

Directing Stem Cell Fate by Controlling the Affinity and Density of Ligand–Receptor Interactions at the Biomaterials Interface**

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Mesenchymal stem cells (MSCs) are adherent multipotent stem cells from bone marrow and potentially numerous other tissues^[1] that serve as an attractive model system for evaluating the influence of extracellular cues on stem cell differentiation. MSCs have been shown to commit to several lineages including: bone, cartilage, fat, and smooth muscle, as well as transdifferentiation to skeletal muscle and neural fates.^[1b,2] Our research group^[3] and others^[2a,b,4] have used MSCs to demonstrate the importance of cytoskeletal tension during lineage specification and commitment. For example, MSCs that were cultured either on stiff substrates^[2b] or patterned on surfaces that promote cell spreading or cytoskeletal tension^[3,4b] all favored an osteogenic program that depended on increased contractility of the actomyosin cytoskeleton. Other reports have demonstrated the use of materials that are modified with cell adhesion ligands to promote MSC osteogenesis;^[5] we reasoned that the molecular characteristics of the adhesion ligands—including the affinity and density—would influence the cytoskeleton of the cell and may, therefore, serve to direct the differentiation pathways of adherent MSCs. Herein we report that the biomolecular interactions between cells and their substrates can be tuned to promote osteogenesis, myogenesis, or neurogenesis of cultured MSCs. This work provides an example of the use of molecular engineering to control the influence that materials have in regulating cell function.

We used self-assembled monolayers (SAMs) of alkanethiolates on gold (anchored through the thiol group) as model substrates, because these surfaces allow excellent control over the ligand–receptor interactions that mediate cell adhesion, in part, because they are structurally well-defined and, in part, because the use of monolayers that are terminated with an oligo(ethylene glycol) group are highly effective at preventing nonspecific adsorption of proteins.^[6] Monolayers, to which short peptide-adhesion ligands are immobilized, have been

used to study several aspects of cell adhesion, including that of embryonic and mesenchymal stem cells, and are an established model for these applications.^[7] We prepared substrates by immobilizing either the linear peptide GRGDSC (linRGD) or the cyclic peptide RGDfC (cycRGD, where f denotes an F residue having the D configuration) to monolayers presenting a maleimide group at a density of 1 % (high density) or 0.1 % (low density) against a background of tri(ethylene glycol) groups (Figure 1a).^[8] The cyclic peptide has approximately two orders of magnitude higher affinity for the $\alpha_v\beta_3$ integrin—an important receptor in the adhesion and osteogenesis of MSCs^[9]—than does the linear peptide.^[10] We used self-assembled monolayer desorption ionization (SAMDI) mass spectrometry to confirm immobilization of the peptides to the maleimide group (Supporting Information, Figure S1).

We cultured MSCs under standard growth conditions (see Supporting Information) for ten days on substrates having a bare gold film, a fibronectin-coated gold film (Fn), or on monolayers presenting linRGD, cycRGD, or a scrambled form of the linear peptide that we have demonstrated to be inactive (KR DGVC).^[11] For the monolayers, peptides were present at a density of 1 % relative to total alkanethiolate (high density) or 0.1 % (low density). We then fixed and stained the cells to observe alkaline phosphatase (AP) expression, which is an early marker for osteogenesis. We detected elevated AP expression for cells on the fibronectin (48 % of cells stained for AP) and cycRGD substrates (44 % on high density, 30 % on low density) compared to a lower level of expression for cells adherent to the bare gold film (25 %; Figure 1b). MSCs cultured on monolayers presenting linRGD, at either high or low density, expressed AP at levels that are comparable to cells cultured on the unmodified bare gold. Control experiments, which used monolayers presenting no peptide or the scrambled RDG peptide, showed insignificant levels of cell adhesion and were not included in the analysis. We confirmed these trends by using reverse transcriptase PCR (RT-PCR) to quantitate the amount of AP mRNA transcript. Again, we found higher expression for cells cultured on monolayers presenting cycRGD and the fibronectin-coated substrate relative to those cultured on the linRGD-terminated monolayers (Figure 1c). These results suggest that fibronectin and the monolayers presenting cycRGD promote osteogenesis.

To further investigate the influence of the monolayers on differentiation, we performed immunofluorescence staining of several lineage-specific markers. For example, cells that differentiate into osteoblasts express the runt-related transcription factor 2 (Runx2). MSCs cultured on fibronectin and cycRGD surfaces show a higher level of nuclear Runx2

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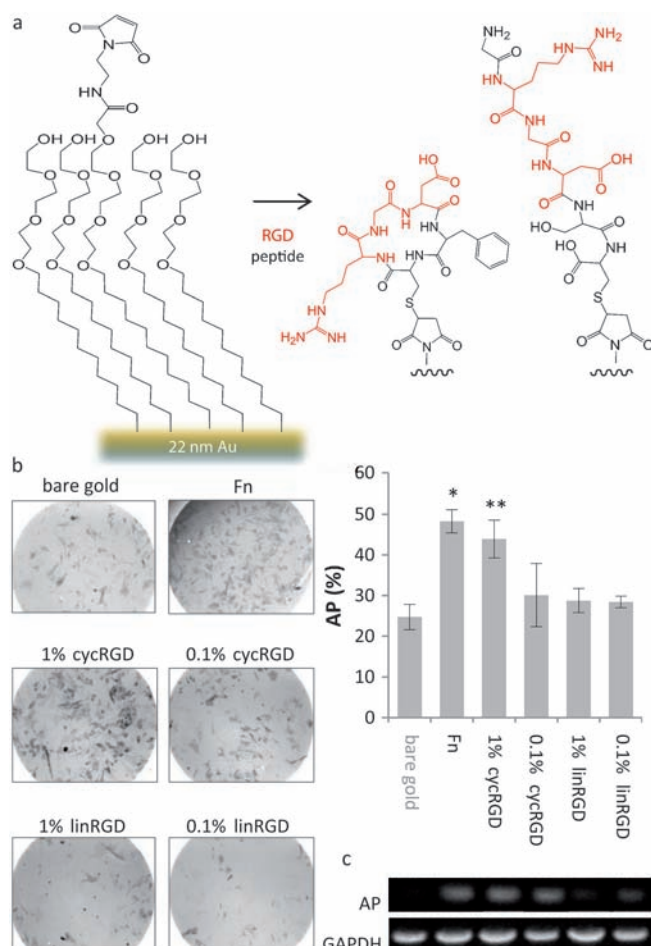


Figure 1. Monolayers presenting a high-affinity adhesion ligand promote osteogenesis in adherent mesenchymal stem cells. a) An array of monolayers were prepared on a glass slide having gold islands and present a maleimide group to allow immobilization of RGD peptides (red). b) Left: Phase contrast images of mesenchymal stem cells stained for the osteogenesis marker, alkaline phosphatase (AP, dark gray), after 10 days of culture on monolayers presenting the linear and cyclic RGD peptides at two surface densities. Right: Quantitation of the percentage of cells expressing AP on the different monolayers. c) RT-PCR of AP transcript expression compared to control glyceraldehyde-3-phosphate dehydrogenase (GAPDH) transcripts. Error bars represent standard deviations of at least three experiments. Statistical significance compared to bare gold: * p value < 0.001, ** p value < 0.02.

compared to cells cultured on the linRGD and bare gold surfaces (Figure 2a, b), which is in agreement with the results from staining and RT-PCR of alkaline phosphatase (Figure 1b, c). To determine if the linRGD surface promotes differentiation towards other lineages, we immunostained cells for a skeletal-muscle marker, the early myogenic regulatory transcription factor MyoD. MSCs cultured on monolayers presenting linRGD at high density showed the highest level of MyoD expression. Cells cultured on monolayers presenting cycRGD (at both high and low densities of ligand) or monolayers presenting the linRGD peptide at low density show elevated expression of MyoD compared to cells cultured on bare gold and fibronectin-coated gold but

significantly less than cells cultured on monolayers presenting linRGD at high density (Figure 2b). We also immunostained cells for the neurogenic marker β -tubulin to assess whether certain combinations of peptide affinity and density could promote differentiation to neuron-like cells. As the affinity and density of the peptide ligand were decreased, we observed an increase in β -tubulin expression, with the highest percentage of β -tubulin-expressing cells occurring on the monolayers that present linRGD at low density (11 %, Figure 2a, b).

We performed RT-PCR to verify the trends in immunofluorescence and found a decrease in Runx2 expression as the affinity and density of the immobilized ligands were reduced (Figure 2c). In contrast, expression of the myogenic marker genes (Desmin, MyoD) increased in these samples and showed a maximum expression in cells that were cultured on monolayers presenting linRGD at high density. Expression of the neuronal markers (β -tubulin, CEND1) was detected primarily in cells cultured on monolayers presenting linRGD at low density, although we did detect expression of CEND1 in cells cultured on bare gold. In addition to neural specific markers, we note that the expression of myogenic markers is elevated on the low-density linRGD surface, which suggests that a fraction of cells on this peptide surface are specifying markers associated with muscle differentiation. Taken together, these results suggest that the monolayers that present high-affinity ligands promote osteogenesis, and monolayers that present low-affinity ligands primarily promote myogenesis at a high density and neurogenesis at a low density of ligand (Figure 2d).

Since cell morphology has previously been suggested as a qualitative marker of differentiation,^[2b] we characterized the changes in the shape of MSCs that were cultured on different substrates. A greater number of cells initially adhered to the bare gold, fibronectin-coated gold, and cycRGD surfaces (Supporting Information, Figure S2). Cells showed a higher proliferation rate on surfaces coated with fibronectin, when compared to bare gold or peptide-modified surfaces. MSCs cultured for one week on monolayers presenting cycRGD remained well spread with a cuboidal phenotype, while cells cultured on linRGD displayed an elongated morphology, similar to myoblasts (Figure S2a). For the monolayers that present linRGD at low density, many of the cells extended long processes characteristic of neuronal-like cells. These morphological differences are consistent with commitment of the MSCs to osteogenesis, myogenesis, or neurogenesis programs.^[2b] While the cells continued to proliferate, we noted a decrease in total cell area and nuclear area after one week in culture for cells adherent to the linRGD surfaces (Figure S2b). Changes in nuclear area have previously been shown to influence gene expression and cell differentiation.^[12]

The influence of ligand affinity and density on differentiation is consistent with a model wherein substrates presenting ligands of higher affinity and at higher density lead to more spreading of cells and greater tension in the cytoskeleton, which favors an osteogenic outcome. Previous studies have demonstrated the importance of cytoskeletal tension and focal-adhesion assembly in directing the differ-

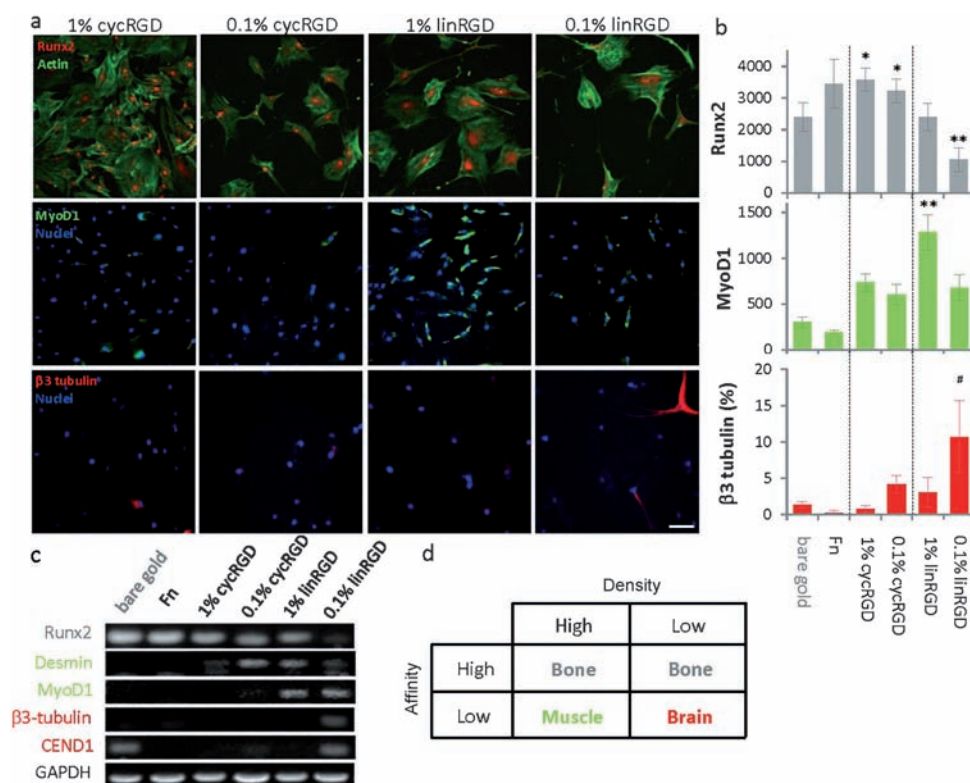


Figure 2. The density and affinity of an adhesion ligand influence the differentiation of mesenchymal stem cells. Cells were cultured on monolayers presenting either the cyclic or linear RGD peptides at a high or low density and differentiation was analyzed using a, b) immunofluorescence imaging of markers for osteogenesis (Runx2), myogenesis (MyoD1), and neurogenesis (β3-tubulin) and c) expression analysis by RT-PCR of lineage-specific transcripts. d) Table summarizing the preferred differentiation outcome for cells cultured on the four monolayer surfaces. Error bars represent standard deviations from three replicates. Statistical significance compared to bare gold: **p* value < 0.02, ***p* value < 0.002, #*p* value < 0.05 as determined using Student's *t*-test. Scale bar is 20 μm.

entiation of MSCs.^[2b,3,4b] We immunostained MSCs for filamentous actin and the focal-adhesion protein vinculin. Cells that were adherent to monolayers presenting cycRGD displayed a higher degree of spreading, more stress fibers, and more focal adhesion structures as compared to cells on monolayers presenting the linRGD peptide (Figure 3). We also immunostained MSCs for non-muscle myosin IIa and IIb to evaluate differences in contractility in cells on the different substrates. After normalizing the fluorescence data, a greater fraction of cells expressed high levels of myosin IIb when cultured on monolayers presenting cycRGD (36% on high density, 40% on low density) as compared to cells cultured on monolayers presenting linRGD (13% on high density, 25% on low density; Figure 4a). For myosin IIa, we observed a comparable level of total fluorescence in cells cultured on the various monolayers (Supporting Information, Figure S3). This result agrees with a report that showed expression of this isoform to be relatively insensitive to variations in substrate elasticity.^[2b] Since cells cultured on the linRGD surfaces display the highest expression of myogenic markers, we immunostained the cells for the muscle-specific myosin heavy chain (MYH). The number of cells expressing high levels of MYH increased significantly as the affinity and density of the cell-adhesion peptide decreased, consistent with a previous

report that demonstrated increased myogenesis with a decrease in cell contractility using substrates with variable mechanical properties.^[2b]

To confirm the important role that cell contractility plays in differentiation, we cultured MSCs in a medium that was supplemented with blebbistatin, an inhibitor of non-muscle myosins that has little effect on muscle-specific isoforms.^[2b] Cells were allowed to fully adhere and then the medium was exchanged with medium containing blebbistatin at a concentration that does not significantly perturb cell shape. We found that treated cells showed a decrease in osteogenesis, as determined by alkaline-phosphatase staining and an increase in expression of MyoD and β3-tubulin (Figure 4b and Figure S4). This result supports our hypothesis that the monolayers having a high-affinity ligand promote a more contractile cytoskeleton in adherent cells, which is

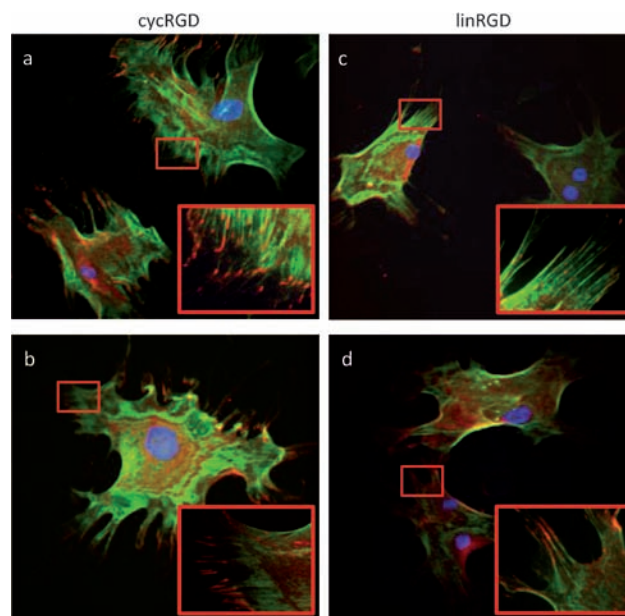


Figure 3. The affinity and density of a cell-adhesion peptide influence focal adhesion and stress-fiber formation. Immunofluorescence images of mesenchymal stem cells stained for filamentous actin (green), vinculin (red), and nuclei (blue). Surfaces presenting 1% (a) and 0.1% (b) cyclic RGD peptide and surfaces presenting 1% (c) and 0.1% (d) linear RGD peptide.

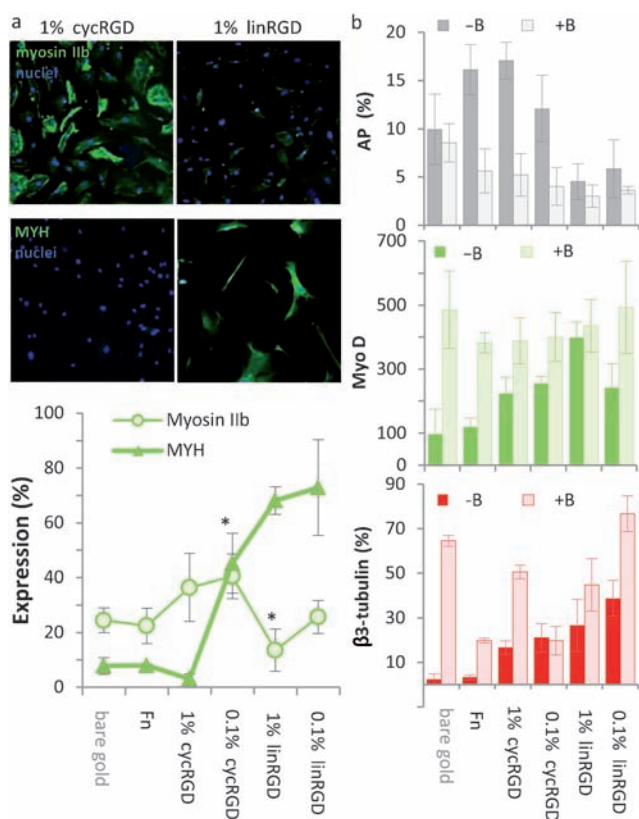


Figure 4. The differentiation of mesenchymal stem cells on peptide surfaces is influenced by non-muscle myosin. a) Immunofluorescence images and quantitation of cells stained for non-muscle myosin IIb and muscle-specific myosin heavy chain (MYH). *statistical significance p value < 0.005. b) Lineage marker expression as determined by immunofluorescence for MSCs exposed to the non-muscle myosin inhibitor blebbistatin (+B) and control cultures that were not exposed (–B): markers for osteogenesis (AP), myogenesis (MyoD1), and neurogenesis (β3-tubulin). Error bars represent standard deviations of at least three replicates.

necessary for the osteogenic preference exhibited by these cells.

Engler, Discher et al. demonstrated the influence of substrate stiffness on MSC differentiation and found that stiff matrices promoted osteogenesis, those of intermediate stiffness promoted myogenesis, and soft matrices promoted neurogenesis.^[2b] Our study demonstrates a similar trend in lineage specification, where monolayers presenting high-affinity peptides promote osteogenesis, those presenting a low-affinity peptide at high density promote myogenesis, and those that present a low-affinity peptide at low density promote neurogenesis. In both examples, the differentiation outcomes depend on the ability of the substrate to oppose the traction forces exerted by the adherent cells. Substrates that present ligands that have a higher affinity for the integrin-adhesion receptors are known to increase the spreading of adherent cells, with a corresponding increase in the traction forces applied by the cell.^[13] This idea is further supported by a recent report by Mooney and co-workers that demonstrated the importance of traction-mediated reorganization of the matrix to direct MSC fate within 3D hydrogels.^[9a] Osteo-

genesis was found to occur preferentially at an optimal stiffness where MSCs can maximize the number of adhesive contacts with the surrounding matrix.

In conclusion, we have illustrated a molecular approach to engineering substrates used in stem cell cultures; herein, we have shown that the affinity and density of ligands at the cell-biomaterial interface can be engineered to influence the fate of stem cells. This demonstration provides another method to the set of materials science based approaches that have been used to direct stem cell fate. The relationships identified in this study might also serve as design rules for the modification of other materials used in stem cell cultures. It further suggests that optimal adhesive microenvironments that favor specific differentiation outcomes may exist. Understanding the relationship between adhesion and lineage could assist efforts in defining the MSC niche, as well as studies aimed at elucidating the factors that control stem cell lineage and differentiation. We expect that the use of structurally well-defined mimics of the extracellular matrix will be important for controlling cellular activities in a broad range of contexts.

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