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Microfluidic assay for simultaneous culture of multiple cell types on surfaces or within hydrogels

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Abstract

This protocol describes a simple but robust microfluidic assay combining three-dimensional (3D) and two-dimensional (2D) cell culture. The microfluidic platform comprises hydrogel incorporating chambers between surface-accessible microchannels. Using this platform, well-defined biochemical and biophysical stimuli can be applied to multiple cell types interacting over distances of <1mm, thereby replicating many aspects of the in vivo microenvironment. Capabilities exist for time-dependent manipulation of flows and concentration gradients as well as high-resolution real-time imaging for observing spatial-temporal single cell behavior, cell-cell communication, cell-matrix interactions and cell population dynamics. These heterotypic cell type assays can be used to study cell survival, proliferation, migration, morphogenesis and differentiation under controlled conditions. Applications include the study of previously unexplored cellular interactions, and have already provided new insights into how biochemical and biophysical factors regulate interactions between populations of different cell types. It takes 3 days to fabricate the system and experiments can run for up to several weeks.

INTRODUCTION

Development of the protocol

Three-dimensional (3D) models of cellular microenvironments mimicking the functions of intact tissues are recognized to have numerous advantages over well-plate based two-dimensional (2D) cell culture approaches. A robust in vitro assay enabling 3D cell culture is beneficial for cell-based drug and toxicity screening to closely mimic the

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AUTHOR CONTRIBUTIONS

Y.S. and S.H. were equally contributed to this protocol, designing and carrying out the experiments and writing the paper. J.S.J, K.Y., I.K.Z., R.S. and S.C. designed and carried the experiments. S.C. is responsible for all the experiments described in this article and preparation of the paper. R.D.K. is responsible for providing guidance for the experiments and for editing the paper.
microenvironment of living tissues, and to successfully isolate and define specific contributions of single and/or multiple factors to the overall processes of cellular morphogenesis including cell migration, angiogenesis, and cancer metastasis\textsuperscript{2-5}. Cells cultured using traditional 2D tissue culture methods are morphologically different from cells in 3D environments, and moreover, show different gene expression levels of a variety of proteins\textsuperscript{2,3} compared to their native environment \textit{in vivo}. Approaches that employ cells suspended in 3D gels have also been used, but these typically lack the ability to culture in defined locations cells of different types\textsuperscript{6}. For example, Transwell\textsuperscript{®} systems enable the study of 3D chemotactic responses of the cells cultured on filters or in gels placed on the filters, but lack the ability to image at high resolution in real time, or to control many aspects of the local cellular microenvironments. Other 3D assays suspending cells in a gel or plating cells on beads within a gel provide a simple and effective means for monitoring cellular morphogenesis within an extracellular matrix (ECM)\textsuperscript{7,8}. However, sprouts or migration of cells from a solid bead differ from cellular morphogenesis \textit{in vivo}. It is also difficult to produce a well-controlled microenvironment with similar dimensions to tissue structures \textit{in vivo}.

Recognizing the need to better replicate the \textit{in vivo} environment, we developed a novel hydrogel-incorporating microfluidic cell culture assay, which facilitates the interaction of cells in 3D ECM scaffolds, and thus, better captures the complex behaviors that occur within the \textit{in vivo} microenvironment compared to 2D \textit{in vitro} assays\textsuperscript{9}. Similar microfluidic platform protocols designed for neuroscience research\textsuperscript{10}, gelatin-based cellular interaction\textsuperscript{11} and bacterial chemotaxis\textsuperscript{12} have enabled integration of complex environmental factors and high resolution \textit{in situ} imaging. Our system is a multipurpose platform for local containment of ECM-mimicking hydrogels to supply precisely controlled conditions to the cells cultured on and/or inside. It also provides the flexibility necessary to study 3D cellular morphogenesis\textsuperscript{13}, 3D homotypic/heterotypic cell-cell and cell-ECM interactions\textsuperscript{9,14,15}, and 3D responses to biochemical gradients\textsuperscript{16-18}, interstitial flow\textsuperscript{15}, diffusion of soluble factors, and various mechanical stimuli\textsuperscript{19} with the same batch of cells at the same time. Furthermore, due to the smaller length scales compared to traditional cultures it requires lower volumes of reagents and media.

**Applications of the method**

The microfluidic platform described in this protocol has been widely applied to 3D cell culture in various applications, including 3D angiogenic responses under the gradient of growth factors introduced in the media\textsuperscript{13,16-18} or secreted by co-cultured tumor cells\textsuperscript{14}. The angiogenic structure of networks of endothelial cells in the hydrogel was confirmed as 3D and regulated by surface treatment of the microfluidic channel\textsuperscript{13}. We also found that primary hepatocytes cultured on the hydrogel under interstitial flow formed 3D tissues, enabling detoxification and regulating capillary morphogenesis of co-cultured endothelial cells\textsuperscript{20}. We investigated the endothelium-dependent migration of smooth muscle cells co-cultured with ECs\textsuperscript{21}, 3D axonal responses under various growth factor gradients\textsuperscript{22}, and 3D cancer metastatic processes\textsuperscript{14,23}. Cancer cells invaded and migrated in 3D into the hydrogel and finally intravasated through the endothelial monolayer cultured in a microfluidic channel\textsuperscript{14}. We modified the microfluidic design to create a more stable concentration
gradient that was better able to represent the metabolic activity of cells\textsuperscript{18,24} and also successfully manufactured a platform of similar design suitable for high volume manufacture using hot-embossed plastics\textsuperscript{25}. The structure, results and possible applications of the platform technology have been described in two recent reviews\textsuperscript{9,26}. In this protocol, we describe the basic procedure for using a hydrogel-incorporating microfluidic assay using angiogenesis studies as an example. Based on the protocol, the readers can easily integrate conventional soft lithography technology\textsuperscript{27} to explore a wide variety of applications with small modifications in cell seeding and culturing procedures.

**Experimental design**

Microfluidic systems are fabricated in PDMS using standard soft lithographic methods. Other materials such as standard tissue culture plastics can also be used in combination with either hot-embossing or injection molding methods. It is preferable if all features are of uniform depth to enable simple single-stage fabrication. Drilling or coring holes through the full thickness of the device provides direct access to the gel-containing and media channels, and bonding a coverslip to the side with imprinted features produces closed channels. The entire process of microfluidic cell culture including fabrication, cell culture and analysis is explained here. From various microfluidic device designs, we chose to focus on the one most widely used for 3D cell culture assay, which is composed of 4 gel regions and 3 channels, each of which is individually accessible (Fig.1). The cells are usually cultured in the center channel and the other two side channels act as control channel and condition channel, respectively. The interior of the channels is coated with ECM proteins to promote cell adhesion. In particular, the described method of coating with Poly-D-lysine Hydrobromide (PDL) has been confirmed to promote 3D capillary morphogenesis into the hydrogel\textsuperscript{13}. Finally, we illustrate the method for introducing a hydrogel solution (type I collagen gel as the prototypical example) controlling gel stiffness, seeding cells inside the gel region and immunostaining the cells after the experiment using a method that is specially optimized for the micro-scale. In addition to general instructions we give specific details for how to evaluate the angiogenic response to a gradient in vascular endothelial growth factor (VEGF).

The part of the protocol describing the manufacture of the microfluidic device can be divided into the master fabrication (photolithography; steps 1-8) and mold fabrication (soft lithography stages). The first stage only needs to be performed once per master type, as a master can be used multiple times in the subsequent stages. For the general biology community, the photolithography stage may require equipment not available in the typically stocked lab plus assistance from someone experienced in microfluidics or microfabrication. Further details can also be acquired from other protocols (see procedure 1-7\textsuperscript{12} & procedure 1-5\textsuperscript{27}). However, the soft lithography stage only requires equipment (vacuum desiccator and dry oven) that is commonly available in a biology laboratory and is easier to follow without specialist knowledge. Further details about how to perform soft lithography can be found in other protocols (see procedure 8-12\textsuperscript{12} & procedure 6-11\textsuperscript{27}).
Limitations

Whereas the small dimensions within the device are essential for realistic cell-cell signaling to occur, it also imposes certain inherent limitations. Typical biochemical assays could be difficult to perform due to the small cell number (typically <1000 cells), so methods of interrogating biological function are generally restricted to antibody staining for the distribution, concentration or activation of various intra- or extracellular biomolecules. Channel structure has been modified in similar devices to increase cell number to enable conventional biochemical analysis of ELISA or RT-PCR. Loading the system with hydrogel, media and cells requires training, but can usually be mastered in a day. Imaging is typically performed by conventional microscopy including confocal microscopy, but other methods such as transmission EM are also possible.

MATERIALS

REAGENTS

Photoresist SU-8 2050 (for photolithography, Microchem)!

CAUTION Wear protective goggles and gloves and suitable protective clothing when working with photoresist.

Photoresists developer (for photolithography, PGMEA, Microchem)

Isopropyl alcohol (for photolithography, IPA)

Sylgard 184 silicone elastomer base and curing agent (for soft lithography, Dow Corning)

Sterile 1X phosphate buffered saline (PBS), pH 7.4 (Thermo Scientific, cat. no. SH30256.01)

Deionized water (Direct-Q® 3; water purifying system, Millipore)

Ethanol: 70% in sterile deionized water

Methanol

Poly-D-lysine (PDL) hydrobromide solution (see REAGENT SETUP) (Sigma-Aldrich, cat. no. P7886)

Trypsin 0.25% (1X) with EDTA (Gibco, cat. no. 25200-056)

Collagen type I, rat tail (BD Bioscience, cat. no. 40236)

10X phosphate buffered saline (PBS) with phenol red (see REAGENT SETUP) (Gibco, cat. no. 14080-055)

0.5N NaOH solution (see REAGENT SETUP) (Sigma-Aldrich, cat. no. 655104)

Human microvascular endothelial cells (hMVEC) culture medium, EGM-2MV Bulletkit (Lonza, cat. no. CC-3202)

Human vascular endothelial growth factor (VEGF; R&D systems, cat. no. 293-VE)
Trypan blue stain 0.4% (Gibco, cat. no. 15250-061)！CAUTION Wear suitable protective clothing and gloves when working with trypan blue.

DAPI, dilactate (Sigma-Aldrich, cat. no. D9564)

Rhodamine Phalloidin (Molecular probe, cat. no. R415)

Paraformaldehyde (PFA) (USB-Affymetrix, cat. no. 199431 LT)

Triton X-100 solution (see REAGENT SETUP) (Sigma-Aldrich, cat. no. T8787)

**EQUIPMENT**

4 inch Silicon wafers (for photolithography)

UV mask aligner (for photolithography, SUSS MicroTec)

Photo mask (for photolithography, printed transparent film)

Spin coater (for photolithography, spin coating SU-8 photoresists, Dong Ah Trade)

Nitrogen gas (for photolithography, removal dust particles adhering on silicon wafers and glass cover slips)

Hotplates (for photolithography, baking SU-8 films, DongSeo Scientific)

Optical microscope (for characterizing patterns on masters at the microscale, SOMETECH)

Sonicator (for cleaning, Branson)

Alpha step (for photolithography, profiling thickness of SU-8 pattern, KLA-Tencor, ASIQ)

Vacuum desiccator (for soft lithography, removal bubbles in PDMS solution, GAST)

Dry Oven (for soft lithography, drying sterile cured PDMS and glass cover slips, DongSeo Scientific)

Glass cover slips (24×24 mm, Paul Marienfeld)

Plasma cleaner (for surface modification, Femto Science)

Disposable Biopsy Punches (4mm & 1mm diameter, Miltex)

Autoclave (Daihan Scientific, DS-PAC80.)

Incubator at 37°, 5% CO₂, MCO-15AC (Sanyo)

Water bath at 37° (Jeio Tech)

Inverted microscope (for imaging cells, Olympus)

Syringe filters (Pall, cat. no. PN4612)
REAGENT SETUP

Poly-D-lysine (PDL) hydrobromide solution—Dissolve PDL (Sigma-Aldrich, cat. no. P7886) in sterile deionized water (1 mg/ml) using a stirrer at room temperature (23°C). Sterilize the solution with a 250 ml bottle top 0.2 μ filter. Dispense the sterile solution into 1 ml each and store at –20°C for future use.

Triton X-100 solution—Dilute 0.1% of Triton X-100 (Sigma-Aldrich, cat. no. T8787) (v/v) in distilled deionized water (DDW) and store at room temperature (23°C).

10X phosphate buffered saline (PBS) with phenol red—Dissolve 63.6 mg of Phenol red sodium salt (Sigma-Aldrich, cat. no. 114537-5G,) in 400ml of 10xPBS (Gibco, cat. no. 14080-055) using a vortex generator. Sterilize the solution using a 250 ml bottle-top 0.2 μ filter and store at room temperature (23°C).

0.5N NaOH solution—Dissolve 5 g of NaOH tablets in 230 ml DDW by stirring. ▲ CRITICAL STEP Add no more than 1g of NaOH tablets into sterile deionized water each time. If NaOH tablets are added into sterile deionized water at once, the solution will become hot. Add sterile deionized water up to 250 ml into the solution. Sterilize the solution using a 250 ml bottle-top filter (0.2 μ) and store at room temperature (23°C).

PROCEDUREI

Fabrication of the master (photolithography) ● TIMING 3h

1| Dispense about 4 ml of SU-8 100 on a cleaned and dehydrated 4 inch silicon wafer.

2| Coat the dispensed SU-8 uniformly on the wafer using a spinner by ramping up to 500 rpm at 100 rpm/sec for 10 seconds (for 150 μ thick patterns).

3| Ramp to 2,000 rpm at an acceleration of 300 rpm/second for 30 seconds (for 150 μ thick patterns).

4| Pre-bake the wafer for 20 minutes at 65 ° and for 50 minutes at 95 ° on a hotplate (for 150 μ thick patterns). ! CRITICAL STEP Keep the coated wafers level (Fig.2a).

5| Use a transparency photomask printed at 20,000 dpi to produce the master by exposing the wafer to an appropriate dose of UV light as per Microchem’s protocol (Fig.2b).

6| After exposure, post-exposure bake the wafer for 1 minute at 65 ° and for 12 min at 95 ° on the hotplate (for 150 μ thick patterns).

7| Develop for 15 minutes in Photoresist developer (for > 150 μ thick patterns) (Fig.2c).

8| Rinse the substrate briefly with isopropyl alcohol (IPA) and gently dry with pressurized nitrogen gas.

■ PAUSE POINT The master can be stored until further use.
Fabrication of PDMS molds (soft lithography) ● TIMING 7 h

9| Tape (using scotch tape) the master securely on a petri dish (100 mm in diameter), in order to prevent the PDMS solution from percolating down through the wafer.

10| Blow dust particles off the wafer using pressurized nitrogen gas.

11| Take a disposable plastic cup and fill it with Sylgard 184 silicone elastomer base and the curing agent in a 10:1 weight ratio and mix the solution using metal spatula.

12| Pour the mixed solution into the petri dish containing the master to the desired thickness (4-5 mm) and degas to remove air bubbles trapped in patterns.

13| Cure the solution in an oven at 80 ° for 1 hour 30 min (Fig.2d). 

14| Use a scalpel to cut out the cured PDMS and remove PDMS from the wafer (Fig.2e).

15| Punch holes through the reservoir pattern at appropriate points using a dermal biopsy punch (Disposable Biopsy Punches 4mm). To make inlets for hydrogel filling, punch holes using another punch (Disposable Biopsy Punches 1mm) (Fig.2f).

16| Apply and detach scotch tapes on the PDMS to remove small particles, which can cause contamination and leakage of the bonded device.

17| Place the devices in a glass beaker with deionized water and cover the flask with aluminum foil. Using an autoclave, sterilize them at 120 ° for 20 min (a wet autoclave). Make sure that the devices are not stuck to each other for effective sterilization.

18| Place the devices in a clean box using a pair of sterilized tweezers (do this step in a sterile laminar flow hood), the channel side up and sterilize again using an autoclave at 120 ° for 20 min (a dry autoclave).

! CRITICAL STEP Both wet and dry autoclaves are recommended for complete sterilization.

19| Dry the box containing sterile devices in an oven at 80 ° for at least 4 hours.

■ PAUSE POINT These devices can be stored at room temperature 23 ° for up to a month before bonding to glass coverslips.

Cleaning glass coverslips ● TIMING 5 h

20| Using pressurized nitrogen gas, blow off particles adhering to the glass coverslips.

21| Autoclave the glass coverslips in a clean pipette tip box at 120 ° for 20 minutes and dry them in an oven at 80 ° for more than 4 hours. ! CAUTION All steps after sterilization, the autoclaved coverslips and devices should be handled in sterile environment (in laminar flow hood with sterilized gloves).

Device assembly ● TIMING 10 min
22| Place the devices and glass coverslips in a plasma cleaner, the channel side up (Fig.3a).

23| Turn on a vacuum pump and wait until pressure drops below 600 mTorr. It takes about 30 seconds. In this bonding step, the working conditions may vary depending on specifications of plasma cleaner and vacuum pump. Working conditions (vacuum time and working pressure) should be optimized.

24| Turn on plasma to treat the surfaces of PDMS and glass coverslips for 50 seconds. A pale violet color should be observed in the vacuum chamber during this process.

25| Vent the chamber immediately.

26| Bond the device to the glass coverslip and press the device softly with a sterile pair of tweezers, avoiding pressure on the channel area (Fig.3b).

TROUBLESHOOTING

PDL coating of the microchannels ● TIMING 5h

27| Fill the channels with 60 μl of surface coating solution (Fig.3c). As a surface coating solution, various polymer (PDL, and Poly-L-lysine etc.) and ECMs (fibronectin, collagen, and matrigel etc.) can be used according to desired cell types. Table 1 summarizes available surface coating solutions for various cell types. In example application of angiogenesis with type I collagen as hydrogel, PDL (1 mg/ml) coated channel surface is preferred for endothelial cell culture, which can prevent the detachment of collagen gel, resulting in 3D capillary morphogenesis into the type 1 collagen13.

28| Put the devices in an incubator for > 4 hours.

29| Remove the PDL solution in the channels thoroughly using an aspirator.

30| Fill the channels with 150 μl of sterile deionized water, and then wash more than 3 times with a micropipette. Repeat this washing process twice. Excess PDL molecules can cause cell damage. ! CRITICAL STEP This is different from a conventional surface coating protocol. Do not wash with PBS solution as this makes white stains in the microfluidic channel and interrupts hydrogel filling in the desired position.

31| Aspirate all microfluidic channels completely (Fig.3d).

Surface hydrophobicity restoration ● TIMING 1 day

32| Place the coated devices in a sterile dish and place at 80 ° in an oven for at least 24 hours, but no longer than 72 hours. It takes over 24 hours to restore hydrophobicity of the bonded surfaces. ▲ CRITICAL STEP Surface modification for the PDMS bonding (see the Device assembly steps) may lead to unintended breakage of optimal surface tension during the gel filling step. Therefore, hydrophobicity of the surfaces should be obtained.
Collagen gel solution preparation  ●  TIMING 5 min

33] To make 200μl of 2.0mg/ml, prepare 10X phosphate buffered saline (PBS) with phenol red, 0.5N NaOH solution, Collagen type I, and deionized water. Keep all solutions on ice. [Note that cells can also be added to the collagen gel solution for the experiments requiring 3D cell seeding or co-culture with another cell type.]

34] Calculate the exact amount of collagen type I solution needed based on the collagen solution stock density. 400μg of collagen is needed to make 200μl of 2.0mg/ml density collagen type I gel. The protocol described below is, as an example, for pH 7.4 collagen solution using one common stock density of 3.67mg/ml, but any density in the range of 3mg/ml to 4mg/ml may be used.

35] Add fixed amount (20μl) of 10X phosphate buffered saline (PBS) with phenol red solution into a 1ml tube.

36] Add 7.9μl of 0.5N NaOH solution into the same tube. The amount of NaOH solution can be adjusted according to the desired pH. ▲ CRITICAL STEP Be careful not to leave any residue in the pipet tip as slight differences in amount of 0.5N NaOH solution may alter the pH.

37] Add 109μl [3.67mg: 1ml = 400μg: x] of type 1 collagen stock solution and carefully swirl and pipet up and down the entire mixture until there are no variations of color in the gel. From the point when the gel seems to be uniform, thoroughly mix for about 20% time longer than the duration required to mix the solution. ▲ CRITICAL STEP Because of high viscosity of collagen, it is easy to create undesirable bubbles while mixing. Mix the solution with special care to avoid creating bubbles.

38] Add deionized water (option A, in the example application of angiogenesis) or cell suspension and deionized water (option B, when cells need to embedded within hydrogen) to the solution. The amount of water should be added to meet total amount (200μl).

(A) Collagen gel solution without embedding cells

(i) Add 63.1μl deionized water to the solution to meet the total amount. Mix thoroughly by swirling around the pipet tip and pipetting the solution in and out multiple times.

(B) Collagen gel solution embedding cells

(i) Make cell suspension with a cell density of 10 times that required on seeding.

(ii) Place the cell suspension in ice to cool down for 5 min. ▲ CRITICAL STEP This step is required for avoiding pre-polymerization of collagen gel suspension

(iii) Add 20μl of pre-chilled cell suspension to the solution and stir it gently using a large pipette tip. ▲ CRITICAL STEP Use a large pipette tip to reduce shear stress which will have an effect on cell function and/or viability.
(iv) Add 43.1ul deionized water to the cell embedded solution to create the total required volume (200ul) and carefully mix. ▲ CRITICAL STEP Use a large pipette tip to prevent cell damage.

39) Check the pH of the media is correct using a pH meter. If you follow the above process, the pH of the resultant collagen gel should be about 7.4. The color of the collagen solution can be a good indicator for the initial pH (Fig.4a). In addition to using collagen type 1 as scaffold, various hydrogels (matrigel, hyaluronic acid(HA), and hydrogel mixture etc.) can be used.

Collagen gel filling ● TIMING 40 min

40) Prepare a humid chamber and put it in a CO₂ incubator to keep warm before gel filling.

41) Gently and slowly fill the collagen gel solution into all four hydrogel scaffold regions with a 10ul micropipette. Collagen gel solution is easily fixed in hydrogel scaffold regions because of the surface tension caused by posts in the hydrogel region (Fig.3e, Fig.4c&d).

?TROUBLESHOOTING

42) Place the gel-filled device in the prepared humid chamber with the channel side facing up.

43) Put the humid chamber in a CO₂ incubator for 30 minutes to allow the collagen to gel (Fig.4b).

44) After incubation, place the device in a petri dish and fill medium in all channels.

45) Place the device in a CO₂ incubator.

■ PAUSE POINT If cells are not already seeded, the device can be stored in a CO₂ incubator for 2~3 days before cell seeding.

Cell suspension preparation ● TIMING 10 min

46) Remove media from cells to be used in the microfluidic device and wash cells once with PBS. Subsequently add warm trypsin 0.25% with EDTA solution to a cell culture dish that is about 70% confluent with hMVECs and incubate for 5 minutes. [Note that other cell types could be used in place of the hMVEC cells, depending on the nature of the experiment.]

47) Add 10 times volume of media containing FBS to dilute trypsin.

48) Centrifuge at 200×g for 5 minutes to obtain hMVEC pellet and remove supernatant.

49) Add appropriate amount of medium to obtain the proper density of hMVECs for seeding. For hMVECs, 2 million cells /ml is generally suitable.
hMVEC loading and culture in the microfluidic platform ● TIMING 1 h

50] Aspirate media from all reservoirs of the device and add 50–60 μl of the hMVEC suspension in one reservoir of the cell channel (Fig.5a&c) (the upper middle reservoir). The pressure difference between cell channel (middle) and condition / control channels (right / left channels) enables interstitial flow and the hMVECs are consequently more likely to connect and adhere to the side of the scaffold (Fig.5b). Cells can be seeded into one or both side channels as well as in the center channel. Since the channels are independent and separately accessible, different cell types can be introduced for various applications (HMVECs with cancer cells, cancer spheroids, SMC, and neutrophil etc.). For most of cell types, 50–60 μl of cell suspension should be applied to the reservoirs. The optimal concentration of cell suspension lies between 0.5 million cells/ml and 5 million cells/ml.

51] Put the device in a CO₂ incubator for 30 minutes to allow hMVECs to attach to the surface of the cell channel and form a monolayer on the side of scaffold. Incubation time can be adjusted depending on cell types, from 30 minutes to several hours.

52] After incubation, aspirate medium from the cell channel carefully. ▲ CRITICAL STEP Aspirate medium carefully by placing aspiration tip at the outside edge of reservoir to prevent cell detachment.

53] Add 50-60 μl of medium and place droplets in all six reservoirs (100-120 μl per channel). ▲ CRITICAL STEP The amount of media added to each reservoir should be the same (by pipette) to minimize initial fluctuations across the collagen gel. Such initial fluctuations may result from small differences between reservoirs in the volume of media added, and from device handling, but the induced perturbations generally stabilize in about 30 mins.

54] Place device in a CO₂ incubator (Fig.5d) for one day. In one day, seeded hMVECs should form a confluent endothelial monolayer (Fig.5b&e).

? TROUBLESHOOTING

Chemoattractant treatment and media exchange ● TIMING 10 min

55] Remove media from all six reservoirs and fill 100-120 μl of fresh media in the cell channel and control channel. ▲ CRITICAL STEP The total amount of fresh media applied to two reservoirs connected by one channel should be more than 100 μl to avoid cell death due to evaporation of media. The amount of media in all reservoirs should be the same (by pipette) to avoid initial fluctuation.

56] Add the same volume of media with chemoattractants, chemorepelsents or growth factor in the condition channel. In case of human vascular endothelial growth factor (VEGF), fill the EGM2mv media supplemented with 50ng ml⁻¹ recombinant VGEF gradient across the gel scaffold. Culture cells for a further 24h.

57] Replace media daily by aspirating out media in the channels and adding 100-120 μl of fresh media. The gradient reaches a steady state after about 30 min for a protein of 40 kD
molecular weight (VEGF) but will dissipate over time if not replenished by either another media change or by introducing a steady, slow flow through the channels by peristaltic or syringe pump. Without slow perfusion, linearity can be maintained for 24 hours and easily restored by daily replacement of media, which enables multiple experiments with the same batch of cells. ▲ CRITICAL STEP The time to establish a gradient is determined by the diffusion coefficient of the applied protein in the hydrogel, and can also be affected by matrix binding. Slope can be visualized by a fluorescent molecule (e.g., dextran) and also simulated by conventional CFD software (Fig. 6). After applying a chemical gradient, cellular responses can be monitored and quantified daily by conventional microscopy (Fig. 6c&d). Media or cells in the channel can be collected for biochemical or PCR analysis.

Cell staining

58] At the desired cell culture endpoint, remove the remaining cell culture media in all reservoirs by vacuum aspiration. ▲ CRITICAL STEP when aspirating media or other solutions from the reservoirs, aspirate media carefully by placing aspiration tip at the outside edge of reservoirs to prevent cells from detaching.

59] Rinse the channels of the device with PBS. Fill 90–100 μl PBS in one side of the reservoirs, then let it flow along the channels and check to ensure that it also filled the opposite reservoirs. Remove PBS in all reservoirs of the device. Repeat this step twice.

60] Add 50–60ul of 4% PFA in all reservoirs and leave it to stand for 15 min for cell fixation. ! CAUTION Wear suitable protective clothing and gloves when working with 4% PFA.

61] After 15 min of incubation, remove 4% PFA from each reservoir using a 1000 μl pipette. ! CAUTION Properly dispose of the PFA.

62] Wash the channels twice with PBS as described in step (60). ! CAUTION The first wash of PBS should be disposed into the PFA waste bottle.

63] Add 50–60ul of 0.1% Triton-X in each reservoir for solubilization and permeabilization of cell membranes. Let incubate on bench top for 5min.

64] Aspirate 0.1% Trion-X and wash twice with PBS as in step (60).

■ PAUSE POINT The device filled with PBS can be stored for 1-2 days at 4° in the refrigerator before fluorescent staining.

65] If fluorescent staining, follow option A. If using antibodies, follow option B. These methods can be combined.

(A) Fluorescent staining ● TIMING 2 h

(i) Dilute Rhodamine Phalloidin in the ratio of 1:40 and DAPI in the ration of 1:1000 with PBS to make a working concentration of fluorescent staining solution mixture. ▲
**CRITICAL STEP** Do this step and flowing steps in the darkroom and wrap all dishes and tubes with aluminum foil to protect them from light.

(ii) Remove PBS using a device with vacuum aspiration.

(iii) Add 50–60 μl of DAPI/ Rhodamine Phalloidin fluorescent staining solution mixture in each reservoir. **CRITICAL STEP** Immediately cover the individual dish with aluminum foil to protect it from light.

(iv) Maintain the device at room temperature for 1 hour.

(v) Aspirate remaining DAPI/ Phodamine Phalloidin fluorescent staining solution and rinse the channels with PBS as in step (59).

(vi) Wrap all dishes with foil and store at 4° until imaging. Imaging should be performed within the next 2–3 days after staining to ensure optimal fluorescent intensity signal.

(B) Immunofluorescent staining ● TIMING 2–3 d

(i) Remove PBS from device. Then add 40–50ul of Block Ace in each reservoir and incubate on bench top at room temperature for 1 hour.

(ii) Remove Block Ace from each reservoir and rinse the channels with PBS as in step (59) **PAUSE POINT** The device filled with PBS can be stored for 2–3 days at 4° in refrigerator until antibody staining.

(iii) Dilute the antibodies in PBS to the desired working concentrations for primary antibody incubation.

(iv) Remove PBS from the channels of the device with vacuum aspiration and add 50–60ul of primary antibody solution in each reservoir.

(v) Place the device at 4° in the refrigerator for >6 hr.

(vi) Aspirate the remaining primary antibody solution and rinse the channels with PBS as in step (59).

(vii) Fill PBS in all channels and place the device on a shaker in a a 4° refrigerator overnight.

(viii) Prepare a working concentration of secondary antibody solution in a conical tube by diluting in PBS. **CRITICAL STEP** Do this step and following steps in the darkroom and wrap all dishes and tubes with aluminum foil to protect them from light.

(ix) Remove PBS from device with vacuum aspiration and add 50–60ul of secondary antibody solution.

(x) Maintain the device at 4° in the refrigerator for >6 hr.
(xi) Aspirate remaining secondary antibody solution and rinse the channels with PBS as in step (59).

(xii) Fill the channels of device with PBS and place on a shaker in a 4°C refrigerator for overnight then store device at 4°C until imaging.

■ PAUSE POINT In this cell staining step, the incubating time with staining solution may vary depending on the cell numbers and amount of cells in hydrogel. Working time should be optimized.

● TIMING

Steps 1-8, Fabrication of the master: 3h
Steps 9-19, Fabrication of PDMS molds: 7h
Steps 20-21, Cleaning glass coverslips: 5h
Steps 22-26, Device assembly: 10 min
Steps 27-31, PDL coating of the microchannels: 5h
Step 32, Surface hydrophobicity restoration: 1 day
Steps 33-39, Collagen gel solution preparation: 5 min
Steps 40-45, Collagen gel filling: 40 min
Steps 46-49, Cell suspension preparation: 10 min
Steps 50-54, hMVEC loading and culture in the microfluidic platform: 1 h
Steps 55-57, Chemoattractant treatment and media exchange: 10 min
Steps 58-64, Cell staining: 2h or 2-3 d

?TROUBLESHOOTING

Troubleshooting advice can be found in Table 2.

ANTICIPATED RESULTS

The described platform allows for a large number of different experimental cell culture designs depending on which channels or gel regions contain cells, the types of cells used, and the nature of biochemical gradients or flow that are imposed. We first focus on the results obtained for the specific case described above, that of endothelial cells being seeded in the center channel with no cells seeded elsewhere in the system (either the side channels or the gel regions), and in the case with a gradient of growth factor applied on one side to induce angiogenesis. Fig.7 shows successful culturing of hMVEC in the cell channel and a 3D sprouting angiogenic response in the direction of increased VEGF concentration.
right side). Cells can be observed to progressively invade 3D into the type 1 collagen hydrogel towards the right side where a higher VEGF concentration has been introduced. Sprouting of the endothelial monolayer is observed only on the right side of the center channel, leaving the left side (no added VEGF) stable. Using confocal microscopy, the monolayer has been demonstrated to be confluent over the face of the gel and the 3D nature of the capillaries has been confirmed. Other cell types exhibit a variety of responses when seeded in the center channel and put in contact with a collagen hydrogel (Fig. 8). Gradient of growth factors or chemoattractants can regulate both the morphology and the degree of cellular morphogenesis into the collagen hydrogel. Various experimental modes of cell seeding and morphogenesis in hydrogel incorporating microfluidic assay are listed in Fig. 9.

When hMVEC are seeded in the center channel and another cell type (e.g., cancer cells, 10T 1/2) is seeded and cultured in the side channel, paracrine interactions occur across the ~1 mm hydrogel region. Various responses can be observed. In some cases, cancer cells induce a sprouting response in the endothelial monolayer, similar to that found with added VEGF. In other cases, the added cells migrate through the gel and interact with the other cells or monolayer inducing cellular morphogenesis in some instances and in others suppressing it (Fig. 10a). The microfluidic cell culture assay also allows the study of 3D cell-cell interactions and cell-ECM interactions, especially for the evaluation of new synthesized biodegradable polymers (hyaluronic acid, synthetic peptides, etc.) or other ECM materials (Matrigel, fibrin etc.). Co-cultures of cancer cells-endothelial cells (Fig. 10b&d), hepatocyte-endothelial cells, and smooth muscle cells (SMCs)-endothelial cells (Fig. 10c) have already been realized, to enable study of cellular morphogenesis and interactions. We proved that the assay can generate well-defined biochemical and biophysical stimuli to multiple cell types interacting with each other, thereby replicating many aspects of the in vivo microenvironment with significant implications for drug discovery and artificial organ systems in vitro and with time and cost benefits for drug screening.

Acknowledgments

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REFERENCES


Figure 1.
(a) Schematic and (b) photograph of the microfluidic cell culture assay incorporating hydrogel and colored liquids in the side channels (red) and central channel (blue). All scales are microns.
Figure 2.
Schematic of the photolithography (a-c) and soft lithography (d-f) procedure. (a) SU-8 is spin-coated and pre-baked on a bare wafer. (b) Use a transparency photomask (black), UV light is exposed on the SU-8. (c) Exposed SU-8 is then post-exposure baked and developed to define channel patterns. (d) PDMS mixed solution is poured on the wafer and cured. (e) Cured PDMS is then peeled from the wafer. (f) Device is trimmed, punched and autoclaved ready for assembly.
Figure 3.
Procedure of hydrogel incorporating microfluidic assay preparation. (a) Autoclaved PDMS device and coverslip are assembled with plasma treatment (b) to close the microfluidic channels. (c) PDL solution is filled and device is placed in an incubator. (d) After washing and aspiration, coated device is stored in dry oven for 24 hours to render the microfluidic channel surface hydrophobic. (e) Hydrogel is filled into the hydrogel region, and (f) medium is added into the microfluidic channel. Device is ready for cell seeding in incubator.
Figure 4.
(a) Color of type 1 collagen solution (before gellation) with different pH; about 5.0 (left), 7.4 (center) and 11.0 (right). The prepared type 1 collagen solution was then filled into the hydrogel region. (b) SEM image of gelled filbers of type 1 collagen gel (after gellation) with different initial pH; about 5.0 (left), 7.4 (center) and 11.0 (right). Scale bars indicate 1 microns. (c) In hydrogel filling, use 10 ul pipette and slowly inject hydrogel into the gel region. (d) Injected hydrogel perfectly filled into the hydrogel region (red) and stoped by surface tension between post and PDMS wall.
Figure 5.
Cell culture in microfluidic assay. (a) After aspiration of medium in the reservoir, 50–60 μl of the hMVEC suspension is added into one reservoir connected to the center channel. (b) Hydrostatic pressure difference in the reservoirs pushes the suspended hMVECs onto the collagen gel to facilitate attachment. hMVECs right after seeding (right top) and tight monolayer at 1 day (right bottom). Scale bar indicates 250 microns. (c) To add cell suspension or media into the reservoir, use a pipette and gently inject medium into one reservoir after aspiration. The filled liquid automatically fills the channel and replaces the contained medium. (d) Cell seeded assays can be stored in incubator or 6 well plate with various experimental conditions. (e) In 4 days of culture, hMVECs form intact monolayer in channel and on type I collagen wall. Actin filaments and nuclei were then stained with rhodamine-phalloidin (yellow; Sigma-Aldrich) and 4',6-diamidino-2-phenylindole (DAPI; blue; Sigma-Aldrich), respectively. Scale bar indicates 50 microns. EC monolayer was imaged by confocal microscope\textsuperscript{13} and its diffusion coefficient was measured by fluorescent dextran diffusion\textsuperscript{13,14,18}. 

Nat Protoc. Author manuscript; available in PMC 2014 May 27.
Figure 6. Characterization of diffusion profile of applied VEGF. (a) 4 hours of VEGF diffusion into type 1 collagen is simulated (left). Diffusion coefficient of VEGF in type 1 collagen in the simulation was $5.8 \times 10^{-11}$ m$^2$/sec. Arrows indicate diffusive flux. Steady-state concentration profile is shown along the black line (right). Blue color corresponds to 0 ng/ml VEGF while red corresponds to 50 ng/ml VEGF. (b) Gradient (left) and normalized intensity plot from 0 to 30 minutes (center) and from 30 minutes to 10 hours (right). Intensity was measured from right to left sides of type 1 collagen (2.0mg/ml, polymerized at pH 7.4), and then normalized to unity in each figure. (c) Angiogenic response from ECs monolayer was induced by the VEGF diffusion gradient from the right channel (red arrowhead). (d) Perimeter, area and length of angiogenic response were measured (segmented in pale red) and quantified in the captured images.
Figure 7.
Confocal image of 3D angiogenic response into the type 1 collagen (2.0 mg/ml, polymerized at pH 7.4) induced by VEGF diffused from the right channel. (left) 3D projection image of the capillary-like structures formed by hMVEC on day 12 of culture, which was reconstructed from confocal z-stack images. hMVECs formed 3D capillary-like networks extending into the collagen gel, which were stained with CD-146 (hMVEC, red) and DAPI (nucleus, blue). (right) A single z-plane image and x-z and y-z cross-sectional images of the capillary-like networks. Image field corresponds to the left image. Arrowheads indicate a capillary-like structure which is continuous from the monolayer in the channel. An arrow in the x-z image indicates a lumen of the capillary-like structure.
Figure 8.
3D interaction of various types of cells with incorporated hydrogel. (a) Cells are seeded onto hydrogel, and then migrate in individually. (top) Migration of mouse smooth muscle precursor cells (10T 1/2) into type 1 collagen (2.0 mg/ml, initial pH 9.0), on day 6 co-cultured with hMVECs (right channel). (center) Sheet like migration of mouse breast cancer cells (MTLn3) into type 1 collagen (2.0 mg/ml, initial pH 7.4) in day 3 co-cultured with hMVECs (right channel). (bottom) Migration of human breast cancer cells (MDA-MB-231) into type 1 collagen (2.0 mg/ml, initial pH 7.4) on day 3 under 5 % FBS gradient. (b) Cells formed 3D tissue-like structures on hydrogel. (top) Axonal growth into type 1 collagen (1.0 mg/ml, initial pH 7.4) from primary mouse cortical neurons cultured 2 days under 50 μM Forskolin gradient. (center) Primary hepatocyte cultured on type 1 collagen (2.0 mg/ml, initial pH 7.4) in 7 days under interstitial flow condition. Cells were fixed at day 7 and stained for actin filaments (rhodamine-phalloidin; red), nuclei (DAPI; blue) and SEC fraction (GFP; green). Tight tissue-like hepatocyte structure can be found, without migration. (bottom) Human breast cancer cells (MCF-7) cultured on type 1 collagen (2.0 mg/ml, initial pH 9.0) for 3 days under 5 % FBS gradient formed tight tumor colony without migration. (c) Endothelial cells form intact monolayer on hydrogel, showing capillary morphogenesis into hydrogel. hMVEC cells form complex capillary structure in type 1 collagen (2.0 mg/ml, initial pH 7.4) for 5 days under VEGF gradient and ANG-1 supplement. (d) Floating cells in the media migrate into hydrogel under chemoattractant gradient. T lymphocyte cells (Jurkat) migrating into type 1 collagen (2.0 mg/ml, initial pH 7.4) on day 3 under SDF-1 gradient. (e) Co-culture of neutrophil (HL60) in microfluidic channel covered by hMVEC monolayer cultured 3 days. HL60 migrated through the nMVEC monolayer and then into type 1 collagen (2.0 mg/ml, initial pH 10). Cells were fixed at day 4 and stained for actin filaments (rhodamine-phalloidin; red), nuclei (DAPI; blue) and neutrophil (CFDA-SE cell tracker; green). In all other figures not mentioned, actin filaments and nuclei were stained with rhodamine-phalloidin (yellow) and DAPI (blue), respectively. White lines indicate outline of incorporated hydrogel. Scale bars indicate 100 μm.
Figure 9.
Various experimental modes of cell seeding and morphogenesis in hydrogel incorporating microfluidic assay.
<table>
<thead>
<tr>
<th>cell seeding</th>
<th>morphogenesis</th>
<th>Purpose of Experiment</th>
</tr>
</thead>
<tbody>
<tr>
<td>individual cells</td>
<td>hydrogel</td>
<td>migration of various cell types</td>
</tr>
<tr>
<td>epithelial monolayer</td>
<td>channel hydrogel</td>
<td>angiogenesis (surface coating regulates 3D capillary morphogenesis)</td>
</tr>
<tr>
<td>tissue-like 3D structure</td>
<td></td>
<td>metastasis of cancer cells, EMT, hepatocytic tissue, neurons</td>
</tr>
<tr>
<td>embedded cells in hydrogel (spheroid / individual cells)</td>
<td></td>
<td>EMT, metastasis of cancer cells, differentiation of stem cells</td>
</tr>
<tr>
<td>floating cells</td>
<td></td>
<td>transendothelial migration (neutrophils, CTCs)</td>
</tr>
<tr>
<td>co-culture</td>
<td></td>
<td>co-culture with endothelial monolayer (cancer angiogenesis)</td>
</tr>
</tbody>
</table>

**Figure 10.**
Examples of cell-cell 3D interaction in hydrogel. (a) Co-culture of SMCs (10T 1/2; right channel) and ECs (hMVEC; center channel). Active 3D sprouting angiogenesis into type 1 collagen is monitored (into left hydrogels), while the hMVEC monolayer is stabilized by the presence of SMCs (right hydrogels). 10T 1/2 migration was also activated by the existence of hMVEC monolayer in the center channel. Actin filaments and nuclei were stained with rhodamine-phalloidin (yellow) and DAPI (blue), respectively. (b) GFP transfected U87MG (white arrowheads) transmigrated through the hMVEC monolayer. Colors indicate actin (white), nucleus (blue) and GFP (green). Scale bars 50 μm. (c) Interaction between SMC (white arrowhead) and hMVEC (blue arrowheads). A SMC touches the hMVEC monolayer with its filopodia, and becomes adhered to the monolayer. Colors indicate actin (white), nucleus (blue) and EC-specific primary antibody (green). Scale bars 50 μm. (d) GFP transfected breast cancer cell MDA-MB-231 (white arrowhead) interacting with hMVEC (blue arrowheads) sprouts in type 1 collagen scaffold. Colors indicate actin (white), nucleus (blue) and GFP (green). Scale bars 100 μm. (e) Primary neurons generating axons towards GFP transfected glioblastoma cells U87MG (white arrowheads). Colors indicate actin (white), nucleus (blue) and GFP (green). Scale bars 100 μm.
Table 1
Examples of surface coating solution and preferred cell types in hydrogel incorporating microfluidic assay.

<table>
<thead>
<tr>
<th>Surface type</th>
<th>Preferred cell types to culture on cell culture plate</th>
<th>Preferred cell types to culture in hydrogel incorporating microfluidic assay</th>
<th>Purpose of experiment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without coating</td>
<td>-</td>
<td>Neuronal cells, hepatocytes in Fig.8b Stem cells Floating cells (Jurkat in Fig.8d)</td>
<td>To culture cells only on the hydrogel surface To culture cells embedded in hydrogel To make cells only interact with hydrogel surface</td>
</tr>
<tr>
<td>Type 1 collagen</td>
<td>Endothelial cells, hepatocytes, fibroblasts, tumor cells, neuronal cells</td>
<td>Tumor cells</td>
<td></td>
</tr>
<tr>
<td>Poly-Lysine (PDL, PLL)</td>
<td>Neuronal cells, tumor cells</td>
<td>Endothelial cells in Fig.8c, Fig.8e Tumor cells in Fig.8a, Fig.8b</td>
<td>To generate 3D capillary morphogenesis into type 1 collagen To generate 3D migration into type 1 collagen *Enhanced adhesion to type 1 collagen hydrogel</td>
</tr>
<tr>
<td>Fibronectin, laminin</td>
<td>Neuronal cells, tumor cells</td>
<td></td>
<td>*Weak adhesion to type 1 collagen hydrogel</td>
</tr>
<tr>
<td>Matrigel</td>
<td>Endothelial cells</td>
<td>Endothelial cells</td>
<td>To mimic basement membrane</td>
</tr>
</tbody>
</table>

*Note: Fig.8a, Fig.8b, Fig.8c, Fig.8d, Fig.8e represent images or figures in the manuscript.
Table 2

Troubleshooting table.

<table>
<thead>
<tr>
<th>Step</th>
<th>Problem</th>
<th>Possible reason</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>26</td>
<td>Defect in bonding</td>
<td>Mishandling in bonding process</td>
<td>Press the defective region using a pair of tweezers until the unbounded PDMS surface adheres to the glass surface.</td>
</tr>
<tr>
<td></td>
<td>Weak bonding</td>
<td>Weak plasma treatment</td>
<td>Increase plasma treatment time or place the bonded device at 80 °C in an oven for several hours.</td>
</tr>
<tr>
<td>41</td>
<td>Collagen solution flows out into the media channel</td>
<td>Hydrophobicity of the device not yet recovered to the required level</td>
<td>Let device dry for a longer time (e.g., 48 hrs after PDL coating) in an 80 °C oven.</td>
</tr>
<tr>
<td>54</td>
<td>Non-confluent endothelial monolayer on one day</td>
<td>Low hMVECs seeding density</td>
<td>Wait one more day and increase cell seeding density.</td>
</tr>
<tr>
<td></td>
<td>Unhealthy hMVECs</td>
<td>Poor surface coating</td>
<td>Use fresh surface coating solution.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Media evaporation in reservoirs</td>
<td>Maintain a stable humidity in the incubator and add more medium into the reservoirs.</td>
</tr>
</tbody>
</table>