Controlled Heterogeneous Stem Cell Differentiation on a Shape Memory Hydrogel Surface

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The success of stem cell therapies is highly dependent on the ability to control their programmed differentiation. So far, it is commonly believed that the differentiation behavior of stem cells is supposed to be identical when they are cultured on the same homogeneous platform. However, in this report, we show that this is not always true. By utilizing a double-ion-triggered shape memory effect, the pre-seeded hMSCs were controllably located in different growth positions. Here, we demonstrate for the first time that the differentiation behavior of hMSCs is highly sensitive to their growth position on a hydrogel scaffold. This work will not only enrich the mechanisms for controlling the differentiation of stem cells, but also offer a one-of-a-kind platform to achieve a heterogeneously differentiated stem cell-seeded hydrogel scaffold for complex biological applications.

Stem cells can differentiate into various cell lineages with different functions, thus holding great potential in tissue engineering and regenerative medicine. Recent studies have shown that the fate of stem cells can be controlled by both chemical and physical factors. For the biochemical factors, it is well-established that several chemicals including hormones, growth factors and small molecules can modulate the biological activity of stem cells and direct their differentiation down specific pathways. On the other hand, the physical properties including stiffness, porosity, surface pattern and geometry of the extracellular matrix have been recognized as crucial factors in regulating the differentiation of stem cells. For instance, human mesenchymal stem cells (hMSCs) are shown to specify lineage and acquire phenotypes with extreme sensitivity to the elasticity of the substrate. A deeper analysis reveals that the stem cells are able to sense the mechanical property of their surrounding environment via integrin binding and initiate the specific differentiation process.

In the transition from fundamental studies to clinically relevant applications of stem cells, it is very important to elucidate the behavior of stem cells in complex geometrical scaffolds in which the stem cells are located in different positions and supported by the platform in different ways. However, the influence of the growth position on the differentiation behavior of stem cells is unknown, but of great implication to understanding regenerative phenomena as well as directing design of tissue engineering scaffolds. It is generally difficult to achieve various culture positions for stem cells on the same platform since normal 2D cultures only enable the cells to grow on the top of platforms. In our previous work, we demonstrated polyacrylonitrile (PAN)-based copolymer hydrogels were able to exhibit a shape memory effect by manipulating the reversible formation of dipole-dipole interactions in response to zinc ions. Furthermore, the self-rolling/folding and arbitrary 3-D shape changes could be achieved and well maintained if the gel was pre-programmed with a surface pattern. The goal of this work was to explore the differentiation behavior of hMSCs when they were grown on different places of a single piece hydrogel. The hMSCs were cultured on the surface of PAN based hydrogels, while their growth location was altered by a shape memory effect. Finally, the hydrogel-hMSC constructs were cultured in a bipotential differentiation medium (osteogenic and adipogenic). As a result, we were able to examine the direct influence growth location could have on the fate choice of hMSCs. This work demonstrates that the location of a scaffold where stem cells grow may be used to control complicated cellular functions such as stem cell differentiation. These results are not only fundamentally interesting, but also could lead to simple and controllable scaffold formulations for the application of hMSC-based regenerative medicine.

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Results

Double-ion controlled triple shape memory. In this study, on the basis of the Zn$^{2+}$ responsive behavior of PAN hydrogel reported in our previous work$^{23}$, the hydrogel was further incorporated with carboxyl groups which were used to fix the temporary shape of the hydrogel due to their coordination with calcium ions$^{26}$. We copolymerized monomers of acrylonitrile (AN), acrylic acid (AA) and a biodegradable crosslinker, N,N'-bis(acryloyl)cystamine (BAC), to form dipole-dipole reinforced (DDR) hydrogels whose molecular structure is depicted in Scheme S1. Similar to what we reported previously$^{23}$, due to the reinforcement from strong non-covalent dipole-dipole interactions among cyano groups, the poly(acrylonitrile) (PAN) hydrogels presented here exhibited 1.5 MPa tensile strengths, 160% elongation at break, and 9.3 MPa compressive stresses at 80% strain, with equilibrated water contents ranging from 78% to 83% (Supplementary Fig. S2 and Table S1). In order to avoid stiffness-favored differentiation of stem cells as reported$^{27}$, the moduli of all selected samples shown in Table S1 are beyond the normal tissue-level which may specifically induce osteogenesis or adipogenesis. Besides, the gels are degradable in a redox environment (Supplementary Fig. S3).

We began to investigate the influence of divalent ions on the presented hydrogels. Here, Zn$^{2+}$ and Ca$^{2+}$ were selected since they were relatively biocompatible and can selectively bind to CN and COO$^-\text{2}$ groups under certain circumstances. We first tested the influence of Ca$^{2+}$ ions on the equilibrium water content (EWC) of the AN-AA-PBS hydrogel (equilibrated in PBS). The hydrogel samples were equilibrated in either 3.7 M ZnCl$_2$ (denoted as AN-AA-Zn) or 5 mM CaCl$_2$ (denoted as AN-AA-Ca) solution for at least three days prior to testing. Since Ca$^{2+}$ can serve as chelation center among carboxyl groups and further densify the hydrogel network, the EWC of AN-AA-Ca is about 50% lower than AN-AA-PBS (Fig. 1a). Conversely, for AN-AA-Zn hydrogels, Zn$^{2+}$ can selectively disassociate dipole-dipole interactions and thus significantly loosens the hydrogel network. As presented in Fig. 1a, the EWC of AN-AA-Zn (96.7%) is 10% higher than that of AN-AA-PBS. Additionally, the influence of divalent ions on the moduli of selected hydrogels was observed. When dipole-dipole interactions were disassociated by Zn$^{2+}$, the modulus of the selected hydrogel decreased sharply to 179 KPa (Fig. 1b). Also, we found that the strong chelation formed among carboxyl groups and Ca$^{2+}$ was able to significantly enhance the modulus of the selected hydrogel to 9.53 MPa (Fig. 1b). It should be noted that both dipole-dipole and Ca$^{2+}$ ion reinforcement can be reversibly built-in or removed from the system. The shape memory efficiency of AN-AA-PBS hydrogel was characterized by an angle-controlled experiment. First, we put one straight strip AN-AA-PBS hydrogel into a 3.7 M ZnCl$_2$ solution. After reaching adsorption equilibrium, the AN-AA-Zn hydrogel was bent into a “U” shape with the angle between the two arms being fixed at 180 degrees. 0.01 mM EDTA solution (pH = 7.4) was then used to remove the divalent ions from the hydrogel network. As a result, the AN-AA-Zn hydrogel was transformed into D$^-$/C$^+$/D$^+$/C$^-$ state. In a following step, the straight strip shape could be recovered again by equilibrating the AN-AA-PBS hydrogel in a 3.7 M ZnCl$_2$ solution.
hydrogel was then transferred into a 3.7 M ZnCl₂ solution to into a 5 mM CaCl₂ solution. After reaching equilibrium in CaCl₂ as demonstrated above, the U-shape AN-AA-PBS gel was transferred (Phase II). After bending AN-AA-PBS hydrogel into a “U” shape (Phase I) experiments was continued to be used for exhibiting the shape memory effect. Herein, the above hydrogel after the associating/disassociating the dipole-dipole interaction. (Phase I) can be repeated for another two times with selectively switching between D⁺/C⁻ and D⁻/C⁺ states or between D⁺/C⁻ and D⁻/C⁺ states could realize a triple shape memory effect. In the Fig. 1d, an as-prepared hydrogel was cut into a cross-shaped sheet and purified by dialysis in PBS (pH = 7.4) for three days. The hydrogel was then transferred into a 3.7 M ZnCl₂ solution to disassociate the dipole-dipole interactions. At the state of D⁻/C⁺, the gel became soft and able to readily fold into a cubic box (temporary shape 1). Subsequently, the hydrogel was converted into the D⁺/C⁻ state by removing Zn²⁺ with an EDTA solution (pH = 7.4, 0.01 mM). As a result, the box shape was fixed by the re-formation of dipole-dipole interactions. The box-shape hydrogel was further unfolded into a flat shape (temporary shape 2) again and fixed in 5 mM CaCl₂ (D⁻/C⁺). The hydrogel with temporary shape 2 was transferred into a basal medium. After equilibrium in the basal medium and being coated with fibronectin, hMSCs were seeded on the surface of the flat hydrogel. After 24 hours, unattached cells were washed away by refreshing the medium and the cell-seeded construct was folded automatically (D⁺/C⁻ state) by removing Ca²⁺ with EDTA-containing basal medium (0.01 mM). The whole process was completed in five hours. Consequently, the hMSCs were homogeneously seeded on the 6 sides (Detailed characterization can be found in Supplementary Fig. S5, 6). We noted that the EDTA treatment did not cause the detachment of hMSCs.

**Achieving heterogeneous distribution of hMSCs via a shape memory effect.** Based on our discussion above, it is expected that selectively switching between D⁺/C⁻ and D⁻/C⁺ states or between D⁺/C⁻ and D⁻/C⁺ states could realize a triple shape memory effect. In the Fig. 1d, an as-prepared hydrogel was cut into a cross-shaped sheet and purified by dialysis in PBS (pH = 7.4) for three days. The hydrogel was then transferred into a 3.7 M ZnCl₂ solution to disassociate the dipole-dipole interactions. At the state of D⁻/C⁺, the gel became soft and able to readily fold into a cubic box (temporary shape 1). Subsequently, the hydrogel was converted into the D⁺/C⁻ state by removing Zn²⁺ with an EDTA solution (pH = 7.4, 0.01 mM). As a result, the box shape was fixed by the re-formation of dipole-dipole interactions. The box-shape hydrogel was further unfolded into a flat shape (temporary shape 2) again and fixed in 5 mM CaCl₂ (D⁻/C⁺). The hydrogel with temporary shape 2 was transferred into a basal medium. After equilibrium in the basal medium and being coated with fibronectin, hMSCs were seeded on the surface of the flat hydrogel. After 24 hours, unattached cells were washed away by refreshing the medium and the cell-seeded construct was folded automatically (D⁺/C⁻ state) by removing Ca²⁺ with EDTA-containing basal medium (0.01 mM). The whole process was completed in five hours. Consequently, the hMSCs were homogeneously seeded on the 6 sides (Detailed characterization can be found in Supplementary Fig. S5, 6). We noted that the EDTA treatment did not cause the detachment of hMSCs.

**Morphology and gene activity of hMSCs on each side.** To verify if the growth position (top, side and bottom surface) of hMSCs influences the differentiation of hMSCs, the cell-hydrogel construct was transferred into a bipotential differentiation medium whose formulation was designed for inducing both adipogenesis and osteogenesis. After a 7-day differentiation process which is widely used⁹,¹⁰, the cell-hydrogel construct was cut into 6 different pieces according to their position. The morphology of the hMSCs was visualized by immunostaining F-actin and nuclei. It was clear that after the 7-day differentiation, hMSCs attained significantly different morphologies on different sides (Fig. 2a–c). Hanging from the top surface, most hMSCs formed a round shape (Fig. 2a). In contrast, the hMSCs on the side and bottom surface were prone to assume an outstretched shape (Fig. 2b–c). As reported previously⁹,¹⁰, the shape of hMSCs is able to modulate cell traction and thus directs them to differentiate into different lineages. The round-shape hMSCs are inclined to differentiate down the adipogenic pathway, while outstretched-shape hMSCs are inclined to differentiate down...
the osteogenic pathway. Herein, we characterized the heterogeneous differentiation by RNA expression and immunocytochemistry (ICC) assays.

We first tested the differentiation behavior of the hMSCs on the cell-hydrogel construct based on their RNA expression. RNAs were isolated according to vendor instructions and quantitative real-time PCR (qRT-PCR) was performed to determine the expression of characteristic adipocyte and osteocyte genes. In the Fig. 2d–e, the expression was normalized to cells cultured on tissue culture polystyrene (TCPs) under the same conditions, using β-actin as the reference gene. Among the expression of a panel of adipogenic genes from hMSCs on top, side and bottom surfaces, peroxisome proliferator-activated receptor gamma (PPARG) is a crucial regulator of adipogenesis and highly related to the uptake and storage of lipids. Fatty acid binding protein 4 (FABP4) is a carrier protein for fatty acids that is mainly expressed in adipocytes. Adiponectin, which regulates fatty acid breakdown, is encoded by the ADIPOQ gene in the human body. Lipoprotein lipase (LPL) is an enzyme that can hydrolyze triglycerides in lipoproteins. As presented in Fig. 2d, all of these four adipogenic biomarkers were expressed at significantly higher levels in the hMSCs from the top compared with those on the side or the bottom. At the same time, one can find that the hMSCs on the bottom expressed a comparable level of adipogenic biomarkers to those on the top surfaces. When they were grown on the side surfaces, expression of the adipogenic biomarker was much more pronounced compared to that on the top. When the MSCs are cultured on the side surface, expression of these biomarker expressions from the hMSCs on the side surface. Meanwhile, one can find a reverse scenario when examining osteogenesis was greatly inhibited due to the competition of adipogenesis while the ‘‘wall-hanging’’ growth condition promotes the process of osteogenesis.

**Histological staining for differentiated hMSCs on each side.** We further aimed to explore certain chemicals and enzymes expressed within specifically differentiated hMSCs. Osteogenesis can be reflected by alkaline phosphatase (ALP), stained with Fast Blue salts. Likewise, adipogenesis in hMSCs can be demonstrated by the presence of lipids, visualized by staining with Oil Red O. We conducted a histological staining for both lipid and ALP in hMSCs cultured in the three positions. Staining for nuclei confirmed that the lipid droplets corresponded to single cells rather than clusters (Supplementary Fig. S8). As presented in Fig. 3a–c, stem cells differentiated positively on all surfaces. While nearly all the differentiated cells committed adipogenesis on the top, osteogenesis dominated on the side and bottom. It is also observed that hMSCs on the bottom are more prone to choose the osteogenic pathway rather than the adipogenic pathway. The quantification of the heterogeneous differentiation is summarized in Fig. 3d. In addition, we quantified the calcium deposition with alizarin red staining in order to confirm the osteogenic differentiation. As presented in the Supplementary Fig. S9, the results were consistent to the ALP staining observed above.

**ICC assay for hMSCs on hydrogel surfaces.** The characteristic antigens on differentiated hMSCs were visualized via ICC assay. Herein, PPARG was used as a biomarker to visualize adipogenesis and RUNX2 was used as a biomarker to observe osteogenesis. Nuclei were stained by DAPI. Again, the hydrogel-cell construct was cut into different parts according to the position (i.e. top, side and bottom surface). In Fig. 4, it is observed that the hMSCs expressed PPARG more robustly when they were cultured on the top. In contrast, RUNX2 was weakly expressed on the top, indicating the osteogenesis was greatly inhibited due to the competition of adipogenesis. Meanwhile, one can find a reverse scenario when examining these biomarker expressions from the hMSCs on the side surface. When the MSCs are cultured on the side surface, expression of the osteogenic biomarker was much more pronounced compared to that on the top. We also measured the expression of osteogenic genes from hMSCs on the top, side and bottom surfaces (Fig. 2e). Runt-related transcription factor 2 (RUNX2) is a key transcription factor essential for osteogenic differentiation. COL1A1 (Collagen, type I, alpha 1) is a human gene that encodes the major component of collagen I, which can be found in osteoprogenitor cells. Osteocalcin (OCN) is one kind of noncollagenous protein secreted exclusively by osteoblasts. Osteopontin (OPN) is expressed predominantly by osteoblasts, osteocytes and osteoclasts and has been implicated as an important factor in bone remodeling. In contrast to our finding on the expression of adipogenic biomarkers, the expression of these osteogenic biomarkers was restrained when the hMSCs were grown on the top surface. It is noteworthy that hMSCs on the side surface demonstrate significantly higher expression of osteogenic biomarkers, while hMSCs on the bottom surface present lower expression. The above-mentioned RNA expression assay demonstrates that growth position could significantly influence the differentiation direction of cultured hMSCs. Compared with hMSCs under normal growth conditions, the ‘‘up-side-down’’ growth condition promotes the process of adipogenesis while the ‘‘wall-hanging’’ growth condition promotes the process of osteogenesis.

**Figure 3 |** Representative bright-field images of histologically stained hMSCs on the top (a), side (b) and bottom (c) following 7-day bipotential differentiation medium incubation. Scale bar, 50 μm. (d) Percentage differentiation of hMSCs on different sides of cubic AN-AA-PBS hydrogels. Asterisks denote statistical significance compared with top surfaces (**p < 0.001, t-test). Error bars represent standard error of the mean from 5 individual experiments.
of adipogenic biomarker PPARG (Fig. 4). Comparatively, osteogen-
sis dominated the differentiation behavior of hMSCs on the bottom
surface. The above results clearly demonstrate the growth position
has a strong influence on the differentiation behavior of hMSCs
under normal culture conditions.

Modulating differentiation by cytoskeletal manipulation. It has
been proved that the differentiation of hMSCs depends on the
cytoskeletal contractility7. In this work, we further tested the
influence of several pharmacological agents known to modulate
the cytoskeleton on the differentiation behavior of hMSCs on top,
side and bottom via histological staining as mentioned above.
The inhibitors were added into the media when the cells were seeded
on the top of AN-AA-PBS hydrogel. The concentration of the added
inhibitors were same in both basal and differentiation media. A 10-
day culture (3 days in a basal media and 7 days in a bipotential media)
was employed before the histological staining. The percentage of
hMSCs differentiating to osteoblasts or adipocytes in media
without the addition of pharmacological agents is shown in Fig. 5a.
Nocodazole is able to interfere with the polymerization of
microtubules, thus increasing the cytoskeletal contractility. As seen
in Fig. 5b, the drug indeed reversed the fate choice of hMSCs cultured
on top and gave a high rate of osteogenesis in all cases. Three other
drugs that can directly inhibit contractility of the cytoskeleton—Y-
27632, which inhibits ROCK, Cytochalasin D, which inhibits F-actin
polymerization, and blebbistatin, which inhibits myosin II—resulted
in a significant decrease in osteogenesis in all cases. These three other
drugs could inhibit the cell from forming a stretched shape by downward
dragging force, thus reducing the cytoskeletal tension. In contrast to
that, the cytoskeletal tension could be amplified due to the gravity-
traction induced stretched shape when the cells were grown on the
bottom and the inner cytoskeletal tension drove the most hMSCs
to adopt osteogenic differentiation. The cells hanging upside down
on the top of the platform were not supported and its own gravity
could inhibit the cell from forming a stretched shape.

Discussion

Current studies on stem cells are mostly focused on their behav-
iors on the top of 2D platforms or within 3D hydrogels9,28–34. While
some experimental techniques like hanging drop method has
interesting implications that the growth position may influence
cellular functions of stem cells, it is still not clear whether the
specific fate of stem cells is able to be harnessed by their culture
position35. At the same time, it is highly desirable to achieve
heterogeneous differentiation of stem cells on/in a single piece
scaffold for many clinically relevant applications such as the con-
struction of artificial organs. However, it is generally difficult to
implement heterogeneous biophysical or biochemical cues in one
homogeneous scaffold.

In this study, we developed a novel double-ion-triggered shape
memory hydrogel. Introducing and removing two types of divalent
ions could contribute to reversible dissociation of dipole-dipole pair-
ings and divalent ion-chelation, thereby giving rise to a multiple
shape memory effect. One remarkable feature of the reported hydro-
gel is its capability of automatic shape-changing, eliminating the risk
of contaminating cells via manual manipulation. This shape memory
effect could specifically modulate the differentiation behavior of stem
cells via manipulation of the cytoskeletal contractility. During the
fixation of a temporary shape by dipole-dipole interaction and cal-
cium-chelation, hMSCs were seeded on the hydrogel surface; after
extracting Ca²⁺, the hydrogel-hMSCs construct could automatically
reshape into a cubic box where hMSCs were shown to exhibit het-
erogeneous differentiation behavior at different growth positions.
The “up-side-down” growth position promotes the process of adi-
pogenesis while the “wall-hanging” growth condition promotes the
process of osteogenesis.

One hypothetic mechanism underlying this work could be the
microgravity-induced organization of cytoskeleton36. The gravity
of cells is generally overlooked since the weight of a cell in medium
can only exert a small force of around 0.4 pN37. However, relevant
findings reveal that the cytoskeleton could act as an antenna for
mechanical force and amplify these small forces38. As a result, the
forces in the actin cytoskeleton alone are over 1 million times than
the weight of cells37. In this work, the gravity of cells was counter-
balanced by the supporting force when they were grown on the
bottom and the inner cytoskeletal tension drove the most hMSCs
to adopt osteogenic differentiation. The cells hanging upside down
on the top of the platform were not supported and its own gravity
could inhibit the cell from forming a stretched shape by downward
pulling force, thus reducing the cytoskeletal tension. In contrast to
that, the cytoskeletal tension could be amplified due to the gravity-
traction induced stretched shape when the cells were grown on the
side surface. Cell contraction regulated by biofactors9, signal path-
ways39 and substrate stiffness40 have been reported previously. In this
work, we, for the first time, observed the distinct contraction beha-
viors of cells at different growth sites of a 3-D scaffold. Therefore, it is
expected that the cytoskeletal contraction in cells grown on different
sides of our hydrogels could be dramatically different. The involve-
ment of additional cytoskeletal-disrupting agents further confirms
our speculation that the different cytoskeletal contraction in hMSCs
on different sides could dramatically influence their differentiation
behavior. This is consistent to a previous model reported by Chen
and others8. For example, nocodazole increases the contractility of
the cytoskeleton and was able to drive the majority of hMSCs on all
surfaces towards osteogenic differentiation. In contrast, cells on all
surfaces exposed to molecules (Cytochalasin D, Blebbistatin and Y-
27632) that inhibit contractility were prone to differentiate into adi-
pocytes. Furthermore, we could reduce the tension exerted by the cell
on all surfaces with a corresponding inhibition of osteogenesis by
blocking a portion of cell surface integrin receptor with an antibody.
These results suggest the heterogeneous distribution of hMSCs by the
reported shape memory effect could delicately modulate the cyto-

Figure 4 | After 7 days of culture in bipotential differentiation medium,
cell-hydrogel constructs of different positions (Top, Side and Bottom
surface) were immunostained for adipogenic (PPARG) and osteogenic
(RUNX2) biomarkers. Scale bar, 30 μm.
skeletal contractility within cells and thus manipulate the differentiation behavior of hMSCs.

Collectively, the use of a shape memory effect can be employed to enhance and rationally control differentiation-specific signaling, thereby guiding differentiation in engineered scaffolds. In addition, the effect of the cells' gravity on their differentiation behavior is also firstly reported and employed. Therefore, this study not only provides one-of-a-kind shape memory hydrogels and methodologies for designing tissue engineering scaffolds, but also benefits the fundamental understanding of the differentiation behavior of stem cells in complex geometrical space. It is necessary to point out that we only provide a phenomenal interpretation for the observed heterogeneous differentiation of stem cells on a 3-D scaffold in this study. Nonetheless, the finding reported here hints to us that the different growth positions of cells in a 3D scaffold may influence their other physiological functions, and more experimental and theoretical work is needed to uncover this mechanism from materials science, cell biology as well as biophysics perspectives.

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