Spatial Organization of Functional Groups on Bioactive Supramolecular Glycopeptide Nanofibers for Differentiation of Mesenchymal Stem Cells (MSCs) to Brown Adipogenesis

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Supporting Information

ABSTRACT: Spatial organization of bioactive moieties in biological materials has significant impact on the function and efficiency of these systems. Here, we demonstrate the effect of spatial organization of functional groups including carboxylate, amine, and glucose functionalities by using self-assembled peptide amphiphile (PA) nanofibers as a bioactive scaffold. We show that presentation of bioactive groups on glycopeptide nanofibers affects mesenchymal stem cells (MSCs) in a distinct manner by means of adhesion, proliferation, and differentiation. Strikingly, when the glutamic acid is present in the glycopeptid backbone, the PA nanofibers specifically induced differentiation of MSCs into brown adipocytes in the absence of any differentiation medium as shown by lipid droplet accumulation and adipogenic gene marker expression analyses. This effect was not evident in the other glycopeptide nanofibers, which displayed the same functional groups but with different spatial organization. Brown adipocytes are attractive targets for obesity treatment and are found in trace amounts in adults, which also makes this specific glycopeptide nanofiber system an attractive tool to study molecular pathways of brown adipocyte formation.

INTRODUCTION

Spatial organization of bioactive groups in biological materials is important in function and mechanism of cellular events. The peptide amphiphile (PA) molecules are a versatile class of materials that can self-assemble into a variety of bioactive nanostructures. The PAs integrate structural features of amphiphiles with functions of peptides that can be biologically active. The PAs can be designed for different tasks, as they have tailorable bioactivity depending on the peptide sequence, which can be used in diverse biomedical applications including regenerative medicine and drug delivery. Self-assembly of PA molecules can be triggered by electrolyte addition, changing pH, or coassembling with oppositely charged systems. Through self-assembly, the PAs can form high-aspect-ratio nanofibers, which are powerful and exclusive tools for fabrication of scaffolds due to their unique bioactive extracellular matrix (ECM) mimetic properties, which direct cellular responses and cell fate. In addition, the PA nanofibers present bioactive peptide epitopes with a much higher density compared to single bioactive peptide molecules, and high density epitope presentation is known to strongly enhance the activity of bioactive sequences. Spatial organization of functional groups on PA nanofibers affects induction of biological mechanisms, and understanding the factors affecting bioactivity of PA nanofibers is vital for the successful engineering of supramolecular architectures with desired property and function and for better understanding of the biological processes. Although there are a few studies showing the advantages of spatial organization of peptide functionalities in two dimensions, the effect of supramolecular spatial organization has not been addressed in detail.

In this study, we show the influence of supramolecular spatial organization of bioactive groups in PA nanofibers in cellular processes. For this purpose, the PA nanofiber systems were designed with bioactive glycopeptide amphiphiles. Bioactivity of glucose residues has been shown by investigating the roles of glycosylation in proteins and peptides, the roles of oligosaccharides attached to cell surface, or in a glucose unit conjugated to a serine amino acid side chain was used as a bioactive group in the PA system. Two different amino acids were used as flanking amino acid: glutamic acid and lysine residues for E-Glc-PA and K-Glc-PA, respectively (Figure 1). Flanking amino acid refers to the amino acid type that is located around the bioactive sequences. The K-PA and E-PA molecules lack the bioactive glucose units. The E-Glc-PA was mixed with K-PA and K-Glc-PA was mixed with E-PA and two nanofiber systems with the same amino acid composition and the same number of bioactive groups were obtained; however, as supramolecular...
systems, the spatial organizations of the bioactive groups were distinct in these two systems. Mixture of K-PA and E-PA was used as a control for testing the bioactivity of glucose residues on cellular responses. Rat mesenchymal stem cells (MSCs) were cultured on the PA nanofiber scaffolds, and the effect of spatial organization of functional groups on cellular responses of MSCs was studied. Adhesion and proliferation of the cells exhibited different profiles between the E-Glc-PA/K-PA and K-Glc-PA/E-PA surfaces. We further observed that the E-Glc-PA/K-PA nanofiber system is a striking inducer of brown fat adipogenesis of MSCs, which was significantly much stronger than the K-Glc-PA/E-PA and other control surfaces. This result reported the importance of spatial organization of functional groups in the self-assembled PA systems and demonstrated that different conformational signals create a significant difference on cell behavior of MSCs. In addition, this study introduced a novel scaffold to culture stem cells in order to trigger differentiation into brown fat cells. Brown fat cells are in the focus of obesity related studies; however there is limited access to brown fat cells because they are rarely found in the body. Thus, our results also provide a method for obtaining brown fat cells from stem cells with high yield.

## RESULTS

### Synthesis and Characterization of Peptide Amphiphile Molecules

The E-Glc-PA (lauryl-VVAGES(β-D-Glc)-Am) and K-Glc-PA (lauryl-VVAGKS(β-D-Glc)-Am) molecules were designed as glycopeptide amphiphiles (Figure 1). The K-PA (lauryl-VVAGK-Am) and E-PA (lauryl-VVAGE) were synthesized to form supramolecular nanofibers with the glycopeptides through coassembly, forming two distinct nanofiber systems with the same number of functional groups and amino acids (Figure 1). The coassembly of K-PA and E-PA was formed as a non-glucose bearing PA nanofiber control. The PA molecules are composed of a hydrophobic alkyl tail (lauryl moiety), a β-sheet forming amino acid sequence (-VVAG-), and a charged amino acid for solubility and charge control. Glucose bearing PA molecules (K-Glc-PA and E-Glc-PA) have additional Ser-linked β-D-glucose glycoamino acid as bioactive unit (Figure 1 and Table S1 in Supporting Information). The E-Glc-PA and E-PA bear a negative charge due to glutamic acid (E) residue, and K-Glc-PA and K-PA bear a positive charge due to lysine (K) residues. All PA molecules were synthesized by solid phase peptide synthesis method, purified by preparative high performance liquid chromatography (HPLC), and characterized by liquid chromatography–mass spectrometry (LC–MS).
**PA Nanofiber Characterization.** The oppositely charged PA molecules were mixed and characterized with scanning electron microscopy (SEM) (Figure 2A). SEM images showed that PA nanonetworks were porous and composed of nanofibers, which highly resemble the native ECM found around the cells. This resemblance makes the PA nanofibers a suitable microenvironment for the cells to attach, live, and proliferate. The closer visualization of PA nanofibers was performed by transmission electron microscopy (TEM) or scanning transmission electron microscopy (STEM). TEM and STEM images revealed that ECM-like structure was formed by high-aspect-ratio nanofibers with diameter of 8−10 nm and lengths reaching several micrometers (Figure 2B). Circular dichroism (CD) analyses were performed to understand the secondary structure of PA nanostructures formed upon mixing oppositely charged PA molecules. The CD spectra revealed that mixed PA systems show β-sheet structure, displaying a minimum around 220 nm and a maximum around 204 nm (Figure 2C). The PA molecules at neutral pH showed disordered conformation, except E-Glc-PA (Figure S2) molecule, which showed β-sheet indicating CD peaks. Overall, all three nanofiber systems (E-Glc-PA/K-PA, K-Glc-PA/E-PA, and K-PA/E-PA) showed similar morphologies and molecular organizations.

**Figure 2.** Morphological and structural properties of the peptide amphiphile nanofibers. (A) SEM images show ECM-like structure of PA nanonetworks. (B) TEM images (left and right) and STEM image (middle) show PA nanofibers. (C) Circular dichroism reveals the secondary structure of PA nanofibers.

**Early Cellular Response on the PA Networks: Viability, Adhesion, and Proliferation.** Biocompatibility of the PA nanofiber systems was assessed by monitoring the viability of cells cultured on the nanofiber networks. Cellular viability was checked after 12 h of incubation on the PA nanonetworks in order to discriminate the interference of proliferation on viability results. Viability results showed that all PA nanofiber systems were biocompatible for MSCs, comparable to uncoated tissue culture plates (Figure 3A).

Adhesion behavior of cells on peptide nanofiber network systems is an important initial sign of bioactivity. Relative adhesion data were obtained by normalization of total number of adhered cells on peptide nanofiber networks to total number of cells on bare surface, which was bare tissue culture plate. Total number of cells on bare surface was accepted as one, and cell amount in other groups were normalized to cells on bare surface. Results showed that 5 h after cell seeding, cells adhered more on peptide nanofiber networks compared to bare surface (Figure 3B). This result indicated that PA nanofiber networks provide a favorable environment for cells to adhere. The first significant effect of supramolecular spatial organization of functional groups was also observed during adhesion assay. The number of cells adhered on K-Glc-PA/E-PA was significantly higher than the E-Glc-PA/K-PA, although both nanofiber systems contained the same functional bioactive groups in equal amounts.

**Figure 3.** Early cellular responses on peptide nanofiber networks. (A) Cell viability on peptide nanofiber networks compared to bare surface, after 12 h cultivation period. (B) Number of adhered cells on different peptide nanofiber networks 5 h after cell seeding, in proportion to number of adhered cells on bare surface. (C) Proliferation rate of cells on different peptide nanofiber networks compared to bare surface, after 48 h cultivation period. Different letters denote significance at $p < 0.05$. 742
Effect of culturing on the PA nanofibers on proliferation profile of MSCs was studied by BrdU incorporation assay after 48 h of incubation. When compared to cells cultured on tissue culture plate, proliferation percentages of MSCs were found to be 62% for E-Glc-PA/K-PA, 85.54% for K-Glc-PA/E-PA, and 114.8% for K-PA/E-PA group (Figure 3C). Although E-Glc-PA/K-PA and K-Glc-PA/E-PA nanofibers were observed to affect the proliferation at different levels, this difference was not statistically significant (difference between the two groups, 23%).

Investigation of the Differentiation Response of MSCs to Glucose Bearing Nanofiber Scaffolds with Different Spatial Organizations. Morphological changes of the cells that were seeded on PA nanofiber network groups were tracked in order to observe any changes triggered by different PA networks. Initially, accumulation of lipid droplet-like structures was observed in E-Glc-PA/K-PA system (Figure 4), whereas they were rarely observed in other groups. Therefore, investigation of the influence of different spatial organizations of PA nanofibers on MSC differentiation was specifically focused on adipogenesis.

Oil red-O is a fat-soluble diazol dye and stains neutral lipids such as triglycerides, diacylglycerols, and cholesteryl esters but not polar lipids (i.e., phospholipids, sphingolipids, and ceramides). Oil red-O staining demonstrated that MSCs accumulated lipid droplets in significantly high numbers when cells were cultured on E-Glc-PA/K-PA nanofiber network after 7 days of culturing (Figure 5A). On the other hand, the number of cells having lipid droplets and amount of total lipid droplets in K-Glc-PA/E-PA and E-PA/K-PA combinations and bare surface were very few compared to E-Glc-PA/K-PA group.

To better evaluate the differentiated cells, gene expression profiles of MSCs cultured on peptide nanofiber networks and on bare surfaces were analyzed. Expression levels of two widely used adipogenesis markers adiponectin (ADIPOQ) and fatty acid binding protein 4 (FABP4) were examined. On day 7, a remarkable increase in expression of this adipogenesis gene markers was observed for E-Glc-PA/K-PA compared to three other groups (Figure 5B). There was a slight increase in expression of both genes on K-Glc-PA/E-PA group compared to K-PA/E-PA and bare surface groups. However, this increase was much less than the one that was observed on E-Glc-PA/K-PA. In terms of gene expressions, the K-PA/E-PA and bare surface groups were comparable for both genes on both days. This result showed that the glucose has to be included in peptide nanofiber system in order to obtain this specific bioactivity. Overall mRNA expression results support the lipid droplet formations which were observed by oil red-O staining.

Oil red-O staining, adiponectin, and FABP4 mRNA expression analyses were also performed on day 11 in order to track the trend observed on day 7. The results were similar on both experimental days. Day 11 oil red-O staining underlines the importance of spatial organization of PA nanofibers. E-Glc-PA/K-PA and K-Glc-PA/E-PA had significantly different effects of on MSC adipogenesis (Figure S3). Like oil red-O, the trend in relative change of adiponectin and FABP4 gene expressions on day 7 and day 11 was also comparable to each other (Figure S5), where E-Glc-PA/K-PA group had a strikingly higher level of expression.

Determination of the Type of Fat Cells Which Were Differentiated from MSCs on E-Glc-PA/K-PA Nanofiber Scaffolds. There are mainly two types of different fat cells, white and brown fat cells. White adipocytes contain single or a few large lipid droplets that appear to comprise the majority of cell volume. The cytoplasm and nucleus of white adipocytes are found at the cell periphery. Brown adipocytes, on the other hand, are characterized by small and multilocular lipid droplets. The type of fat cells committed from MSCs on E-Glc-PA/K-PA nanofiber systems was further investigated by staining with Nile red and TO-PRO-3, and stained cells were imaged with a high resolution confocal microscope. Nile red is a hydrophobic and fluorescent dye that selectively stains the intracellular lipid droplets. TO-PRO-3 sensitively stains the double stranded nucleic acids and is widely used as a nuclear counterstain. Most of the cells cultured on the E-Glc-PA/K-PA nanofiber formed multilocular lipid droplets (Figure 6A). Multilocular lipid droplets were observed throughout the cytoplasm. Multilocular lipid droplets are a characteristic feature of brown fat cells, and other scaffolds did not induce high amount of lipid droplets (Figure S4).

Leptin and UCP1 are characteristic proteins of white and brown fat, respectively. Uncoupled protein 1 (UCP1), also called thermogenin, is a mitochondrial protein, which is uniquely expressed by the mitochondrion-rich brown adipocytes, and its function is to uncouple respiration from ATP synthesis. Leptin is an endocrine hormone, which is primarily produced by adipose tissue and is well-known as a satiety factor. Leptin is also determined as a marker gene to distinguish white fat cells and brown fat cells. Therefore, expression levels of leptin and UCP1 genes of MSCs cultured...
Figure 5. Determination of adipogenesis. (A) Oil red-O staining of cells cultured in maintenance medium for 7 days, on different networks, showing the lipid droplet accumulation. Red colors indicate lipid droplets inside cells. Scale bar is 100 μm and equal for all images. (B) Expression analyses of adipogenesis marker genes, adiponectin and FABP4, in mesenchymal stem cells cultured in maintenance medium for 7 and 11 days. The expression level of each gene was normalized against GAPDH and bare surface. Different letters denote significance at $p < 0.05$. 

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on PA nanofiber scaffolds were analyzed. Results revealed that while leptin expression levels were not significantly different from each other for all experimental groups, expression level of UCP1 on the E-Glc-PA/K-PA nanofibers was remarkably high compared to K-Glc-PA/E-PA system and the control groups (Figure 6B). Variation in leptin levels among the different experimental groups was not more than 0.7-fold change on both day 7 and day 11. The expression levels of UCP1 on K-Glc-PA/E-PA, K-PA/E-PA, and bare surface were comparatively in the same range on day 7 and day 11, compared to the bare surface. On the other hand, expression of UCP1 on E-Glc-PA/K-PA on day 7 was 61.8-fold more than the bare surface and was also more than all other groups. UCP1 expression of cells cultured on E-Glc-PA/K-PA further increased on day 11, with a prominent 114-fold change compared to bare surface level. UCP1 expression analysis itself is striking in showing the importance of spatial organization.

**DISCUSSION**

The bioactive PA molecules with opposite charges self-assemble into nanofiber networks. The self-assembly process is driven by hydrophobic collapse, electrostatic interactions, and β-sheet formation, and the nanofibers are stabilized by van der Waals and hydrophobic forces and hydrogen bonding. SEM and TEM imaging showed that PA molecules create ECM-like highly porous environment formed by nanofibers. Circular dichroism spectroscopy revealed that coassembly of PAs has β-sheet secondary structure due to the incorporation of β-sheet driving amino acids into the PA backbone.

Biocompatible PA nanofibers exhibited different capacities to alter cell adhesion and proliferation among different PA coassembly combinations. Higher number of cells adhered on nanofiber scaffolds compared to bare surface indicating that PA niche provides more adhesion epitopes than that of tissue culture plate. The number of adhered cells was significantly higher on glucose-containing nanofiber systems compared to E-PA/K-PA system, which shows the importance of glucose
residue for cell adhesion. On the other hand, cellular adhesion on E-Glc-PA/K-PA was significantly less than K-Glc-PA/E-PA group, although they have the same amount of bioactive functional groups and glucose. This difference could be due to the cooperative effect of glucose residue and lysine when placed in the same peptide backbone. Lysine residues create an adhesive environment for cells, and polylysine is widely used to modify glass or plastic surfaces to culture specific cell types that are less prone to adhesion.\textsuperscript{50,51} Glucose group is not present in K-PA/E-PA group, and it is conjugated to glutamic acid and adipogenic gene marker expression analyses with glucose group, although they have the same amount of bioactive groups of PAs in nanostructures further demonstrates that different spatial organizations of PA nanofibers trigger different cellular responses.

By revealing the multicellular lipid droplets, Nile red staining also indicated that E-Glc-PA/K-PA induces MSC differentiation into brown fat adipocytes, not to the white ones. This conclusion was further confirmed by brown and white fat specific gene expression analyses. Brown character of the adipocytes that differentiated from MSCs was measured by leptin and UCP1 gene expression levels. UCP1 is a mitochondrial protein that dissociates oxidative phosphorylation from energy production, leading to increased thermogenesis.\textsuperscript{33} Brown adipose tissue is specialized for thermogenic energy expenditure, and it maintains animal body temperature by increasing mitochondrial protein that dissociates oxidative phosphorylation from energy production, leading to increased thermogenesis.\textsuperscript{33} Brown adipose tissue is specialized for thermogenic energy expenditure, and it maintains animal body temperature by increased mitochondrial protein that dissociates oxidative phosphorylation from energy production, leading to increased thermogenesis.\textsuperscript{33}

**CONCLUSION**

Early cellular response, lipid droplet formation, and expression levels of adipogenesis marker genes demonstrated that the spatial organization of the functional groups in PA nanofibers determines cellular differentiation profiles of MSCs. In addition, this study introduced a simple, straightforward method to differentiate MSCs to mature adipocytes with brown adipocyte characteristics. As there is no need to use any other differentiation inducing agents, this scaffold eliminates the high cost efforts for obtaining brown fat cells from stem cells. This scaffold system not only can be used for brown fat formation/stem cell differentiation studies but also can be used to obtain brown fat cells from individual’s own stem cells with high yield, and these brown fat cells can be used as obesity fighting and thermogenesis agents in future. Overall, the present study shows the importance of spatial organization of bioactive groups of PAs in nanofiber systems and introduced a novel scaffold for brown fat cell production from MSCs.

**EXPERIMENTAL SECTION**

**Synthesis of Peptide Amphiphile Molecules.** The PA molecules were synthesized by standard SPPS (solid phase peptide synthesis) method. The PAs, except E-PA, were constructed on MBHA Rink amide resin, and E-PA [lauryl-VVAGE] was constructed on Wang resin preloaded with Fmoc-Glu(O’Bu). The resins were swelled in DCM (dichloromethane) for 30 min. Following resin swelling, DCM solvent was exchanged to DMF (dimethylformamide), in which all remaining reactions were carried out. All amino acid couplings, except Fmoc-Ser[β-D-Glc(OAc)\textsubscript{4}]-OH, were performed with 2 equiv of Fmoc protected amino acid, 1.95 equiv of HBTU, and 3 equiv of N,N-diisopropylethylamine (DIEA) in DMF. Coupling duration was at least 3 h but varied depending on the type of amino acid that was coupled. Fmoc-Ser[β-D-Glc(OAc)\textsubscript{4}]-OH coupling was performed with 1.2 equiv of amino acid, 1.1 equiv of HBTU, and 1.8 equiv of DIEA in DMF. Equivalences are based on the resin that was used for construction. Fmoc removal was performed with 20% piperidine/DMF solution for 20 min. After each coupling reaction, resin was treated with 10% acetic anhydride in DMF for 30 min to block any remaining free amino groups. Before each succeeding event, washing was performed by DMF, DCM, and DMF three times each subsequently. Cleavage of the peptides from resin was carried out with a mixture of trifluoroacetic acid (TFA)/triisopropylsilane (TIS)/water in the ratio of 95:2.5:2.5 for 2 h. Excess TFA was removed by rotary evaporation. The remaining viscous peptide solution was treated with ice-cold diethyl ether overnight at −20 °C. Ether decantation was performed after centrifugation at 4 °C and 8000 rpm for 15 min. After complete evaporation of diethyl ether via air drying, the resulting pellet was dissolved in ddH\textsubscript{2}O, sonicated for 30 min, freeze-dried at −80 °C, and lyophilized.

Deacetylation of glucose conjugated PAs, which are K-Glc-PA [lauryl-VVAGKS(β-D-Glc(OAc)\textsubscript{4})-Am] and E-Glc-PA [Lauryl-
Scanning Electron Microscopy. For SEM imaging, peptide hydrogels were prepared on cleaned silica wafers by mixing 10 mM PA solutions at 1:1 ratio except combinations including E-PA. PAs that were combined with E-PA were mixed at 2:1 ratio, where E-PA was 1 equiv. Obtained gels were incubated for 20 min at room temperature to allow gelation and then dehydrated. Dehydration was performed in 20, 40, 60, 80, and 100% ethanol solutions sequentially. Dehydrated gels were dried with a Tousimis Autosamdi-815B critical-point-drier to preserve the network structure. The dried samples were covered with Au/Pd at 5 nm thickness and visualized under high vacuum with a FEI Quanta 200 FEG scanning electron microscope equipped with ETD detector.

Transmission Electron Microscopy. TEM samples were prepared on a 200-mesh copper TEM grid. Nanostructures were formed as explained above, and these solutions were 10 times diluted. An amount of 30 μL of the solutions was dropped on a parafilm, which is a hydrophobic surface, and the grid was placed onto the droplet. After 5 min of incubation, grid was taken, excess PA solution was removed, and the sample was stained with 2 wt % uranyl acetate for 3 min. Immediately after 3 min, uranyl acetate was removed and the grid was rinsed with ddH2O and left for air drying. TEM images were acquired with FEI Tecnai G2 F30 TEM at 300 kV.

Cell Culture and Maintenance. Rat mesenchymal stem cells (MSCs) (Invitrogen) were cultured as monolayer cultures in maintenance medium, DMEM (Dulbecco’s modified Eagle medium) supplemented with 10% FBS and 1% penicillin-streptomycin, in tissue culture plates under standard conditions (at 37 °C under 5% CO2). Cells were seeded at 3000–5000 cells/cm² and passaged at 80% confluence by trypsin–EDTA (ethylene diamine tetraacetic acid) (0.025%). The culture medium was replaced every 3–4 days. In all experiments, cells between passages 6–9 were used. All cell culture reagents were purchased from Invitrogen.

Viability Assay: MTT Assay. Viability of MSCs seeded on nanoﬁber networks was quantiﬁed by MTT assay (Sigma, catalog no. TOX-1). In order to perform MTT assay, cells at a density of 1.2 × 10⁴ cells/cm² were seeded on PA coated wells and on bare wells of 96 well-plates. Then, plates were incubated under standard conditions for 12 h. After incubation period, medium was replaced with phenol red free medium containing (3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide) (MTT) reagent and cells were incubated for 4 h at standard cell culture conditions. Following the incubation period, plates were removed from the incubator, and the resulting formazan crystals, formed by viable cells through cleavage of tetrazolium ring, were dissolved in MTT solubilization solution [M-8910]. Spectrophotometric measurement was performed at a wavelength of 570 nm, and obtained results were converted into % viability against bare surface.

Cell Adhesion. Adhesion capacity of MSCs on PA coated and on bare surfaces was analyzed after 4 h of incubation. To perform adhesion test, growth medium of the ﬂask was replaced with adhesion medium, which is serum-free low glucose phenol red including DMEM supplemented with 3 wt % BSA and 0.05 wt % cycloheximide, and cells were incubated for 1 h at standard cell culture conditions. To detach cells from the ﬂask surface, trypsin-EDTA was added and incubated for 30 s. Later 6 × 10³ cells/cm² were seeded on PA coated wells and on bare wells and incubated under standard conditions for 5 h. After 5 h, the medium was removed by aspiration, the wells were washed with PBS, and cells were stained with 2 μM calcein AM.
(Invitrogen) in PBS for 40 min. Stained cells were imaged by an inverted fluorescent microscope and counted using ImageJ software. The experiment was carried out on a 96-well plate.

**Cell Proliferation.** Proliferation of cells was determined by cell proliferation ELISA, BrdU (colorimetric) (Roche) according to the manufacturer’s protocol. A 96-well plate was used for the proliferation assay. Briefly, 1.2 × 10⁴ cells/cm² were seeded on PA coated wells and on bare wells and incubated for 48 h under standard conditions. After 48 h, medium was replaced with 10 μM BrdU (bromodeoxyuridine) containing medium and cells were incubated under standard conditions for an additional 4 h. Cells were then fixed and stained by using labeled antibody against BrdU. Cells were washed three times with PBS, and substrate solution was added. After 20 min, color development was measured at a wavelength of 370 nm by using a microplate reader (Molecular Devices Spectramax M5) and subtracted from reference wavelength (492 nm) values.

**Oil Red-O Staining.** Oil red-O assay was performed to evaluate deposition of triglycerides and lipids. Nanofiber formation on 96-well plate was done as explained. Cells were seeded at 1.2 × 10⁴ cells/cm² density and incubated under standard conditions until the indicated experiment time points. At the indicated time points, medium was discarded by aspiration and cells were washed with 1x PBS and fixed in 10% formalin in 1x PBS. After 10 min, fixation solution was refreshed and the cells were incubated for an additional 1 h at room temperature. Later, wells were washed with ddH₂O and incubated in 60% isopropanol for 5 min at room temperature. After 5 min, isopropanol was discarded and plate was left open to air-dry. When the wells were completely dried, 100 μL oil red-O working solution per well was added and incubated for 10 min by gently shaking on a shaker. Unbound dyes were washed with ddH₂O 4 times. Finally, ddH₂O was added on wells and images were acquired under an inverted light microscope.

**Real-Time Gene Expression Analysis.** Gene expression profiles of differentiating MSCs were assessed by quantitative real time PCR (qRT-PCR) analyses. For total RNA extraction, cells were seeded at 1.2 × 10⁴ cells/cm² density on PA coated and bare well surfaces. For RNA extraction, cells were grown on 24-well plates. Total RNAs of cells were extracted by using TRIzol (Invitrogen) according to the manufacturer’s instructions. Concentration and purity of isolated RNAs were measured by Nanodrop 2000 (Thermoscientific). All samples were diluted to 50 ng/μL concentration prior to their use in qRT-PCR. Conversion of RNAs to cDNA and amplification of cDNAs were performed by using SuperScript III Platinum SYBR Green one-step qRT-PCR kit (Invitrogen) according to the manufacturer’s protocol. Reaction conditions were briefly as follows: 55 °C for 5 min, 95 °C for 5 min, 40 cycles of 95 °C for 15 s, 60 °C (where X is the annealing temperature of the primers; see Table S2) for 30 s and 40 °C for 1 min, followed by a melting curve analysis in order to confirm product specificity. The reaction efficiencies for each primer were determined by plotting a standard curve with 10-fold dilutions of the total RNA. Primary gene expression levels were normalized to GAPDH (a housekeeping gene), and comparative Ct method (Pfaffl method) was used to analyze results.

**Nile Red/TO-PRO-3 One Step Staining.** Nile red/TO-PRO-3 staining was performed to visualize cells under confocal microscope for high resolution analyses of lipid droplets and nucleus. For Nile red/TO-PRO-3 staining, nanofiber formation was carried out with the same method on glass coverslips that are placed in 24-well plates. Cells were seeded at 1.2 × 10⁴ cells/cm² density on PA coated and bare glass surfaces and incubated under standard conditions. At the indicated time points, cell medium was discarded, cells were washed with 1× PBS twice, and cells were fixed with 4% paraformaldehyde for 20 min. Later, fixative solution was removed, cells were washed with 1× PBS, and 250 μL per well Nile red/TO-PRO-3 solution (100 ng/mL Nile red and 1:1000 dilution of TO-PRO-3 in 1X PBS) was added. After a 20 min incubation period, Nile red/TO-PRO-3 solution was discarded and cells were washed with ddH₂O. Coverslips were then removed from wells, mounted with Prolong gold antifade mounting reagent (Invitrogen), and imaged under Carl Zeiss LSM510 confocal microscope.

**REFERENCES**


