Microfluidic bioreactor for dynamic regulation of early mesodermal commitment in human pluripotent stem cells†‡

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During development and regeneration, tissues emerge from coordinated sequences of stem cell renewal, specialization and assembly that are orchestrated by cascades of regulatory signals. The complex and dynamic in vivo milieu cannot be replicated using standard in vitro techniques. Microscale technologies now offer potential for conducting highly controllable and sophisticated experiments at biologically relevant scales, with real-time insights into cellular responses. We developed a microbioreactor providing time sequences of space-resolved gradients of multiple molecular factors in three-dimensional (3D) cell culture settings, along with a versatile, high-throughput operation and imaging compatibility. A single microbioreactor yields up to 120 data points, corresponding to 15 replicates of a gradient with 8 concentration levels. Embryoid bodies (EBs) obtained from human embryonic and induced pluripotent stem cells (hESC, hiPSC) were exposed to concentration gradients of Wnt3a, Activin A, BMP4 and their inhibitors, to get new insights into the early-stage fate specification and mesodermal lineage commitment. We were able to evaluate the initiation of mesodermal induction by measuring and correlating the gene expression profiles to the concentration gradients of mesoderm-inducing morphogens. We propose that the microbioreactor systems combining spatial and temporal gradients of molecular and physical factors to hESC and hiPSC cultures can form a basis for predictable in vitro models of development and disease.

Bioengineered environments that combine tissue-specific transport and signaling are becoming critical in studies of development, regeneration and disease under settings predictive of human condition.5–8 Technologies reconstructing biologically sound niches along with tight control of the cell environment are starting to offer an entirely new set of tools for stem cell research.5,9–17 In this context, microscale technologies offer potential for conducting highly controllable and highly sophisticated experiments at biologically relevant scales and with real-time insights into cellular responses. Unique advantages of microbioreactors and microfluidic platforms are based on the intrinsically laminar flow in micro-channels and the short transport distances, enabling the maintenance and dynamic changes of well-defined concentration profiles.13,15,18–20

During development, regulatory molecules present themselves in the form of spatial and temporal gradients, rather than at discrete levels to which cell cultures are typically exposed. Concentration gradients guide the formation of the embryo’s axes: Anterior–Posterior (A–P) and Proximal–Distal (P–D), and of the primitive streak (PS), the region in the developing embryo from which mesoderm and definitive endoderm originate.4 Different regions of the PS constitute different signaling environments that are responsible for

Introduction

‘Biomimetics’, ‘cell niche’ and ‘biologically sound environment’ are nowadays among the most used terms in the biological field.3 In our body, cells reside in a complex milieu composed of other cell types, extracellular matrix, and an intricate network of molecular and physical factors that activate signaling pathways and regulate cell fate and function.2 Standard in vitro models lack most of this complexity. Also, relatively large operating volumes and periodic exchange of medium do not allow for the generation of precise spatial and temporal patterns of stimulation. Collectively, these limitations result in unrealistic and uncontrollable biological readouts that fall short of predicting the actual in vivo situation, of relevance both to fundamental research and cell and drug screening for medical applications.1–4

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induction of specific lineages, with morphogens such as ActivinA, BMP4, and Wnt3a playing major roles in these events.4

hESCs are now widely accepted as an ‘ideal’ model for studying the complex developmental processes.21–23 The emergence of iPSCs has added an additional degree of significance: patient-specific cells can be obtained for a multiplicity of studies ranging from drug screening to personalized medicine.24–28

We hypothesized that the application of spatial and temporal gradients of multiple factors to hESCs and hiPSC cultures would provide predictable and realistic in vitro models of development. To test this hypothesis, we designed a microioreactor platform for stem cell culture with spatial and temporal concentration gradients of regulatory molecules, guiding cell development, specification, and commitment to the mesodermal fate. The platform combines some of the advantages of multi-well plates (small volume, high-throughput, independent wells) and perfusion bioreactors (steady state, enhanced mass transport, application of signals) while respecting the constraints dictated by the biological system of choice (e.g., absence of shear forces). Mathematical modeling of flow and mass transport within the bioreactor was used during the design phase to determine the geometry of the cell culture modules and microfluidic channels. The model predictions were experimentally validated using labeled molecular markers. The technology we developed was then applied to test its efficacy in yielding valuable results on a relevant biological issue: the early mesodermal commitment in human pluripotent stem cells.

The main advantage of our system is that it always works at a steady state, so that all established concentrations are invariant with time, a situation resembling homeostasis in vivo. In addition, if the operator introduces a defined perturbation of concentration in the system, a new steady state is quickly established. In contrast, static well plates operate under conditions that vary from one medium change to another. Thanks to this fundamental property, we were able to evaluate mesodermal induction by measuring and correlating the expression of key genes after exposure to tightly controlled concentration gradients of mesodermal-inducing morphogens.

Methods

Microfluidic bioreactor design, fabrication and assembly

The microioreactor platform was designed to meet a set of specific requirements: (i) generation of multiple concentration gradients, (ii) support of long-term culture of EBs formed from hESCs and hiPSCs in a no-shear environment, (iii) high-throughput studies with large numbers of replicates, (iv) compatibility with on-line imaging and standard analytics, (v) ease of retrieval of the cell samples for additional post-processing, and (vi) capability to apply fast dynamic changes of environmental signals (i.e. morphogens and inhibitors). Our platform comprises a matrix of conical microwells each accommodating one single EB. The EBs are protected from hydrodynamic shear forces and exposed to stable concentration gradients generated by an integrated microfluidic platform with independent rows of microwells connected to two lateral flow channels. Microwells within a row communicate between each other, while separate rows represent replicates of the same conditions.

Two configurations were fabricated: (i) microioreactors with 5 microwells per row for preliminary experiments and gradient-validation studies, and (ii) microbioreactors with 8 and 6 microwells per row for all human pluripotent stem cell studies. The overall dimensions of all microbioreactors was kept constant for all prototypes. The final designs are shown in Fig. 1, with the microwells (diameter: 560 μm at the top, 100 μm at the bottom; depth: 850 μm) and the channels (100 μm deep, 150 μm wide) connecting the rows of microwells to the lateral flow channels. A single microioreactor yields 120 data points, with 8 concentration levels across the gradient with 15 replicates.

![Fig. 1 Microbioreactor design. A The microioreactor comprises a matrix of microwells arranged in independent rows and connected to two lateral flow channels. The overall dimensions (45 × 21 mm length × width) are comparable to those of a standard microscope slide. The length and size of the flow channels were kept constant for all prototypes while the number of microwells could be varied. The CAD drawing in panel A contains all relevant dimensions: the top diameter of the 850 μm deep microwells is 560 μm, tapering towards the bottom with a ~18° angle. 100 μm deep × 150 μm wide channels connect rows of microwells to the main lateral flow channels. The corresponding mold was fabricated using standard stereolithography rapid prototyping techniques and used to obtain the final PDMS microioreactor via replica molding. B A color tracer fills the microioreactor, allowing visualization of the channels and microwells; please refer to Fig. 2 for color-coded concentration maps showing the gradient formation. C An exploded view of the individual components of the clamping device: the aluminum base at the bottom, the microioreactor, a polycarbonate manifold and the clamping plate at the top. Two slide latches were used to tighten the manifold against the microioreactor ensuring hydraulic sealing. D The assembled device.

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The platform was designed with the aid of 3D CAD software. The mold was produced via stereolithography rapid prototyping, and the microfluidic device was replica-molded in poly(dimethylsiloxane) (PDMS). Connections were made using 1.5 mm diameter Tygon® tubes (Cole Parmer) and blunt 21 G hypodermic needles (McMaster-Carr).

A reversible hydraulic sealing between the components is ensured by a clamping device, which couples ease of assembly and retrieval of the cell samples. Briefly, the microwell-cell chamber is placed onto a glass slide or PDMS membrane within the aluminum base. A polycarbonate manifold is placed above the cell chamber and the seal is established via a clamping plate, with two slide latches fastened to the clamping plate, with two slide latches fastened to the clamping plate to provide a seal of the manifold against the cell chamber by mating with studs fixed to the base. Thumbscrews are used to position the height of the spring-loaded studs, allowing fine-tuning of the clamping force. The cell culture area in the assembled platform is optically transparent, for microscopy observation. The microbioreactor, and the tubing, connections, and the clamping system were steam sterilized (20 min at 121 °C) prior to use.

Computational models of the flow and transport

The flow regimes and concentration profiles were mathematically described by using the 2D incompressible Navier–Stokes equations and the convection-diffusion equations, that were solved numerically using published parameters.\(^{12}\) These two non-linear partial differential equations were solved simultaneously using a finite element analysis solver (COMSOL Multiphysics). Relevant parameters characterizing the studied morphogens are listed in Table 1. Simulations were performed in the range of flow and transport conditions encountered during the EB culture, to verify that the gradients’ shape was independent of the applied flow rate and only dependent on the molecular weight (translating into different diffusion coefficients) of the dissolved molecules.

Quantitative fluorescent imaging

To validate the theoretical models, fluorescently labeled dextran was used to study mass transport in the bioreactor. Fluorescein and three different dextran molecules were used to cover the range of molecular weights corresponding to the growth factors of interest: 2’,7’-Dichlorofluorescein diacetate (Sigma) (0.4 kDa), Cascade Blue-conjugate (10 kDa), Fluorescein-conjugate (40 kDa), and Tetramethylrhodamine-conjugate (70 kDa), all from Molecular Probes. Each dextran was dissolved in tissue culture medium at 50 μg mL\(^{-1}\), 0.2% fluorescein was dissolved in PBS. Each solution was perfused through one of the main channels, while the solvent alone was used in the other main channel, using 5 mL Becton-Plastik syringes filled with the two solutions and connected to the syringe pumps (Harvard Apparatus) by 1.5 mm diameter Tygon® tubing. The system was assembled under the microscope (Leica CTR6000) and fluorescent images were taken at different time points at 10× resolution. The fluorescence intensities were quantified using commercial software (ImageJ) and compared with the model predictions.

hESC and hiPSC culture

Two lines of hESC and one of hiPS cells were used as follows.

**H1-BAR-Venus (BAR: β-catenin activated reporter) hESCs.** H1 hESC line was modified incorporating a new Wnt/β-catenin reporter system, β-catenin-activated reporter (BAR), and its accompanying control reporter system, found unresponsive BAR (fuBAR).\(^{29}\) hESC colonies were routinely passaged on irradiated MEFs feeder layers (Global Stem) after reaching 80% confluence using 1.75 Units/mL Dispase (Invitrogen, Catalog number 17105-041). Cells were expanded in human ESC Knockout Serum Replacement (KSR) medium: DMEM/F12 (1 : 1) (1 ×, Invitrogen 11320), Knockout Serum Replacement (Invitrogen 10828) 20%, NEAA Solution, 10 mM (Invitrogen 11140) 0.1 mM, l-Glutamine, 200 mM (Invitrogen 25030) 2.0 mM, Sodium Pyruvate (Invitrogen 11360) 1.0 mM, Strep

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solution (Invitrogen 15140) 1%, β-ME, 55 mM (Invitrogen 21985) 0.1 mM, FGF-2 (Invitrogen) 8 ng ml\(^{-1}\). All solutions were sterile filtered and protected from light. Cells were cultured in differentiation medium: DMEM, high glucose (Invitrogen 11995), FBS (Moon lab) 20%, NEAA Solution, 10 mM (Invitrogen 11140) 0.1 mM, L-Glutamine, 200 mM (Invitrogen 25030) 2.0 mM, Sodium Pyruvate (Invitrogen 11360) 1.0 mM, Pen/Strep Solution (Invitrogen 15140) 1%, β-ME, 55 mM (Invitrogen 21985) 0.1 mM.

**hES2 and BMC1.** The second line of hESCs and the hiPSCs were used exactly as described above, except for the compositions of culture media. The growth medium was DMEM/F-12 with KooSr, 20%, MEFs conditioned medium 10%, NEAA Solution 0.1 mM, L-Glutamine 2 mM, Pen/Strep 1 : 100, FGF-2 (Invitrogen) 20 ng ml\(^{-1}\). The differentiation medium was STEMPRO 34 plus supplement component according to manufacture (Invitrogen) with Ascorbic Acid (Sigma) 50 μg ml\(^{-1}\), Transferrin (Roche) 150 μg ml\(^{-1}\), MTG (in medium) (Sigma) 3 μL ml\(^{-1}\) of diluted 3 : 100 BMP4 (Peprotech) 0.5 ng ml\(^{-1}\) plus Rock inhibitor (Tocris Bioscience) 0.5 μM in aggregation low adhesion plates (Fisher), and BMP4 (Peprotech) 10 ng ml\(^{-1}\), FGF-2 (Invitrogen) 5 ng ml\(^{-1}\) and ActivinA (R&D Systems) 6 ng ml\(^{-1}\) at day 1–4.

**Formation of EBs.** EBs were formed in Aggrewell400\(^{\text{R}}\) plates starting from single cell suspensions. Cells were gently detached using TrypLe and counted. Seeding densities were determined as to have 1000–1500 cells/EB. After spinning, detached using TrypLe and counted. Seeding densities were determined as to have 1000–1500 cells/EB. After spinning, EBs were recovered from Aggrewell plates, washed and allowed to gravity settle in a 50 mL Falcon tube to remove debris and single cells, and transferred to Eppendorf microcentrifuge tubes. Culture medium was care-

**Dissociation and immunostaining of EBs.** When required for post-gradient exposure analyses, EBs were dissociated to single cells. Briefly, EBs were collected and transferred to Eppendorf microcentrifuge tubes. Culture medium was carefully removed and a washing step with PBS (Ca\(^{2+}\), Mg\(^{2+}\) free) was performed. EBs were then treated with 200 μL of a 2 mg ml\(^{-1}\) Collagenase I ( Worthington Biochemical) in PBS solution with 20% FCS for 1 h at 37 °C, carefully resuspended and centrifuged at 800 g for 2 min. After removing the supernatant, 50 μL of 0.25% Trypsin solution, diluted 1 : 4 in PBS (Ca\(^{2+}\), Mg\(^{2+}\) free) were added for 5 min while gentle resuspending ensured proper dissociation. Cell suspensions are centrifuged at 1800 g for 3 min to remove Trypsin. Cells were then resuspended in the desired antibody solution and plated for subsequent image analysis. Nuclei were counterstained with a 1.5 μg ml\(^{-1}\) Hoechst 33342 (Invitrogen) nuclear dye solution for 10 min at 37 °C.

**Experimental set-up**

**Exposure of EBs to multiple gradients of morphogens**

Low volumes of high-density suspensions of EBs were carefully dispersed on top of the microwell areas of the microbioreactors, with an excess of approximately 50% EBs, and incubated. After a few minutes the devices were observed under the microscope to confirm proper filling of the microwells. If needed, additional EBs suspension was added to each microbioreactor until ≥90% of the wells were occupied by one single EB each. Typically, after one seeding cycle, the occupancy of the was 75–80%. The clamping plate was then assembled and the platform was transferred to the incubator. Syringes filled with culture media of a desired composition were connected to the inlets of each platform using Tygon tubing. The flow rate was set at the syringe pump and medium perfusion was initiated.

Wnt3a, ActivinA, BMP4 and their inhibitors Dkk1 and SB-431542 were chosen based on their known effect on inducing mesodermal differentiation and on their levels and temporal sequences defined in the protocol developed in the Keller lab.\(^{20}\) The temporal windows of exposure to gradients spanned from day 2 to day 4 following the aggregation of EBs. Combined gradients of Wnt3a, ActivinA and BMP4, with or without opposing gradients of the inhibitors Dkk1 (for Wnt3a) and SB-431542 (for ActivinA) were studied as specified in Table 2 and are indicated as ng ml\(^{-1}\), where not specified otherwise.

**qRT-PCR**

Total RNA from EBs harvested from microbioreactors was isolated using the TRIzol protocol (Invitrogen) or the RNAqueous Micro kit (Ambion, Life Technologies), treated with DNase I (Invitrogen) and reverse transcribed with the High Capacity Reverse Transcription kit (Applied Biosystems). Real-time quantitative PCR was carried out with the StepOnePlus qPCR machine (Applied Biosystems). Each cDNA sample was amplified in duplicate or triplicate by using the Fast SYBR Green Master Mix (Applied Biosystems). Relative expression was obtained from Delta Ct values normalized to those obtained from the amplification of GAPDH transcripts.

The primers used for these experiments were as follows:

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<td>0</td>
</tr>
<tr>
<td>SB-431542</td>
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<td>Dkk1</td>
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<td>0</td>
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<td>0</td>
<td>150</td>
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Table 2 Conditions tested. (a–d) different medium-compositions corresponding to the listed morphogens’ quantities. Left and Right refer to the two streams of fluid entering the microbioreactor. All values, if not specified otherwise, are in ng ml\(^{-1}\).
Quantitative fluorescent imaging of Venus expression in cultured cells

As an online semi-quantitative assay, we evaluated H1-BARV cell response by imaging the activation of the Venus-tagged reporter gene. Fluorescent images of the nuclei were used to determine the total cell number. The fraction of cells expressing Venus signal was determined by the pair-to-pair comparisons of fluorescent images of the two channels: Venus in green, and Hoechst in blue, using a script we developed in Matlab. Exposure times, gain and intensity values were the same in all set of images to prevent thresholding mismatches and validate the comparisons between the different conditions. An average of 3 images per EB and 10 EBs per condition were analyzed.

Results

Validation of the microbioreactor

Computational modeling and experimental validations confirmed that the platform assures the stable establishment of complex patterns of stimulation easily modifiable in space (by changing geometric parameters of the microbioreactor) and time (adjusting perfusion settings). Moreover, the gradients were the same for a range of molecular weights of interest (selected to match the size of growth factors). All large molecules (20–100 kDa) are transported at similar rates, and only the SB-431542, a small 0.4 kDa molecule, had a higher diffusion coefficient (Table 1). Fig. 2 illustrates the experimental validation of computational modeling of mass transport within the microbioreactor.

The theoretical concentration heat-map (zoomed vision of the microwell-cell chamber area) and the profile developed across the rows of microwells are compared with the experimental data in panel B (left). The modeled and experimental values of concentrations were not significantly different.

Table 3 reports approximated average concentrations values, measured as percent of maximum, in the individual microwells for the different microbioreactor configurations used (platforms with 5, 6, and 8 microwells per row). The values were obtained from the modeled concentration profiles.

Yield of EB seeding

EBs were easily seeded into the microbioreactors by dispensing suspensions of excess numbers of EBs (based on microscope observation of the docked vs. floating EBs). After <5 min of gravity settling, EBs docked into the microwells with an average efficiency of 90%, determined as the fraction of total microwells containing EBs (one per microwell) after two cycles of seeding. The conical geometry of the microwells maintains EBs in place and protects them from shear forces during culture (Fig. 3).

EBs made from hESC and hiPSC were cultured within these platforms for up to four days with exposure to the micro-fluidic-driven concentration gradients, after which time they were retrieved for subsequent analyses. A preliminary validation of the device was performed flowing Calcein AM enriched medium (5 μM) through one of the flow channels, thus generating a linear gradient of the membrane dye which resulted in a proportionate fluorescent stain of the EBs inside the microwells (Fig. S1, ESI†).

Responses of H1-BARV-EBs to Wnt3a/ActivinA/BMP4 concentration gradients

After exposure to a concentration gradient (condition ‘a’ in Table 2), H1-BARV cells expressed an on-line detectable Venus signal. Fluorescence could be observed during culture with EBs docked inside the microwells, as well as on harvested EBs (second row of Fig. 4). Qualitative differences were observed right away. Given the three dimensionality of the EBs, additional processing was needed to assess the levels of gene expression. The level of activation of the Wnt3a/β-catenin signaling pathway was evaluated using the image analysis of the Venus signal (at a single cell level on dissociated EBs) and q-PCR for Axin2, a canonical Wnt/β-catenin target gene.31

The obtained results (Fig. 5) showed that the pathway activation was not directly proportional to the concentration of Wnt3a in culture medium (which correlated to the position of each EBs in the row of microwells). Instead, distinct concentration levels were identified that resulted in increased gene expression as assessed by both the fraction of Venus positive cells and by qPCR data. This result is consistent with the notion that the combined action of the three morphogens stimulates complex signaling cascades, resulting in non-proportionate cell responses.

The top histogram plots the obtained results for EBs contained in microwells from 1 to 8 across the gradient. The bottom histogram reports results of q-PCR for Axin2 in corresponding EBs from a parallel experiment; its activation profile confirmed the Venus quantification.

Responses of hES2-EBs and BMC1-EBs to mesodermal-inducing gradients of morphogens

The set of genes that we chose to monitor are markers and regulators of early mesendodermal (Brachyury T) and mesodermal (Mesp1, Isl1, KDR) commitment.32 Mesp1 and Isl1 have been described as ‘master regulators’ of cardiac progenitor specification, where Mesp1 drives cardiac differentiation by inhibiting canonical Wnt signaling (a DKK1-mediated process).33 In turn, Wnt3a/β-catenin is a pathway that was proved to have a biphasic effect on cardiogenesis.34 KDR is a known marker for cardiac mesoderm populations35 and Brachyury (T) is a mesendodermal marker induced by Wnt.

EBs were subjected to a stable gradient of these morphogens (compositions b–d in Table 2) for 24 h starting at either day 2
or day 3 since initiation of differentiation (day 0: the time of EBs formation). These temporal windows for the gradient experiments were determined based on the known sequences of events leading to cardiac mesoderm specification.\textsuperscript{30,36,37}

Parallel static experiments were performed for all conditions tested, exposing EBs populations to different levels of concentrations that were within the range generated by the microbioreactors. EBs were seeded onto the microbioreactors as for the gradients experiments and allowed to settle inside the microwells. The microbioreactors were then operated statically, each exposing the docked EBs to discrete and uniform microenvironment compositions chosen within the range covered by the gradients. The results obtained with hES2-derived EBs (Fig. S2, ESI\textsuperscript{4}) confirm the pattern of expression that will be detected in the case of the graded exposure. The modulation of the local microenvironment composition led to the formation of more complex gene expressions patterns. Also, control experiments exposing EBs to "flat gradients" obtained for identical cocktails of cytokines in both flow channels resulted in flat gene expression profiles (Fig. S3, ESI\textsuperscript{4}).

Fig. 6 shows the expression of representative mesodermal and mesendodermal genes in hESCs (hES2 line) and hiPSCs (BMC1 line) following exposure to a combined Wnt3a/ActivinA/BMP4 gradient. Once again, gene expression profiles changed between the microwells and in a cell-line dependent manner. Major differences could be detected in the expression of early genes (T and Mesp1) while more subtle differences were observed in expression levels of later genes. The relative expressions for T and Mesp1 were bimodal, indicating differential lineage specification mechanisms within the EBs exposed to the particular local microenvironment along the concentration gradient. Notably, significant differences in the relative expressions of these genes can be detected in the case of EBs cultured in the perfused microbioreactors (which operate at steady-state conditions), as shown in Fig. 6. On the contrary, even if resulting in similar expression patterns, the levels measured in statically operated microbioreactors fail to reach statistical significance, as shown in Fig. S2, ESI\textsuperscript{4}.

From the expression levels measured for the BMC1 hiPSC line, we noticed the presence of two local maxima, corresponding to microwells 3 (local maximum) and 6 (absolute maximum). The microenvironment of microwell 6 (20–30% of the initial factor concentrations) led to the highest expression levels of T, Mesp1 and KDR, while the second local maximum was expressed at the higher concentrations of microwell 3 (60–70% of the initial factor concentrations). In this case however, we recorded a lower KDR expression, suggesting a delayed progression along the mesodermal lineage (KDR is expressed later than T and Mesp1 in the cascade of events leading to mesoderm induction). For the hES2 hESC line, a similar behavior could again be observed in microwell 6, but an evident local maximum corresponding to microwell 3 failed to show. This evidence suggests that similar expression levels could be triggered by different medium compositions, further hinting at the potential role of endogenous mechanisms activated by increased/decreased levels of stimuli.

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Fig. 3 Cultivation of embryoid bodies (EBs) formed from human stem cells in the microbioreactor. A Uniformly sized EBs are formed from human ES cells (hES2 and H1-BARV lines) and hiPS cells (BMC1 line) using Aggrewell\textsuperscript{1} plates; magnifications 4 x and 10 x, respectively. B An entire microbioreactor seeded with EBs (images taken at the stereoscope). Magnified inserts: independent rows of microwells with individual EBs.
In another set of experiments, we exposed hES2-EBs and BMC1-EBs to opposing gradients of ActivinA/BMP4 and either SB-431542 (ActivinA inhibitor) or Dkk1 (canonical Wnt signaling inhibitor). The histograms in Fig. 7 summarize the most relevant findings and compare the gene expression profiles in control experiments, performed without opposing inhibitors-gradients, with those in double-gradients runs. The effects of exogenously inhibited Activin signaling via SB counter-gradient treatment are shown in the left column; Wnt3a inhibition via Dkk1 counter-gradient is in the right column. Again, significant differences were observed in the early vs. late genes expression profiles. The cells were differentially activated by the local microenvironment, in response to concentration gradients.

Interestingly, we observed that Activin inhibition resulted in increased expression levels of all earlier genes, suggesting that: (i) these cells might endogenously produce ActivinA, thus further increasing its concentration in the local microenvironment and bringing it to sub-optimal levels, and that (ii) partial inhibition of this signaling pathway might be required to optimally induce early mesodermal/mesendodermal specification. Wnt inhibition via Dkk1 appeared to elicit an almost linear correlation with the expression levels, with a direct proportionality for T and KDR, and indirect proportionality for Mesp1.

Conclusions and discussion

We designed a microbioreactor platform capable of generating stable concentration gradients over a large number of EBs (120 in a system comparable by size to a microscope slide) in a shear protective environment. These microbioreactors provide multiple gradients of molecular factors in 3D cell culture settings, with complex sequences of time- and space-resolved gradients, and the application of fast dynamic changes of environmental signals. In developmental studies of human embryonic (hESC) and induced pluripotent stem cells (hiPSC), we were able to screen for a large number of variables (concentration levels, cocktails of factors and their inhibitors, space and time gradients). Due to the fast establishment of steady-state conditions, these microbioreactors ensured maintenance of precise and time-invariant compositions of the soluble microenvironment to which individual 3D stem cell aggregates are exposed.

We show the modulation of pathway activations by the local microenvironment to result in non-linear cell responses to linear concentration gradients. This is a physiologically relevant behavior, since the mesodermal/mesendodermal differentiation programs are tightly regulated by sequences...
of factors evolving in space and time. The EBs can respond by ‘proper’ activation of the mesodermal key genes only if and when exposed to the correct levels of cytokines. The use of inhibitors confirmed the non-proportional response in the cultured EBs. Correlations and mismatches between expression levels of activated genes following precise spatial-temporal stimulation suggest induction of specific differentiation programs. Snapshots of relative gene expression levels at defined times post-exposure to the concentration gradients were compatible with previous data.

Several publications reporting devices for screening of molecular factors in cell culture can be found in the literature, among which we cite the work of Kim et al. However, the biological questions we investigated required the development of the novel microfluidic device we describe here that is different from previously reported systems in several respects. First, we performed 3D culture while most other systems allow only for 2D culture. Second, our seeding method by docking of EBs does not expose the cells to shear stress, while other systems employ dynamic loading of the cells. In addition, the biological samples are easily retrieved from our device at any time during culture, something that is not typically possible with other systems. Finally, the high throughput characteristics of our system reside with the large number of replicates: ≥10 EBs per condition were tested in each microbioreactor, whereas most of the existing devices generate matrices of different conditions with one single sample per condition. For a study like ours, the intrinsic variability of the cell phenotypes requires significant numbers of replicates for assessing statistical differences between the groups.

Some of the interesting future directions include: (i) estimating the effects of compensatory mechanisms and (ii) endogenously produced signaling molecules on upstream and downstream effects in neighboring EBs occupying microwells of the same row. The potential compensatory mechanisms activated by a biological system (a differentiating EB in our case) when exposed to defined levels of induction media, could be partially responsible for the lack of direct proportionality between gene expression and morphogen concentrations. This effect could be EB-autonomous (i.e., with a single EB activating its own compensation) or due to a cross-talk between adjacent EB microenvironments, with secretion of diffusing factors that...
activate feedback loops mechanisms. Future studies will focus on analyzing the gene expression profile of EBs when cultured at defined positions across the gradients without any other EB occupying neighboring microwells (i.e. one EB per row of microwells, multiple reactors in parallel as biological replicates). We are also currently developing a system allowing direct formation of EBs within the bioreactors microwells. Furthermore, we are adapting our microbioreactors to be compatible with the Fluidigm Single-Cell Gene Expression platforms (Fluidigm Corp, South San Francisco, CA 94080), which we have used in studies of directed differentiation of human stem cells. This will collectively enable us to implement longer and multi-step differentiation protocols by culturing EBs in the microbioreactor from aggregation to later stages of mesodermal differentiation while performing different time point analyses along the differentiation process.

In summary, our approach helps recapitulate some aspects of the patterns of stimulation that direct cells differentiation during cardiac development. The main aim of this paper is proving the capability of our device to sustain cell differentiation in a physiologic microenvironment, to apply controlled stimulation paradigms guiding cell differentiation and to finally allow obtaining relevant readouts from the biological samples. To this end, our platform can generate dynamically stimulation paradigms guiding cell differentiation and to finally allow obtaining relevant readouts from the biological samples. To this end, our platform can generate dynamically changing spatial and temporal gradients of multiple factors (promoters and inhibitors) and enable studies of the effects of these factors on lineage specification. We developed a simple, highly controllable device that allows studying complex interactions of signaling factors. We are just starting to see the full potential of such system, and the readouts and outcomes we can obtain. Some of the most interesting future applications will include the optimization of conditions for generating human progenitor cells and the establishment of physiological models for screening of therapeutic cells and drugs.

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