist	T-topping (wider features at the top than at the bottom of the photoresist) is caused by the photoresist absorbing too much short wavelength UV light (<350 nm), which leads to faster crosslinking of the top part of the photoresist before the bottom can be cross-linked
13	DRIE etching does not reproduce the features on the photomask with good fidelity	Severe undercuts occur during the DRIE process	Double-check the DRIE etch recipe. If necessary, adjust the etching parameters to minimize undercuts
		The etch mask (i.e., the photoresist layer) is etched away during DRIE process	Spin a thicker layer of photoresist to use as a protective layer
43	Molds are difficult to peel from the cover slips after curing micropost substrates	Molds may be insufficiently silanized if the surface was not completely activated with plasma	Optimize the plasma cleaning parameters in Step 33. Tune the resonance frequency to obtain the brightest plasma possible. If the specified power setting cannot be achieved, increase the duration
		Molds have been used several times and have lost some of their silane coating	Molds should be re-silanized after two castings. Molds that have been used four times should be discarded in favor of new molds

(continued)



TABLE 1 | Troubleshooting table (continued).

Step	Problem	Possible reason	Solution
		Molds with deeper holes are more difficult to peel than molds with shorter holes	Score the cast PDMS along the edge of the mold with a razor blade to loosen the mold and make it easier to peel. Additionally, molds can be peeled away under ethanol, although this requires that the substrates be supercritically dried
66	Micropost arrays are collapsed after stamping	Too much pressure was applied after stamp was placed on top of arrays	Apply gentle pressure using blunt tweezers. The weight of the tweezers is sufficient pressure
		The act of peeling off the stamp produced compressive forces that collapsed regions of the arrays	Rather than peel it off, flick off the stamp so as to not compress the PDMS against the arrays. Submerge the substrate and stamp in ethanol before flicking off the stamp
	Micropost arrays have missing regions of posts	Taller, thinner microposts can be ripped off by sticky 30:1 stamps that have been left in contact on the order of minutes	Use 10:1 stamps, which are less sticky than 30:1 stamps
78–80	Cells do not adhere to micropost arrays or are adherent but not spreading	This could be a biologically meaningful result. Cells may be responding to the stiffness of the microposts and unable to spread	Test the effect of micropost rigidity on cell adhesion by seeding cells on micropost arrays of varying stiffness
		Fibronectin printing quality was low	Stain substrate for fibronectin as described in Box 3 to check whether fibronectin printing was uniform
		Cells were overtrypsinized	Reduce trypsin concentration and neutralize after the minimal amount of time needed to detach cells. Pellet cells by centrifugation and remove trypsin-containing media
		Micropost array spacing is not optimal for the target cell type	Optimize the micropost array spacing in the mask design before starting master fabrication
84	Imaging quality is low; illumination is uneven with opposite sides of the field of view being over- and underfocused	Substrate is not level, possibly because of casting the negative mold without squeezing out the excess PDMS or not resetting the set screws in the microscope stage adaptor	Only use substrates that appear to be level, which can be inspected after cutting off the overflow PDMS in Step 54. Adjust the microscope stage adaptor if necessary
	Imaging quality is low; micropost labeling is non-uniform with bright spots due to DiI crystals	DiI solution is not properly filtered to remove DiI aggregates	Filter DiI solution again through a membrane filter
	Imaging quality is low; DiI is present in cellular organelles, making it difficult to see microposts under the cell	Certain cell types will uptake DiI from the PDMS. The rate of uptake varies from 1 day to up to a week, depending on the cell type	Manually correct micropost positions in traction force analysis (Fig. 6) to compensate for errors as a result of the cellular uptake of DiI. Alternatively, the unprinted regions of the microposts can be fluorescently coated with BSA–Alexa Fluor 594

(continued)

PROTOCOL

TABLE 1 | Troubleshooting table (continued).

Step	Problem	Possible reason	Solution
Box 2 (Step 5)	Spin-coated photoresist is raised along the wafer edge, leading to poor wafer contact with mask aligner	More viscous photoresists (e.g., SU8 2050 and higher) tend to bead up along the edge of the wafer, known as 'edge bead'	Remove the edge bead by dispensing 10 ml of propylene glycol monomethyl ether acetate through a wide-bore syringe needle onto the edge of the wafer while spinning the chuck at 1,500 r.p.m. Angle the syringe away from the center of the wafer and in the direction of rotation
Box 2 (Step 6)	Soft-baked photoresist has an uneven wavy pattern	Photoresist is either over- or underbaked	Optimize baking time based on manufacturer's guidelines
Box 2 (Step 12)	Photoresist pattern delaminates after hard bake	For very tall structures (usually >250 μm), thermal stresses during baking will cause the photoresist to bow and peel from the silicon	Do not hard-bake the wafer. Only use it for PDMS castings at 65 $^{\circ}\text{C}$ instead of 110 $^{\circ}\text{C}$ where specified

● TIMING

Steps 1–22, Micropost array design and master fabrication: ~2 weeks (Steps 1 and 2: up to 2 weeks; Steps 3–22: 12 h)

Steps 23–34, Micropost array–negative mold production: 2.5 h

Steps 35–44, Micropost array substrate fabrication: 22–24 h

Steps 45–73, Micropost array substrate functionalization: 4 h

Steps 74–80, Cell culture on micropost array substrates: 1–24 h, depends on experiment

Steps 81–89, Imaging cells on micropost array substrates: variable, depends on experiment

Box 1, Supercritical drying of collapsed micropost substrates: 1 h

Box 2, Stamp master fabrication for micropatterned functionalization: 2 weeks (Steps 1–2: up to 2 weeks; Steps 3–13: 5 h to overnight)

Box 3, Fixation and immunofluorescence staining of cells on micropost array substrates: 7–8 h

Box 4, Quantification of gene expression in stem cells on micropost array substrates: 3 h

Box 5, Oil Red O staining for adipogenic differentiation of stem cells on micropost array substrates: 2 h

Box 6, ALP staining for osteogenic differentiation of stem cells on micropost array substrates: 2 h

ANTICIPATED RESULTS

With this protocol, we have established a library of silicon micropost array masters with a uniform post diameter d of 1.83 μm and post heights L spanning from 0.97 to 14.7 μm . These measurements were determined precisely with both the SEM and the surface profilometer, and have standard deviations that are calculated to be <8% across the entire wafer. The replica-molded PDMS substrates therefore have a >1,000-fold range of micropost rigidity from 1,556 $\text{nN } \mu\text{m}^{-1}$ ($L = 0.97 \mu\text{m}$) down to 1.31 $\text{nN } \mu\text{m}^{-1}$ ($L = 14.7 \mu\text{m}$; **Fig. 2c**). In the first example of application of these substrates, the CSK and FA organization of cells is visualized on micropost arrays of varying rigidity (**Fig. 7**). LifeAct-GFP, which labels F-actin, enables live visualization of stress fibers (**Fig. 7a**). To visualize stress fibers as well as FAs in

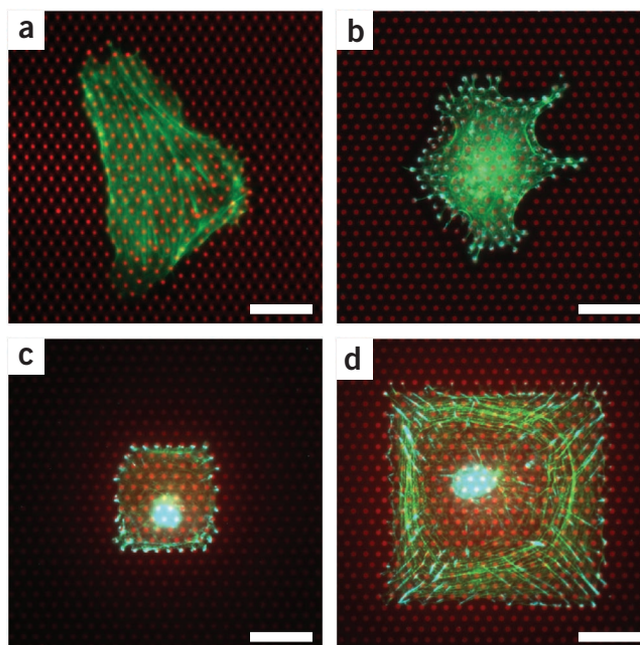
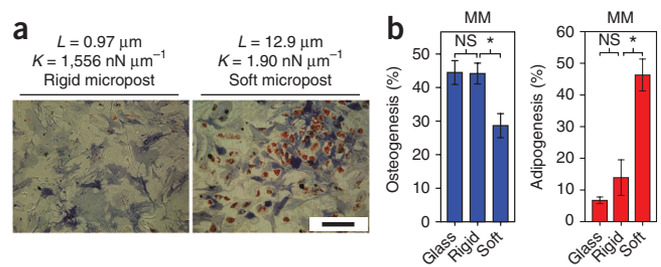


Figure 7 | Representative images of cells on microposts. **(a)** A live cell, expressing LifeAct-GFP to visualize F-actin, on microposts with $K = 7.22 \text{ nN } \mu\text{m}^{-1}$. **(b)** A fixed cell, stained for F-actin (green) and vinculin (cyan). $K = 3.78 \text{ nN } \mu\text{m}^{-1}$. **(c)** A fixed cell constrained to a 30 $\mu\text{m} \times 30 \mu\text{m}$ micropattern and stained for F-actin (green) and vinculin (cyan). $K = 18.19 \text{ nN } \mu\text{m}^{-1}$. **(d)** A fixed cell constrained to a 75 $\mu\text{m} \times 75 \mu\text{m}$ micropattern and stained for F-actin (green) and vinculin (cyan). $K = 15.75 \text{ nN } \mu\text{m}^{-1}$. Scale bars, 20 μm .

Figure 8 | Analysis of stem cell differentiation on micropost array substrates. **(a)** Micrographs of hMSCs on rigid and soft microposts that have been stained for alkaline phosphatase activity (blue) and lipid droplet formation (red) to indicate osteogenic and adipogenic markers, respectively. L and K indicate the micropost height and spring constant for the substrates shown in the corresponding micrographs. Scale bar, 300 μm . **(b)** Quantification of percentage of differentiating cells on rigid and soft microposts, as well as on cover glass as a control. Asterisk (*) indicates statistical significance with $P < 0.05$. NS indicates statistical insignificance with $P > 0.05$. Reprinted with permission from reference 60.



fixed cells, we can use immunostaining (**Fig. 7b–d**). After fixation with 3.7% (vol/vol) paraformaldehyde, cells are labeled with fluorophore-conjugated phalloidin, which stains F-actin. For FA staining, hMSCs can be further incubated with a primary antibody for vinculin followed by a fluorophore-conjugated, isotype-specific, anti-IgG secondary antibody. After obtaining images of either live or fixed cells on the microposts, quantification of subcellular traction forces can be performed using a custom MATLAB program (**Fig. 6**). Moreover, FA areas can be analyzed using suitable segmentation algorithms and applied to further processing of the traction force data^{60,102}.

In the second application, the effect of micropost rigidity on hMSC osteogenesis and adipogenesis is examined. hMSCs are sparsely plated on PDMS micropost arrays with the same post diameter but different heights, and then exposed to mixed differentiation media containing osteogenic- and adipogenic inductive factors. After 2 weeks, the hMSCs are fixed and stained for ALP activity (for osteogenesis) and lipid droplets (for adipogenesis; **Fig. 8a**). The mean percentages of hMSCs expressing osteogenic and adipogenic markers as a function of micropost rigidity are quantified by counting the positively stained cells and divided by the total cell number (**Fig. 8b**). As this example indicates, rigid micropost arrays preferentially induce osteogenic lineage commitment, whereas soft micropost arrays promote adipogenic differentiation in hMSCs.

ACKNOWLEDGMENTS We acknowledge financial support from the National Institutes of Health (EB00262, HL73305 and GM74048); the Army Research Office Multidisciplinary University Research Initiative; the Material Research Science and Engineering Center, the Nano/Bio Interface Center, the Institute for Regenerative Medicine, and the Center for Musculoskeletal Disorders of the University of Pennsylvania; and the New Jersey Center for Biomaterials (RESBIO Resource Center). M.T.Y. was partially supported by the National Science Foundation Integrative Graduate Education and Research Traineeship program (DGE-0221664). J.F. and Y.-K.W. were both partially supported by the American Heart Association Postdoctoral Fellowship. R.A.D. was partially supported by a National Science Foundation Graduate Research Fellowship. We acknowledge the Massachusetts Institute of Technology Microsystems Technology Laboratories for support in microfabrication.

AUTHOR CONTRIBUTIONS M.T.Y. and J.F. designed and fabricated the micropost array masters. M.T.Y. wrote the traction force analysis software. J.F., Y.-K.W. and C.S.C. conceived and designed stem cell experiments with micropost array substrates. J.F., Y.-K.W., M.T.Y. and R.A.D. performed experiments and analyzed data. M.T.Y., J.F., R.A.D. and Y.-K.W. wrote the manuscript.

COMPETING FINANCIAL INTERESTS The authors declare no competing financial interests.

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