1 Defined microenvironments trigger *in vitro* gastrulation in human pluripotent stem cells

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17 Abstract

18 Embryogenesis is orchestrated through local morphogen gradients and endometrial constraints that give rise to the three germ layers in a well-defined assembly. In vitro models of embryogenesis 19 have been demonstrated by treating pluripotent stem cells in adherent or suspension culture with 20 21 soluble morphogens and small molecules, which leads to tri-lineage differentiation. However, treatment with exogenous agents override the subtle spatiotemporal changes observed in vivo 22 23 that ultimately underly the human body plan. Here we demonstrate how microconfinement of pluripotent stem cells on hydrogel substrates catalyses gastrulation-like events without the need 24 for supplements. Within six hours of initial seeding, cells at the boundary show elevated 25 26 cytoskeletal tension and yes-associated protein (YAP) activity, which leads to changes in cell and 27 nuclear morphology, epithelial to mesenchymal transition, and emergence of defined patterns of primitive streak containing SRY-Box Transcription Factor 17 (SOX17)⁺ T/BRACHYURY⁺ cells. 28 Immunofluorescence staining, transcript analysis, and the use of pharmacological modulators 29 reveal a role for mechanotransduction-coupled non-canonical wingless-type (WNT) signalling in 30 promoting epithelial to mesenchymal transition and multilayered organization within the colonies. 31 These microscale gastruloids were removed from the substrate and encapsulated in 3D hydrogels, 32 where biomaterials properties correspond to maintenance and spatial positioning of the primitive 33 streak. Together, this approach demonstrates how materials alone can nurture embryonic 34 35 gastrulation, thereby providing an *in vitro* model of early development.

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1 Introduction

2 One of the most important events in developmental biology is gastrulation, where single layered 3 pluripotent epiblast cells go through a series of carefully regulated cell fate decisions to form the 4 progenitors of the three germ layers ¹. Previous studies in mouse have revealed that the posterior side of the embryo initiates a region of cells undergoing epithelial-to-mesenchymal transition (EMT) 5 6 to form the primitive streak, and delaminate from the epiblast surface after gaining mesenchymal 7 motility². The cells in this primitive streak region are positive for the mesodermal marker T/BRACHYURY, and as they ingress inwards, they segregate towards endoderm progenitors with 8 9 endodermal marker SOX17 as the streak extends. These mes-endodermal cells further proceed to 10 ingress through the primitive streak into the gastrulating embryo³. The dramatic cellular identity 11 changes *in vivo* are attributed to an interplay of the TGF β , WNT and FGF signalling pathways with their respective antagonists⁴ and their order of activation and patterns of gene regulation have been 12 well studied⁵. 13

Recently, there has been an effort to understand developmental signals in the context of 14 15 biophysical forces during gastrulation; specifically, the local mechanical forces and geometric constraints created by surrounding extra-embryonic tissues during embryo implantation into the 16 17 uterine lining⁶. Biophysical regulation of the gastrulation process is difficult to study due to ethical and physiological limitations with handling human embryos after the appearance of the primitive 18 19 streak, approximately 14 days after fertilisation⁷. The anatomy of embryonic development contains 20 several phases where extra-embryonic pressure and the mechanics of the surrounding tissue will 21 impose constraints on morphogenesis. However, current opinion is mixed regarding the importance of the biophysical microenvironment in directing early stages of embryogenesis. Therefore, in vitro 22 23 approaches to culture or encapsulate pluripotent stem cells or epiblast stem cells, using hydrogels, microcarriers, scaffolds and other biomaterials have been used to model gastrulation in vitro towards 24 understanding the role biophysical factors play during embryogenesis^{8,9}. 25

In vitro models for gastrulation using microengineering have allowed the effect of geometric
 confinement during tri-lineage differentiation to be probed^{10,11}. After bone morphogenic protein-4
 (BMP4) stimulation for 48 hours, pluripotent stem cells (PSCs) undergo spatial patterning in response
 to confinement, which is reminiscent of *in vivo* processes. Most *in vitro* studies use rigid glass surfaces
 (~3-4 GPa), and exogenous supplements of soluble morphogens to trigger differentiation. In contrast,
 the signalling gradients *in vivo* are dynamic, involving local gradients of paracrine and autocrine

signals to drive morphogenesis in a confined microenvironment with variable viscoelastic 1 2 properties¹². Pre-streak formation events involve actin-controlled oriented cell movements and 3 deformation on the posterior epiblast, which guide dynamic local gradients of signalling to coordinate primitive streak progression.^{13,14} Recently, Weaver and colleagues reported the appearance of 4 multicellular T/BRACHYURY⁺ "gastrulation-like" mesodermal nodes at the colony edges on soft 5 patterned substrates following BMP4 driven differentiation, highlighting the interplay between 6 mechanics and morphogens¹⁵. These *in vitro* gastruloids show similarities in gene expression and 7 cellular organization with several hallmark features of gastrulation – mes-endodermal Identity (along 8 9 with comparable downregulation of pluripotent identity), evidence of EMT, and collective cell 10 movement after loss of pluripotency¹⁵.

11 In this article we demonstrate how human pluripotent stem cells spontaneously differentiate 12 into SOX17⁺ T/BRACHYURY⁺ gastruloid-like structures within 2 days when microconfined on precise matrix-conjugated deformable substrates, without the need for exogenous supplements. The degree 13 14 of lineage specification corresponds to matrix biophysical conditions, with a decline in pluripotency coinciding with EMT and mes-endodermal gene activation. Treatment with BMP4 enhances this 15 transformation, resulting in the appearance of textured wave-like regions of multilayered mes-16 endodermal SOX17⁺ and T/BRACHYURY⁺ structures within the micropatterned colonies. Gene 17 18 expression analysis coupled with pathway inhibition studies indicates that mechanotransduction 19 triggers gastrulation in confined colonies through non-canonical wingless-type (WNT) signalling. Release from confinement and encapsulation in tailored hydrogels results in spatial patterning of 20 differentiation, thereby providing a "materials-centric" method of forming gastrulation mimics to 21 22 model embryogenesis.

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24 **Results**

25 Protein-conjugated hydrogels facilitate human induced pluripotent stem cell (iPSCs) culture.

To assess the behaviour of hiPSCs on compliant substrates, we chose to work with polyacrylamide (PA) hydrogels with tuneable stiffness that can be further modified using soft lithography to define regions of adhesivity¹⁶. Briefly, the concentration of acrylamide and bis-acrylamide was varied to create PA gel solutions for stiffness of 1, 10 and 100 kPa (10-fold increase spanning physiological tissue) and then polymerised on chemically modified glass coverslips. The surface of the PA was treated with hydrazine hydrate and then imprinted with an oxidized protein using soft lithography to

form the covalent Schiff base using patterned or flat polydimethylsiloxane stamps to mediate cell 1 adhesion on an otherwise inert PA surface. The same stamps were used to assist with physical 2 3 adsorption of the protein on a clean glass surface to define protein islands. For this study, we had 4 four test groups: protein coated glass, protein patterned glass, protein coated hydrogels and protein patterned hydrogels (hydrogels formulated at 1, 10 and 100 kPa). These groups assessed the 5 response of iPSCs towards confinement alone, stiffness alone and geometry combined with stiffness, 6 7 all compared to bare glass controls. To screen optimal proteins for iPSC attachment and proliferation, we assayed common matrix proteins involved in *in vitro* stem cell culture: EHS Laminin, Laminin-521, 8 9 Collagen, Fibronectin, hESC qualified Matrigel and rh-Vitronectin, deposited on substates via soft 10 lithography (Figure S1A). These trials revealed rh-Vitronectin to be the most compatible protein 11 which readily facilitated iPSC attachment and proliferation (Figure 1A), as compared to all other 12 tested proteins where high cell death or little to no cell attachment was observed. Since the 13 microcontact patterning process of the PA hydrogels requires an oxidation of the printing protein 14 using sodium periodate at room temperature, rh-Vitronectin, which is generally stable at room 15 temperature proved to be a better alternative to matrigel, which was unstable in supporting patterned iPSC adhesion under these conditions. 16

Having identified vitronectin as a suitable protein for hiPSC culture, we next asked whether these hydrogel matrices would maintain the pluripotent phenotype. The hiPSCs on glass showed positivity for pluripotency markers (OCT4 and NANOG) and were simultaneously found to be negative for germ layer lineage markers (**Figure S1B**). hiPSCs demonstrated healthy attachment on the nonpatterned and patterned PA substrates, with adherent colonies initiating multilayered growth after 48 hours.

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Substrate properties alone guide human pluripotent stem cell lineage specification towards mes endodermal identity.

Having established optimal conditions for iPSC culture on hydrogels, we next sought to assess the expression of molecular markers of pluripotency and tri-lineage differentiation. For geometric confinement, 250µM and 500µM diameter circles were tested. The hiPSCs were dissociated to single cells and seeded at a uniform cell density while being supplemented with Y-27632 and allowed to grow for 48h before fixation. Optimal cell density was selected based on conditions that foster near confluence on day one.

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To determine the pluripotency status and lineage identity of hiPSCs on these different 1 2 surfaces, we immunostained the cell populations with the pluripotency marker OCT4 (POU5f1), as well as definitive endoderm marker - SOX1717 and mesoderm/primitive streak marker -3 4 T/BRACHYURY⁵. We observed that the cells cultured on glass controls and glass patterns, maintained pluripotency with uniform expression of OCT4 (Figure 1B). However, hiPSCs cultured on non-5 patterned hydrogels across each stiffness condition demonstrated decreased expression of OCT4. 6 Microconfined cells across all three stiffnesses demonstrated OCT4 expression restricted to the 7 colony edges, with complete loss of signal in the centre (Figure 1B,D). This appearance of an OCT4 8 annulus in a confined colony was observed previously during cardiac differentiation on patterned 9 glass using CHIR99021 induction,¹⁸ which was postulated to be a Wnt signaling-mechanics 10 11 relationship. Expression of the endoderm marker SOX17 was negligible in colonies on glass, with 12 evidence for modest expression in colonies on non-patterned hydrogels, indicating a potential role for substrate mechanics in regulating endoderm specification. In contrast, there was a striking 13 14 upregulation of SOX17 in hydrogel microconfined conditions, with the highest expression observed 15 in cells on 10 and 100 kPa, with decreased expression in cells confined on 1 kPa hydrogels (Figure 1C, **D**). Substrate softness has been shown to favour endodermal differentiation.^{19,20} However, here we 16 17 see that confining hiPSCs on softer substrates created a multilayered endodermal structure which contained a small mesodermal cell cluster within. The SOX17 expression in colonies confined to 18 19 micropatterned hydrogels was observed as early as 12h after seeding (Figure S2), on both 250 µM 20 and 500 µM diameter micropatterned hydrogels, with the majority of the cells co-expressing the endodermal marker FOXA2 (Figure S3). To rule out artefacts associated with reprogrammed cells, we 21 22 cultured H9 human embryonic stem cells under the same conditions and observed a similar increased 23 SOX17 expression on 10kPa hydrogels (Figure S4).

A distinct but transient expression of T/BRACHYURY was also identified in a small population 24 of cells towards the colony centre, more commonly observed in the 10kPa condition, in about 40-25 26 50% of experimental replicates (Figure 1E), whereas distinct nuclear punctate were observed across 27 all replicates of confined 100kPa and 10kPa hydrogel conditions, with localisation towards the colony centre (Figure S5). We also stained for the ectodermal marker SOX2, and neuroectoderm progenitor 28 SOX1, with comparable negative staining across all confined hydrogel substrates. This shows that the 29 colonies on PA hydrogels were mostly mes-endodermal in identity, and that primitive 30 streak/gastruloid-like identity was instigated by the imposed biophysical microenvironment. To 31

- 1 ensure reproducibility, all results were reproduced in at least 6 technical replicates with a minimum
- 2 of 3 biological replicates for each experiment.

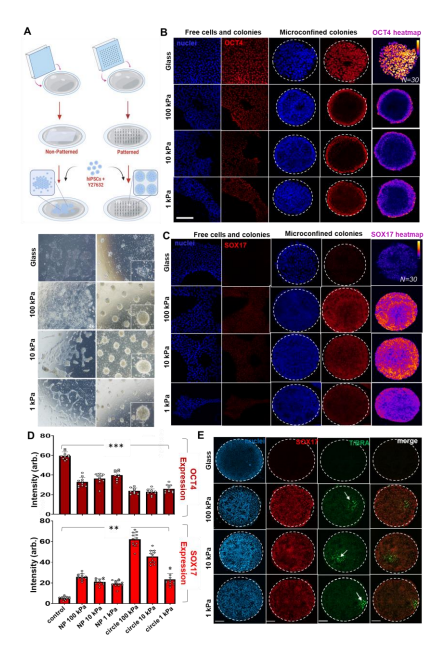


Figure 1: Substrate stiffness directs SOX17⁺ T/BRACHYURY⁺ mes-endodermal population. **A)** Schematic of PA hydrogel substrate preparation and cell seeding procedure, brightfield images from non-pattered and patterned 250µM colonies at 48 hours. All images acquired at 4x objective. **B)** hiPSCs seeded non-patterned and patterned glass and hydrogel substrates immunostained for Oct4. Right –immunofluorescence heatmaps of Oct4 expression. **C)** hiPSCs seeded non-patterned and patterned glass and hydrogel substrates stained for Sox17. Right –immunofluorescence heatmaps of Sox17 expression. **D)** Comparison of expression intensity for OCT4 and SOX17 across all conditions (N=8). *** = p<0.001,**=p<0.01 (Ordinary Oneway ANOVA) **E)** Immunofluorescence images of endoderm marker SOX17 and mesoderm marker T/Bra in cell populations on 500µM circular colonies. Scale Bars: 100um

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Previously we demonstrated how changes in perimeter curvature would influence the behaviour of microconfined cells, where geometry and stiffness both exerted an influence over cell

phenotype in the context of cancer stemness.^{21,22}. To evaluate whether geometry would play a role in the observed differentiation with our microconfined hiPSC colonies, we cultured cells in shapes of the same area approximating a star, flower, square, and capital 'l'—where positive and negative curvature and aspect ratio are varied—followed by immunostaining for germ layer markers. After 48 hours there were no significant differences between mes-endodermal marker expression across the shaped colonies, suggesting changes in geometry at the interface does not play a role in the observed gastrulation-like morphogenesis (**Figure S6**).

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9 Early adhesion stimulates YAP activity to coordinate epithelial-to-mesenchymal transition and 10 differentiation

Having observed how hydrogel microconfinement alone will trigger mes-endodermal differentiation, 11 we next sought to investigate how the surface directs this effect. Epiblast cells at the primitive streak 12 region are widely reported to have undergone EMT, to gain their mesenchymal and mes-endodermal 13 identity before they ingress in the gastrulating embryo²⁸. We immunostained microconfined colonies 14 on 10 kPa hydrogel and glass patterns for EMT molecular markers E-CAD, N-CAD and SNAIL, and OCT4 15 16 to gauge pluripotency. We selected the stiffness 10kPa for our experiments hereafter since we 17 observed a more consistent spontaneous appearance of the SOX17⁺ colonies with the T/BRACHYURY⁺ clusters on 10 kPa as compared to 1 and 100 kPa. As before, we observed OCT4 expression being 18 restricted towards the colony edges with decreased expression towards the centre (Figure 2 A,B). At 19 20 the same time, there is a striking loss of E-CAD in the colony centres, suggesting an EMT prone region starting inwards from the periphery. In conjunction with loss of E-CAD, the population confined on 21 hydrogels expresses uniform SNAIL, with N-CAD expression in the colony centre. This E-CAD to N-22 23 CAD switch is considered a prime indicator of cells undergoing EMT and is considered crucial for specification of the primitive streak and other embryogenesis events²⁹. In comparison, E-CAD and 24 OCT4 show uniform expression throughout the colonies on glass patterns, with no expression of 25 SNAIL (Figure 2A). E-CAD expression on the cell membrane and cytoplasm was discontinuous in 26 27 regions of the 250 µm colonies (Figure S7A). Moreover, E-cad expression was maintained in colonies across non-patterned surfaces, with only a few regions showing discontinuous or punctate E-CAD on 28 the non-patterned hydrogels (Figure S7B). 29

30 We observed a region at the colony edge with a SNAIL⁺ OCT4⁺ population, which is also the 31 region with highest SOX17 expression. We propose that this overlap denotes early stages in embryo development where SNAIL is reported to control EMT at the epiblast via downregulation of E cadherin³⁰. Similar SNAIL expression was also observed in the 250 μm colonies across all patterned
 hydrogels; however, there was no apparent spatial organization (Figure S7A). The E-CAD to N-CAD
 switch and concurrent SNAIL expression implicates EMT as the morphogenetic process that directs
 endodermal/mesodermal identity in the microconfined colonies on hydrogels.

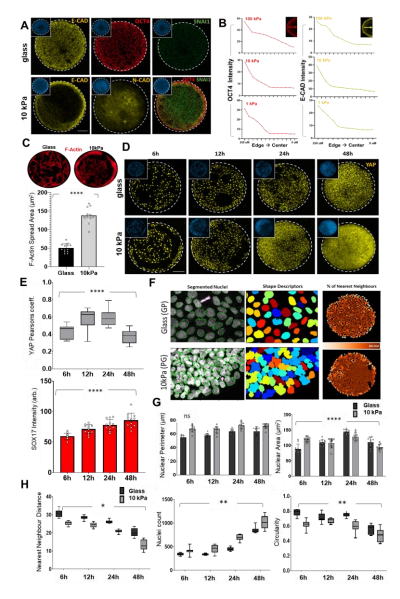


Figure 2: Early adhesion stimulates YAP activity to coordinate epithelial-to-mesenchymal transition and differentiation. A) Immunofluorescence images for E-CAD, N-CAD, SNAIL and OCT4 across colonies on glass and hydrogel patterns. B) Trace plots quantifying OCT4 and E-CAD expression on PA hydrogels from edge to centre. C) Actin spread area comparison for glass and 10kPa 500µM patterns. **** = p<0.0001 (N=13). D) Time course analysis of adhesion pattern and localisation of YAP on glass and 10kPa patterns. E) Quantification of YAP nuclear-cytoplasmic ratio (top) and corresponding SOX17 expression intensity (bottom) at four time points. **** = p<0.0001, ***=p<0.001, N=12 (Ordinary One-way ANOVA). F) Nuclei segmentation as primary objects, nuclei shape description, percentage depiction of nearest neighbours on glass and 10kPa 500µM patterns. *** = p<0.001,**=p<0.01(Two-way ANOVA) G) and H) Comparison of nuclei area and perimeter, distance between nearest neighbours, average cell count and averaged circularity of identified nuclei on glass and 10kPa 500µM patterns *** = p<0.001,**=p<0.01,*=p<0.05, ns=not significant (Two-way ANOVA). Scale bars = 100µM

Next, we analysed differences in cell adhesion and proliferation over time in our 1 microconfined cultures by fixing and staining at 6, 12, 24, and 48 hours with the same initial seeding 2 density. After initial seeding, cells encircle the border and adopt an elongated contractile morphology 3 4 (Figure S2) with elevated F-actin and higher spread area compared to cells in the interior of the pattern or those constrained in glass patterns (Figure 2C). Over time, these contractile cells 5 proliferate to fill the pattern which coincides with lessening of the observed cytoskeletal tension. 6 Confined cells show significantly higher proliferation on the hydrogel patterns compared to glass, 7 which at first glance is counterintuitive since soft matrices are known to limit cell proliferation. 8 9 However, ROCK inhibition with Y27632 has been reported to promote cell proliferation on soft matrices and supress proliferation on glass through modulating actomyosin contractility³¹. This is also 10 11 apparent through a nearest neighbours analysis (Figure 2 G,H). A core protein in sensing matrix 12 stiffness, and a driver of pluripotency maintenance, is the yes-associated protein (YAP)³². YAP activity is involved in controlling the expression of genes associated with the anterior primitive streak³³ and 13 14 has been indicated in coordinating germ layer patterning during *in vitro* BMP4 driven gastrulation³⁴. 15 YAP expression is predominantly nuclear within the first 12 hours, followed by a decrease as the cells proliferate from the perimeter to the centre (Figure 3 D,E). Monitoring SOX17 expression at the same 16 17 time showed that the transition of nuclear YAP to the cytoplasm corresponds with decreased cytoskeletal tension and increase SOX17 expression. During this time course we also measured 18 19 changes in nuclear geometry, where variations in morphology has been demonstrated during EMT, 20 differentiation and de-differentiation.^{35,36} Cells confined on 10 kPa hydrogels show increased nuclear perimeter over time with decreased circularity compared to cells confined on glass (Figure 3 F,G,H). 21 Overall, these results suggest that initial adhesion increases cytoskeletal tension at the boundary, 22 directing YAP activity and stimulating EMT, followed by differentiation to a mes-endodermal 23 24 population.

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Priming pluripotent stem cell colonies on hydrogels augments BMP4 induced germ layer specification

Previous work with micropatterned cultures used BMP4 as a soluble morphogen to initiate differentiation.^{10,11,15} We hypothesised that our cells on micropatterned hydrogels would be more susceptible to differentiation, or bias the population to different outcomes, after induction with BMP4 when compared to the standard condition of culture on glass. To test this, we seeded hiPSCs with BMP4 (50 ng/ml) induction to drive differentiation over 48 hours. In the brightfield images, the

colonies on the glass surface show homogenous distribution with some multilayering at the edges. 1 2 In contrast, the colonies on 10 and 100 kPa hydrogels showed distinct multilayered clustering 3 towards the centre with monolayer organization at the border (Figure 3A). Cells confined on 1 kPa 4 hydrogels did not show this multilayered characteristic. We immunostained all cultures for germ layer markers SOX17, T/BRACHYURY and SOX2. After BMP4 induction, the cells cultured on glass 5 showed periodic patterns of clustered and spread cells with a wave-like morphology, where the 6 7 edges of the clustered regions are rimmed with mes-endodermal SOX17⁺ T/BRACHYURY⁺ cells with slight regions of ectodermal SOX2⁺ (Figure 3B). This result demonstrated how BMP4 induction 8 9 promotes outer endoderm with adjacent mesoderm and ectoderm, with concentric organization when constrained in glass micropatterns.¹⁰ The cells seeded on non-patterned hydrogels similarly 10 11 showed wave-like multilayered texturing with increased density and increased mesoderm 12 specification observed on the 10 kPa hydrogels. Cells cultured on the non-patterned 1 kPa hydrogel 13 showed comparatively less attachment leading to smaller clusters of randomly placed mes-14 endodermal cells (Figure S8A). Strikingly, the concentric spatial patterning of germ layers in 15 microconfined colonies on glass was also observed on 500 µM patterned hydrogels, but with a considerable enhancement leading to a large multilayered cluster in the colony centre (Figure 3C). 16 17 These central clusters were dense with SOX17⁺ cells located towards the edge, intermingled with a population of T/BRACHYURY⁺ cells extending inward, with SOX2⁺ cells exclusively at the centre. We 18 19 observed similar characteristics in cell colonies in the smaller 250 µM diameter patterns (Figure S8B); 20 immunofluorescence quantitation revealed enhanced T/BRACHYURY+ mesoderm specification on the 10 kPa hydrogel micropatterns, confirming the previous observations of softer substrates 21 favouring mesodermal differentiation in response to BMP4¹⁵ (Figure 3D). 22

Since EMT processes were observed in our cultures without BMP4 (**Figure 2**), we also immunostained these cultures for E-cadherin (E-CAD) which demonstrated a pronounced loss towards the colony edges on glass, with the reverse observed for colonies cultured in confinement on hydrogels (**Figure S9A**). These trends in E-CAD expression correspond directly with the expression of markers associated with differentiation. Another characteristic of differentiation is changes in cell and nuclear area. The area and perimeter of the differentiated cell nuclei was considerably lower in clustered cells compared to spread cells across all conditions (**Figure S9B**).

To support our immunofluorescence result, we selected our optimal hydrogel condition (10 kPa) compared to glass for quantitative PCR to assess differences in transcript expression on account

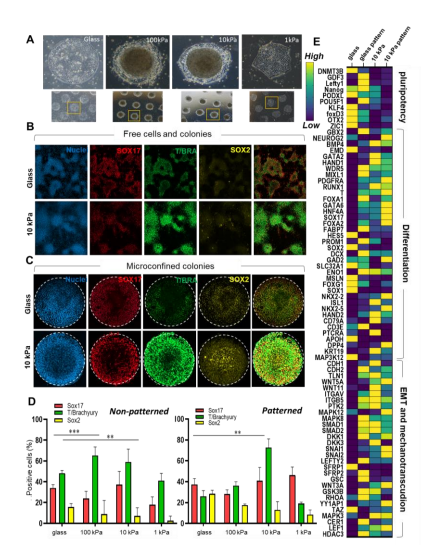


Figure 3: Priming pluripotent stem cell colonies augments BMP4 induced germ layer specification A) Brightfield images of individual representative colonies on each substrate. **B)** BMP4 induction leads to gastruloid-like organized cell clusters on glass and hydrogel substrates (Scale bar – 250μ M). **C)** Circular confinement centralizes the primitive streak-like cluster in the microconfined colonies (Scale bar – 100μ M). **D)** Quantification of the percentage of positive cells for the three markers on non-patterned glass and hydrogels (top) and patterned glass and hydrogels (bottom) *** = p<0.001,**=p<0.01)(Two-way ANOVA) **E)** Heatmap of quantitative expression of genes associated with pluripotency, differentiation, EMT and mechanotransduction (N=2).

1 of substrate stiffens, geometric confinement and both (Figure 3E). Consistent with the 2 immunofluorescence results for microconfined colonies on hydrogels, we observed decreased expression of pluripotency genes with an increase in differentiation markers towards mes-endoderm 3 lineages compared to colonies on glass-e.g., endoderm: SOX17 (x340), FOXA2 (x247) and GATA6 4 (x592), and mesoderm: T/BRACHYURY (x 43), RUNX1 (x33) and MIXL1 (x50). We also observed 5 increased expression of EMT regulators N-CAD (CDH2; x4), SNAIL (SNAI1; x13); SNAI2 (x163), and 6 downstream effectors of BMP signalling SMAD1/2 (x2) in the micropatterned colonies. Canonical 7 WNT signalling member WNT3A (x28) and its inhibitor DKK1 (x72) were also considerably elevated 8

on micropatterned hydrogels; both molecules playing central roles in controlling gastruloid pattern
 extension and meso- vs endo- differentiation²³. Nodal antagonist CEREBRUS 1 (CER1) was
 considerably elevated in micropatterned colonies (x430), suggesting a role contributing to primitive
 streak-like formation in microconfined cultures.

In addition to pluripotency, differentiation and EMT markers, microconfinement on hydrogels 5 led to increased expression of vitronectin binding integrin αV (x2) and $\beta 5$ (x4) and associated 6 downstream effectors of mechanotransduction, consistent with our observations of cytoskeletal 7 tension and YAP activity guiding morphogenesis. Concurrently, we saw an increase in non-canonical 8 9 WNT5A (x59) and WNT11 (x2), which are regulated by several mechanotransduction pathways including planar cell polarity which dictates patterning during embryogenesis in multiple species^{24,25}. 10 11 Furthermore, there was 28-fold higher expression of the WNT inhibitor secreted frizzled related protein (SFRP1) in colonies on glass, suggesting attenuation of WNT signals is a central aspect of 12 maintaining pluripotency. Previous reports demonstrate the importance of WNT in ES cell 13 differentiation²⁶ as well as the role of mechanics favouring mesoderm differentiation on compliant 14 surfaces^{15,27}. Together these results demonstrate how microconfinement on hydrogels enhances 15 EMT and WNT/Nodal activity which leads to increased differentiation upon treatment with BMP4. 16

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18 WNT signalling contributes to differentiation in spatially confined microenvironments.

Pluripotent stem cells seeded on hydrogel micropatterns experience edge stress that increases 19 20 cytoskeletal tension affecting YAP localisation, thereby initiating EMT and differentiation. While our gene expression analysis aligns with these mechanotransduction pathways guiding differentiation on 21 22 hydrogels, there were also considerable changes in numerous paracrine signalling pathways involved in embryogenesis. The formation of primitive streak *in vivo* is guided by the activity of TGF β /Nodal 23 as well as WNT/ β -Catenin pathways^{15,23}. Since these pathways have been shown to synergistically 24 push the epiblast cells towards primitive streak while blocking ectoderm differentiation³⁷, we next 25 investigated the use of small molecule pharmacological disruptors to attenuate WNT and TGFB 26 signalling. 27

28 Canonical WNT signalling involves β -catenin activity and is associated with mesodermal 29 differentiation,³⁸ definitive endoderm progenitors,⁴⁰ as well as EMT related gain of motility 30 supporting primitive streak.³⁹ Since our transcript analysis showed elevated WNT signalling, we 31 supplemented our cultures with the canonical WNT inhibitor IWP2. Treatment with IWP2 leads to complete loss of T/BRACHYURY expression in colonies on the 10kPa hydrogel. However, the average SOX17 expression remains unchanged compared to untreated (**Figure 4A**). This result suggests that canonical WNT signalling is critical for mesoderm but not endoderm differentiation. Our transcript analysis also showed increased expression of non-canonical WNT signals (WNT5A and WNT11), which have previously been shown to coordinate emergence of the primitive streak in model animals.⁴¹⁻⁴³ To impede non-canonical WNT signalling, we selected the soluble protein secreted frizzled related protein 1 (sFRP1) which will bind all extracellular WNT signals, thereby impeding both

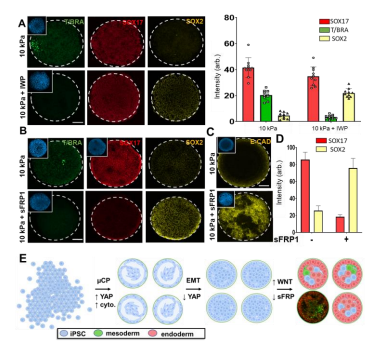


Figure 4 Pharmacological inhibition of canonical and non-canonical Wnt signalling disrupts differentiation. A) Immunofluorescence images of micropatterned iPSCs on 10kPa hydrogels with and without canonical WNT inhibition (IWP2). Right – corresponding quantitation. **B)** Immunofluorescence images of micropatterned iPSCs on 10kPa hydrogels with and without total WNT inhibition (sFRP1). **C)** E-cadherin expression with and without sFRP1 treatment. **D)** Quantification of SOX17 and SOX2 expression with and without sFRP1 supplemented media. **E)** Schematic depicting the mechanism where confinement and stiffness promote cytoskeletal tension and YAP activity, EMT and differentation. Scale bar - 100µM

canonical and non-canonical pathways. Cells were seeded on patterned 10 kPa substrates of 500 μm
diameter with and without soluble SFRP1 (5 μg/ml) for 48h. Immunofluorescence staining of cultures
treated with SFRP1 showed maintenance of pluripotency through SOX2, comparative maintenance
of E-CAD and no T/BRACHYURY or SOX17 expression (Figure 4B,C,D). In contrast to using the
canonical WNT disruptor IWP2, the use of SFRP1 to deplete WNT molecules in the media abolished
mes-endodermal differentiation. This suggests the biophysical microenvironment promotes *in vitro*gastrulation through mechanotransduction initiated non-canonical WNT signalling (Figure 4E).

We also probed the effect of small molecule modulators during BMP4 induction by 1 investigating mes-endodermal differentiation on glass and 10kPa patterns in the presence of several 2 3 modulators of these pathways including the GSKβ antagonist CHIR99021 (CHIR), ALK4/5/7 inhibitor 4 SB431542 (SB) and IWP2 (Figure S10). Treatment with CHIR abrogated spatial patterning in the confined cultures, led to loss of SOX17 expression with a decrease in T/BRACHYURY. Addition of SB 5 along with CHIR results in a further decrease in T/BRACHYURY⁺ cells, demonstrating how combined 6 Nodal and WNT signalling is important for mesodermal differentiation⁴⁴. Addition of SB, either alone 7 or in combination with a WNT activator or inhibitor was expected to abolish endodermal 8 differentiation, as endoderm differentiation is shown to require both WNT and Nodal activity⁴⁵. 9 10 However, a small population of T/BRACHYURY⁺ cells persisted in CHIR+SB and SB conditions, 11 consistent with WNT signalling being indispensable for mesoderm specification⁴⁵. Blocking both WNT 12 and Nodal in SB+IWP conditions abolished all mes-endodermal differentiation following BMP4 13 induction, demonstrating the shared role in BMP4 induced gastruloid formation. In the 14 microconfined colonies on 10kPa hydrogels, we observed partial maintenance of SOX17⁺ populations 15 with CHIR and CHIR+SB treatments compared to the sharp decline for colonies on glass in the same conditions (Figure S10). 16

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18 Primed colonies form gastruloid-like structures in 3D engineered niches.

Since microconfined hiPSCs on gels promote mes-endodermal differentiation without exogenous 19 20 stimulation, we sought to discern subsequent spatial morphology and organization for gastruloid-like 21 structures harvested from the substrates. After culture for 48 hours, primed embryoids were lifted off the PA surface and individually cultured for 14 days in mTeSR media. To control for the primed 22 conditions, cells were seeded in non-adherent plates to form embryoid bodies (Figure 5A). The 23 embryoid bodies show uniform spherical growth while the gastruloids show an unrestricted 24 25 structural growth from the periphery over two weeks (Figure 5B). At day 7, we observe a distinct loss of OCT4 expression in the growing gastruloid compared to embryoid, which is further downregulated 26 by day 14 (Figure 5C). Immunostaining the gastruloid for SOX2 (Ectoderm/ Epiblast cells), SOX17 27 28 (Endoderm) and T/BRACHYURY (Mesoderm/Primitive Streak) shows loss of OCT4⁺ regions at the interface, with partial SOX2⁺ staining and hotspots of SOX17⁺ and T/BRACHYURY⁺, which is not 29 30 observed in the embryoids (Figure 5D). The appearance of positional primitive streak-like populations for gastruloids is reminiscent of embryonic gastrulation; however, the magnitude and directionality 31 32 of the outgrowths vary across samples.

Considering the dynamically changing microenvironment surrounding embryonic tissue 1 2 during gastrulation, we hypothesized that encapsulation of the gastruloids in designer biomaterials 3 that vary ligand presentation, viscoelastic properties, porosity, and diffusion, could be used to guide 4 spatiotemporal patterning. The following hydrogel biomaterials were selected to encapsulate the harvested gastruloid: gelatin methacrylate (GelMA), methacrylated hyaluronic acid (MeHA) and 5 alginate. At day 7, the multi-lobed gastruloids spread out and became more spherical like the 6 embryoid bodies when confined within photocrosslinked gelatin methacrylate (GelMa) or 7 methacrylated hyaluronan (MeHa) with loss of primitive streak hotspots (Figure 5E,F). However, 8 encapsulation within non-covalently stabilised alginate demonstrates elongation of the structure 9 10 with converging SOX17⁺ and T/BRACHYURY⁺ region, reminiscent of the appearance of a streak like 11 population with mesodermal and endoderm progenitor cells for the initiation of human gastrulation. 12 These results highlight how covalent hydrogel networks (e.g., GelMA, MeHa) may inhibit 13 differentiation and limit further development, while viscoelastic materials that accommodate cell 14 migration and reorganization (e.g., alginate) may be well suited to further guide morphogenesis.

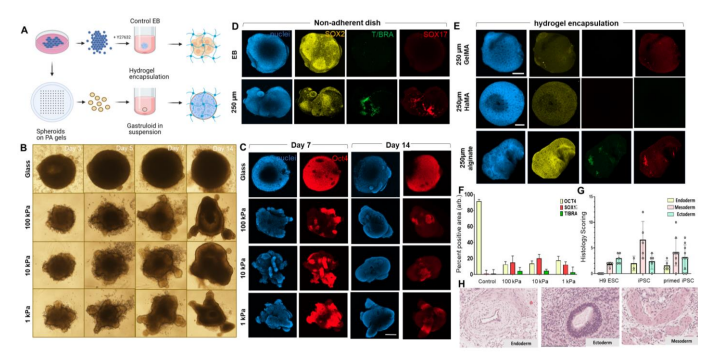


Figure 5: Release from patterns facilitates *in vitro* **gastruloids. A)** Scheme for gastruloid formation and encapsulation within hydrogels. **B)** Brightfield images of embryoid bodies and hydrogel primed gastruloids grown in non-adherent plates. **C)** Immunofluorescence images of embryoid bodies (glass) and gastruloids immunostained for Oct4 at 1 and 2 week culture in non-adherent plates **D)** Immunofluorescence images of embryoid bodies of embryoid bodies and gastruloids on non-adherent plates and **E)** after encapsulation for 14 days in hydrogel biomaterials. **F)** Percentage of positive area for the OCT4, SOX17 and T/BRACHYURY in embryoid bodies and hydrogel cultured gastruloids. **G)** Histology scoring for H&E scans of teratoma sections from H9 hESCs, hiPSCs and primed hiPSCs. **H)** Representative H&E images for germ layer derivatives identified in the teratoma from (tissue from primed iPSCs shown here). Scale bar - 100μM

Detailed live cell imaging during culture will be necessary to accurately assess growth and morphogenesis within 3D hydrogel matrices. Nevertheless, our results demonstrate how materials can trigger and maintain gastrulation-like states in pluripotent stem cell populations.

4 To support our *in vitro* differentiation results, we performed a subcutaneous teratoma assay using immunocompromised SCID mice with control pluripotent stem cells (hESC and iPSC), and cells 5 from 10 kPa hydrogels and followed the teratoma growth for 7 weeks. There was high variability in 6 7 growth across all conditions with no discernible trend in size across starting cell types (Figure S10A). 8 To determin e if the implanted cells undergo differentiation *in vivo*, we performed histology scoring 9 to classify structures associated with different germ layers. We observed duct-like, cartilaginous, bone, loose mesenchyme, smooth muscle, neural rosettes/neuroectoderm, pigmented epithelium, 10 and squamous epithelial-like structures. While the overall histology scoring indicates no significant 11 12 difference across the cell types (Figure 5G,H), there were subtle variations in the occurrence of specific structures that may be related to the primed mes-endodermal population (Figure S11). 13 14 Future work will be invested into tracking the differentiated progeny after implantation. However, this is outside of the scope of the present study. 15

16

17 **Discussion**

Bioengineered *in vitro* models have become powerful tools to study relationships between biophysical microenvironments and intracellular signalling pathways that align to specify cell fate in populations of pluripotent stem cells^{8,23}. However, these models rely on exogenous addition of soluble factors to trigger differentiation. We have demonstrated how careful selection of cell culture materials can serve to guide morphogenesis and foster conditions to drive *in vitro* gastrulation independent of exogenous chemical induction.

24 After initial attachment to the micropatterned islands on hydrogels, cells navigated to the 25 regions of elevated stress at the periphery which led to increased cytoskeletal tension and high 26 nuclear YAP localisation. Over time we see enhanced proliferation on the soft hydrogels, which is 27 consistent with previous reports of cells treated with ROCK inhibitor Y27632.³¹ This is in sharp 28 contrast to the populations confined on glass which show uniform adhesion and proliferation. These early aggregates on hydrogels create an epithelial-to-mesenchymal interface, with subsequent 29 differentiation to a mes-endodermal SOX17⁺ T/BRACHYURY⁺ population in the central region. This 30 behaviour of contractile cells at the boundary coordinating EMT has been observed in vivo⁴³. We 31

propose that hydrogel microconfinement provides analogous initial conditions where the interfacial
 stress leads to EMT, elevated WNT signalling, and formation of primitive streak-like populations.

3 The cells lining the perimeter of the colonies co-expressed OCT4, SOX17 and E-CAD, whereas the central part of the colonies showed loss of both OCT4 and E-CAD but with an increase in 4 expression of mes-endodermal markers SOX17 and T/BRACHYURY. The co-expression of OCT4 and 5 SOX17 at the edge suggests partial differentiation consistent with early phases of endoderm 6 specification.^{46,47,48} SOX17 is one of the first transcription factors to be expressed in the inner cell 7 mass^{50,51} and regulates the dynamics of pluripotency and differentiation.⁵² Moreover, T/BRACHYURY 8 marks the first mesodermal cells that ingress in the primitive streak of a gastrulating embryo.^{53,54} The 9 patterned phenotypes observed under confinement on hydrogels bears a resemblance to the self-10 initiation of primitive streak at the onset of gastrulation. 11

12 There is considerable evidence to support a central role for biophysical cues directing embryogenesis.^{6,12} Embryonic epiblast is essentially a layer of epithelial cells tightly connected and 13 packed at the interface as the embryo implants itself on the maternal uterine lining⁵⁵. During 14 gastrulation, the cells at the posterior side of the embryo undergo epithelial-to-mesenchymal 15 transition (EMT)^{2,56}, a signalling gradient arises between BMP, WNT and Nodal pathways, and the 16 primitive streak appears at the same EMT-prone region^{1,57}. In vivo, the initiation of EMT precedes the 17 appearance of the first multipotent mes-endodermal progenitors at the primitive streak⁵⁶ which 18 19 triggers the epiblast layer towards a dynamic, mesenchymal identity at the onset of gastrulation⁴². 20 EMT leads to loss of apical-basal polarity with concurrent nuclear shape changes that contribute to distinct cell-ECM adhesion patterns and motility⁵⁸. These pre-gastrulation events have been shown 21 to be controlled by non-canonical WNT signalling via tight coordinated movements and intercalation 22 of cells to initiate a region prone to EMT⁴³ with subsequent mes-endodermal differentiation. 23

After treatment of microconfined colonies on hydrogels with BMP4, we see significant 24 enhancement of mes-endodermal differentiation with the appearance of multilayered 25 T/BRACHYURY⁺ nodes, the number and position of which is linked to confined area. Transcript 26 27 analysis indicates that upregulation of mes-endodermal genes is driven by mechanochemical signal transduction that converges with non-canonical WNT signalling. The non-canonical WNT pathways 28 are involved in coordinating EMT^{43,64} with planar cell polarity pathways guiding the mechanical 29 segregation of cells during gastrulation.^{41,65} Of the canonical WNT proteins, WNT5A in particular has 30 been linked to regulation of EMT⁴³, endodermal differentiation⁶⁷ and regulation of planar cell polarity 31

signalling ⁶⁸, where its loss is responsible for defects in embryological axis elongation and tissue-scale
 polarity during development.^{69,70}

3 Mechanotransduction propagates through integrin-mediated adhesion, actomyosin 4 contractility and mitogen activated protein kinase signalling, where numerous downstream effectors are employed by both matrix engagement and soluble morphogens²². Here we showed how adhesion 5 to deformable hydrogels leads to enhanced cytoskeletal tension and YAP activity at the boundary 6 7 which triggers EMT across the colony. Once EMT is underway, differentiation is coordinated through complementary paracrine/autocrine signalling, like the planar cell polarity pathway involving both 8 WNT5A and WNT11. WNT signalling controls differentiation during vertebrate development, and the 9 planar cell polarity serves as a route for organising tissue patterning^{24,25,42,71}. After treating our 10 cultures with small molecule inhibitors for WNT/β-catenin and TGFβ pathways we see some 11 moderate attenuation of mes-endodermal differentiation. The small molecule inhibitor IWP targets 12 canonical WNT signalling involving β -catenin translocation, and does not impede non-canonical 13 pathways. IWP treatment inhibits the expression of mesoderm marker T/BRACHYURY but leads to no 14 change in the expression of endoderm marker SOX17. In contrast, treatment of our cultures with 15 SFRP1, which will inhibit all WNT signalling in the extracellular space, leads to complete abolishment 16 of both markers associated with a primitive streak-like populations. Since differentiation still occurs 17 18 with inhibition of canonical WNT signalling, this result demonstrates a central role for non-canonical 19 WNTs in coordinating *in vitro* gastrulation.

Recently, there have been several reports of pluripotent stem cell colonies patterned on glass 20 substrates for *in vitro* models of neuroectoderm⁶², cardiac tissue¹⁸ and coordination of pre-streak 21 patterning.⁶³ Across all of these studies, geometric confinement was shown to guide cellular 22 assembly, where triggering differentiation with soluble morphogens would lead to biomimetic forms 23 with distinct functions. Weaver and colleagues showed how soft substrates will bias pluripotent stem 24 cells towards mesodermal differentiation following BMP4 induction, providing evidence for the 25 26 importance of matrix mechanics during lineage determination¹⁵. Previously, we showed how 27 combining deformable matrices with geometric confinement promotes EMT via mechanotransduction in populations of cancer cells leading to epigenetic reprogramming and 28 emergence of a stem cell phenotype.^{59,60,61} Here we show that confining pluripotent stem cells on 29 30 deformable hydrogels serves to guide cellular assembly and partitioning, thereby providing the right

context to trigger EMT and catalyse morphogenetic events that mirror *in vivo* processes of
 differentiation.

3 The ability to foster gastrulation-like events through the properties of the culture substrate 4 alone opens opportunities to study the relationships between the biophysical microenvironment and early embryogenesis. Colonies that have been primed for two days in microconfinement were 5 released from the substrate and cultured in suspension to see if further patterning would occur in 6 7 standard conditions for embryoid body growth. Compared to control embryoid bodies which exhibit a uniform morphology with robust staining of pluripotency markers, the primed colonies exhibited 8 9 clear regions of mes-endodermal identity which further evolved into lobed structures reminiscent of embryo patterning over the course of 14 days. The morphology of these gastruloids show similarities 10 11 to approaches involving small molecule stimulation of embryoid bodies⁷². Encapsulation of the gastruloids within a library of hydrogel biomaterials demonstrates materials parameters that 12 promote and prevent continued lineage specification. These finding underscore the importance of 13 materials in nurturing the differentiated phenotype during *in vitro* gastrulation, which draws parallels 14 to the role of the extracellular matrix in orchestrating embryogenesis in vivo. 15

16

17 Conclusions

Modelling human gastrulation in vitro through materials properties without exogenous biochemical 18 induction raises numerous opportunities for probing fundamental stages in embryogenesis. The 19 20 precise control of gastrulation in vivo is afforded by tight coordination of soluble signals with feedback from the biophysical microenvironment. Using microengineered hydrogel substrates, we 21 22 demonstrate a method where controlling the biophysical microenvironment poises a population of pluripotent stem cells to undergo morphogenesis, with integration of soluble signals to orchestrate 23 gastrulation-like processes. These findings provide a new avenue for probing the biophysical and 24 25 biochemical basis of embryogenesis, and a tool to model development for fundamental biology and 26 translational endeavours.

27

28 Methods

29 Cell culture and Maintenance

Hepatic fibroblast derived iPS line ATCC-HYS0103 Human Induced Pluripotent Stem (IPS) Cells were 1 purchased directly from the vendor. hESC-Qualified Matrigel (Corning 354277) Matrigel was used 2 3 coat culture dishes for feeder-free expansion of induced pluripotent stem cells and were routinely 4 cultured and maintained in mTeSR™1 (STEMCELL Technologies, 85850) in a humidified incubator at 37 °C with 5% CO₂. Cells were passaged once a week using selective dissociation reagent ReLeSR™ 5 (STEMCELL Technologies 05872) and seeded using the cell aggregate counting method described in 6 7 'Plating Human ES and iPS Cells Using the Cell Aggregate Count Method' (Appendix 1) from the STEMCELL Technologies-'Maintenance of Human Pluripotent Stem Cells in mTeSR™1' technical 8 9 handbook to assess the size of aggregates and seed them in low, medium or high densities, as 10 described. All cryopreservation of cells was performed in CryoStor[®] CS10 (STEMCELL Technologies 11 07930) freezing media.

For the glass controls in the experiments – sterile rh-Vitronectin (Gibco, Life Technologies, A14700) was used to coat glass cover slips. In case of all experiments, cells were dissociated into a single cell suspension using StemPro[™] Accutase[™] (Gibco A1110501) cell dissociation reagent to facilitate cell counting. In case of single cell dissociation, cells were always seeded with 10uM Rock inhibitor Y27632 (ATCC[®] ACS-3030[™]).

17 **Preparation of PA coated cover slips**

18 Hydrogel based substrates were prepared using chemically modified polyacrylamide and soft lithography. Round glass cover slips (18mm Diameter) were sonicated and individually placed in a 12-19 20 well tissue culture polystyrene plate, treated with 0.5% 3-Aminopropyl triethoxysilane (APTS) (Sigma Aldrich A3648) for 3 minutes then with 0.5% Glutaraldehyde (Sigma Alrich G6257) for 30 minutes. 21 22 The cover slips are thoroughly air dried with the treated surface up. For the polyacrylamide hydrogel coating, 40% solution of Acrylamide (Sigma Alrich A3553) and 2% solution of Bisacrylamide (Sigma 23 24 Aldrich 146072) were prepared in distilled water. Solutions pertaining to various elastic modulus were prepared as described in the table 73 . The stiffness solution was sandwiched between a 25 hydrophobic glass slide and the treated cover slip. 10 % Ammonium Persulfate (APS) and 26 27 Tetramethylethylenediamine (TEMED) (Sigma Aldrich, 1.10732) were used for polymerization in a covered, moist environment. Cover slips were carefully picked up then treated with Hydrazine 28 29 hydrate 100% (Acros organics 196715000) for up to 1 hour to convert amide groups in polyacrylamide 30 to reactive hydrazide groups and then 1 hour incubation in a 5% solution of Glacial acetic acid is done before patterning. DI washes performed between each chemical treatment. 31

1 Microcontact Patterning of PA coated cover slips

2 For microcontact patterning, polydimethylsiloxane (PDMS, Polysciences, Inc.) stamps for 250u and 3 500u diameter circles were prepared by polymerization upon a patterned master of photoresist (SU-4 8, MicroChem) created using UV photolithography using a laser printed mask. Sodium periodate (Univar 695-100G) solution was used to yield free aldehydes in the proteins used for micropatterning. 5 For assisting adhesion of iPS cells, recombinant human Vitronectin (Gibco A14700) was used at a final 6 7 concentration of 25ug/ml. The solution is applied to the patterned or non-patterned PDMS stamps for 30 minutes, air dried and then applied to the air-dried PA surface. The stamps are removed after 8 9 15 seconds and the patterned cover slips are sterilized by transferring to a sterile 12-well TCP dish 10 inside BSCII and 3x Sterile DPBS wash, followed by 12 minutes of UV exposure and then stored in 4°C soaked in a 2% Penicillin/Streptomycin + DPBS solution for minimum 6 hours. The soaking solution 11 to be removed and warm expansion media to be added to the cover slips before seeding the cells. 12

13 Microcontact patterning of glass cover slips

The same PDMS stamps were used to assisnt with physical adsorption of protein on clean, dry glass cover slips. The glass cover slips were sonicated in 100% ethanol and cleaned in a plasma cleaner (PlasmaFlo, Harrick Plasma) to remove all residue. Vitronectin was dissolved in PBS at the final concentration of 25µg/ml and applied on clean stamps 30mins/37°C. The stamps were rinsed, dried and applied on clean cover slips for 5-7 minutes allowing for protein adsorption on defined islands. The patterned glass cover slips were sterilised before cell culture and used within 24 hours of preparation.

21 Cell seeding on the micropatterned cover slips

The cell culture dishes are monitored for confluency before starting the experiment. All experiments 22 were performed using in mTeSR™1 (STEMCELL Technologies, 85850). The cells are dissociated using 23 24 warm StemPro[™] Accutase[™] (Gibco A1110501) cell dissociation reagent for 6-7 minutes. Cells were counted using a haemocytometer and seeded in mTeSR™1 (STEMCELL Technologies, 85850) and a 25 26 density of 5x10⁵ cells/ml as a 1ml/well solution in a 12-well culture dish with 10uM Rock inhibitor 27 Y27632 (ATCC[®] ACS-3030[™]). The medium was replaced with media without Y-27632 at 24h for 10 and 100kPa substrates, and 5uM Y-27632 in case of 1 kPa substrates (This was completely removed 28 at 36h). The cells were allowed to grow on the 4 substrate groups for 48h. 29

For BMP4 induction experiments, same seeding method was used, fresh media supplemented with
rhBMP4 (STEMCELL Technologies, 78211) at a final concentration of 50ng/ml on the 4 substrate
groups for 48h.

For small molecule inhibitor treatments- cells were supplemented with the following concentration
of each small molecule at 6h of seeding– CHIR99021 (3uM), SB431542 (5uM), IWP-2 (2.5uM), sFRP1
(Eug/ml)

6 (5ug/ml).

7 Immunocytochemistry – fixation and staining

Cell media was replaced with 4% solution of Pierce[™] 16% Formaldehyde (w/v), Methanol-free 8 (Thermo Fisher Scientific, 2890) diluted in 1X PBS, for 30 minutes at RT. All washes were performed 9 using Gibco[™] DMEM, no phenol red (Gibco, 31053028). This was done because any contact with PBS 10 would readily detach all cells from the patterned PA surfaces. 0.1% solution of Triton X-100 in PBS 11 12 (v/v) and 3% Bovine Serum Albumin solution (Sigma Aldrich, A3803) in PBS (w/v) was used for 13 permeabilization and blocking. Primary incubation 4°C overnight/secondary incubation for 1 hour at RT in dark. Coverslips were mounted using ProLong[™] Gold Antifade Mountant (Thermo Fisher 14 Scientific, P36930) which contains a DAPI stain. 15

16 **3D Spheroid culture and Immunostaining**

At 48h, the media on the PA gels was removed, and PBS washes were performed 2-3 times, whilst 17 18 collecting the PBS after each wash in a separate labelled tube for each condition. As mentioned above, PBS washes would readily dissociate all colonies, but the spheroid structure of the colonies 19 are maintained. 50ul of warm mTeSR™1 was added to a 96-well Low adherence U-bottom dish. 20 Individual spheroids are picked up and deposited in the 96 well. After checking each well for only one 21 22 spheroid each, the wells are topped up with 100ul of media. The spheroids are grown for 14 days, 23 with media change done every other day. Similarly, spheroids were deposited in the biomaterials and cultured for 7 days. 5000-15000 cells deposited in each well with Y27632 supplemented for controls, 24 cultured 2 days before being deposited in the biomaterials. 25

For fixation, a cut 200ul tip is used to pick up the spheroid and transferred to a glass-bottom 96 well plate and fixed with 4% PFA (Thermo Fisher Scientific, 2890) for 24 hours. 2x PBS washes of 1 hour each was done followed by 0.2% solution of Triton X-100 for 2 hours and blocking for 1 hour. Primary and secondary antibody+Hoescht staining was performed for 24h each on a slow rocker at RT with
 2x1 hour PBS washes in between.

3 In vivo teratoma analysis

4 The intact spheroids were collected off the 10kPa PA gels via PBS washes as described above. The 5 spheroids were allowed to settle in the PBS for 10 minutes at RT, supernatant was removed, and the 6 spheroids were resuspended in 100µl Matrigel-GFR (Corning 354230). ATCC hiPSCs and H9 hESCs 7 were used as controls. This was injected in immunocompromised SCID mice subcutaneously and was observed for growth for the next 7 weeks. The teratomas were collected in ice-cold PBS, fixed in 10% 8 9 neutral-buffered formalin solution for 3 days, and transferred in 70% ethanol for 3 days. Paraffin embedded sectioning, mounting and haematoxylin/eosin staining was performed for all sections. 10 11 Imaging of slides was done using Leica Aperio XT slide scanner. Histology scoring was plotted using 12 Graphpad Prism.

13 RNA isolation and qRT-PCR

14 For RNA isolation, cells on all four groups of substrates were dissociated using StemPro[™] Accutase[™] 15 (Gibco A1110501) for 5 minutes. Cell pellets were washed with RT PBS and then with chilled PBS while being centrifuged at 4C. All existing PBS is removed, cell pellets were snap-frozen using liquid 16 17 nitrogen and stored in -80. RNA isolation is performed using RNeasy Mini Kit (Qiagen 74104) as per 18 kit instructions. cDNA prep was performed using RT2 First Strand Kit (Qiagen 330401) as per kit 19 instructions. The cDNA was added to RT² SYBR Green ROX qPCR Mastermix (Qiagen 330520) and the solution was added to a Custom RT2 PCR Array 384-well plate and qPCR program was run using 20 21 QuantStudio[™] 7 Flex Real-Time PCR System (Applied Biosystems 4485701). Data analysis was 22 performed on <u>https://geneglobe.giagen.com/us/analyze</u>. Heatmaps prepared on GraphPad Prism 23 and Morpheus.

24 Confocal imaging and data quantification

The fluorescence imaging was performed on a Zeiss LSM780 or a Zeiss LSM800 confocal microscope, normally using 10x and 20x objectives and images were acquired using the Zen Pro imaging software from Zeiss. Image analysis was performed using FIJI (Fiji is just ImageJ) software, and extra plugins were downloaded when required. Nuclear characteristics analysis was performed using CellProfiler Cell image analysis software from Broad Institute. Yap localisation and some nuclear measurements

- 1 were performed using a MATLAB based GUI prepared for our data. Data sets were compiled in
- 2 Microsoft Excel and graphs preparation and statistical analysis was done using GraphPad Prism.
- 3

4 Author Contribution

PS, JP and KK developed the ideas. PS, SR, JI, TM, SN, PJ, AY, EP and VC performed the experiments
and analysed the data. All authors contributed to writing the manuscript.

7

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- 14

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Supplementary Information

Defined microenvironments trigger *in vitro* gastrulation in pluripotent stem cells

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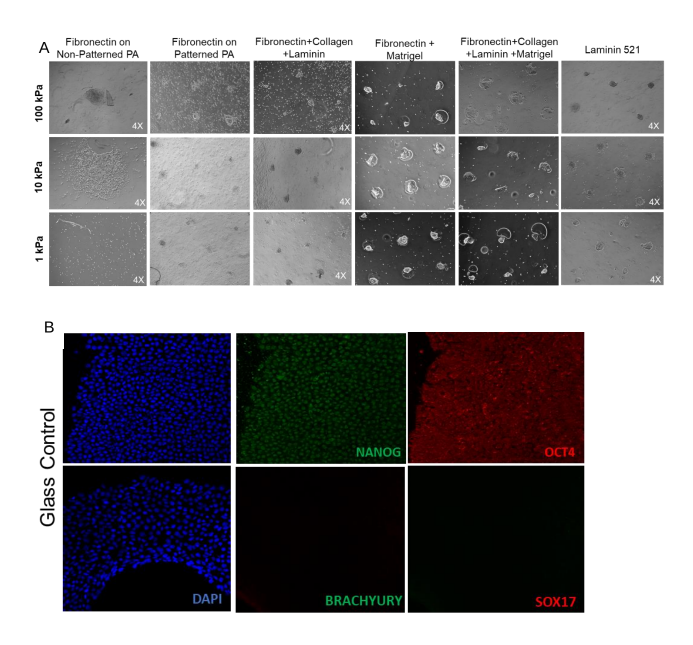


Figure S1: A) The panel of various ECM proteins tested for cell adhesion on polyacrylamide hydrogels, shows inadequate adhesion and proliferation in all combinations at 48 hours.

B) ATCC hiPSC characterisation

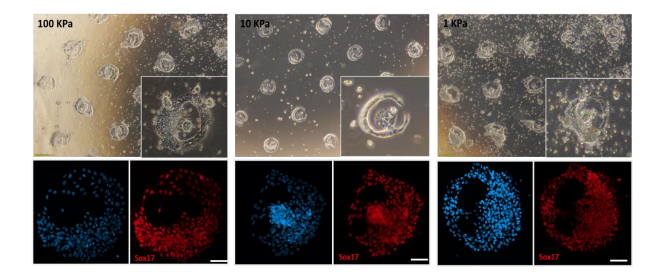


Figure S2: Brightfield and IF images of hiPSCs seeded on PA gel substrates with 250um circular patterns at 12 hours. The cells seem to circle the colony boundary before depositing towards the colony centre. The cells were seen to be of Sox17+ endodermal identity as early as 12h post seeding.

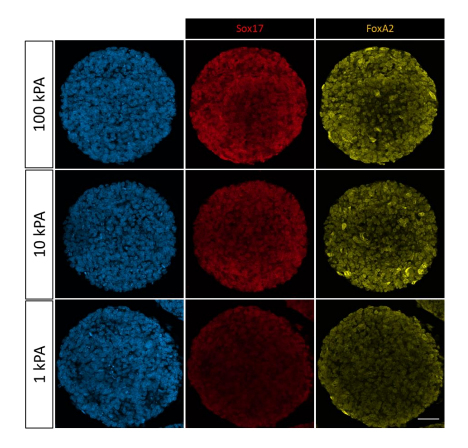


Figure S3 – Colonies 250uM circles on PA gels positive for Sox17 as well as FoxA2 Scale bar – $50 \mu m$

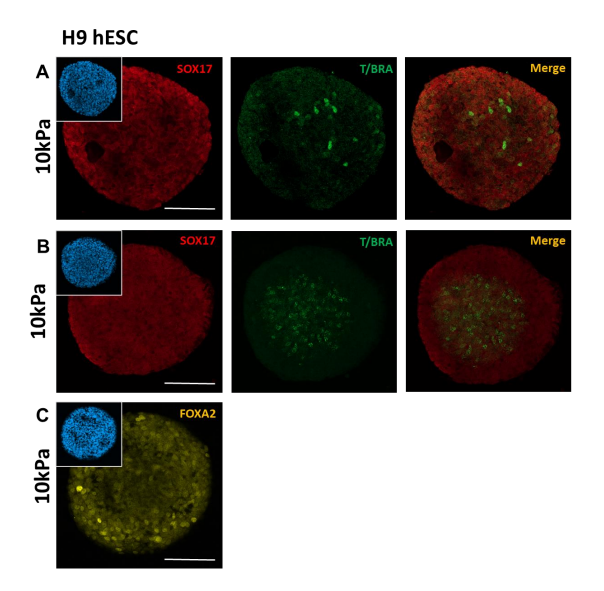
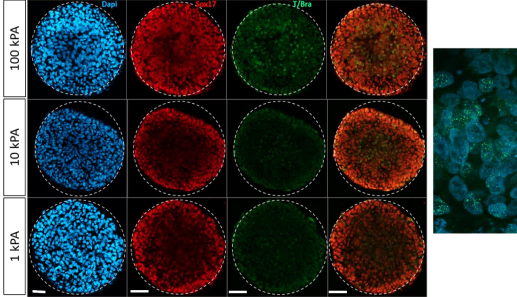


Figure S4: Immunofluorescent images of H9 hESCs seeded on 10kPa PA gel substrates with 250um circular patterns at 48 hours. A) The hESCs assumed an endodermal SOX17+ identity with a mesodermal T/BRACHYURY+ cluster within. B) In absence of a mesodermal cluster, nuclear puncta were observed in colony centres in response to T/BRACHYURY staining. C) Majority of SOX17+ cells co-expressed endodermal marker FOXA2



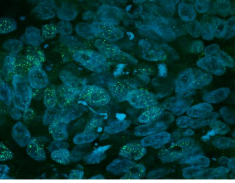


Figure S5: Stiffness and geometrical confinement create distinct nuclear punctates for Mesoderm/Primitive Streak marker T/Brachyury, regularly observed in the cells towards the centre of the hiPSCs colonies seeded on PA hydrogel substrates with circular patterns of 500µm diameter. Right – 63x image to show nuclear puncta.

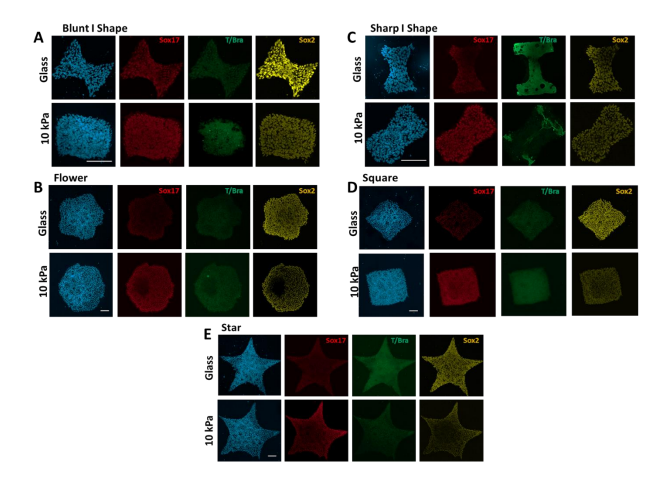


Figure S6: Immunofluorescent images of ATCC hiPSCs seeded on glass 10kPa PA gel substrates patterned using various shapes at 48 hours. A) Blunt 'I' shape B) Flower C) Sharp 'I' shape D) Square, and E) Star.

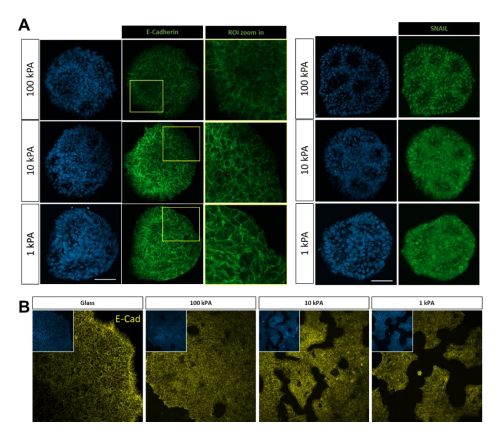


Figure S7: EMT Signifiers on non-BMP4 experiments A) Discontinuous regions of E-Cadherin and Expression of EMT marker snail in the 250um diameter colonies: signifies occurrence of EMT: Significant discontinuity and cytoplasmic localization of E-Cadherin in the colonies on the PA hydrogel substrates. Global Expression of SNAIL on the 3 stiffness substrates, however the clearest expression is observed on 100kPa. Both these act as an important signifier of EMT.

B) E-cad expression on glass and Non-Patterned gels

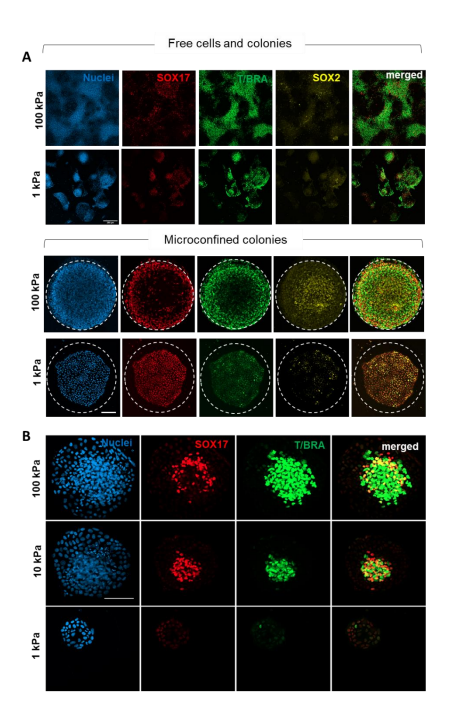


Figure S8: A) Immunofluorescence markers for mes-endodermal differentiation for iPSCs cultured on 1 kPa and 100 kPa hydrogels. B) Immunofluorescence markers for mes-endodermal differentiation for iPSCs cultured on 250 µm² islands.

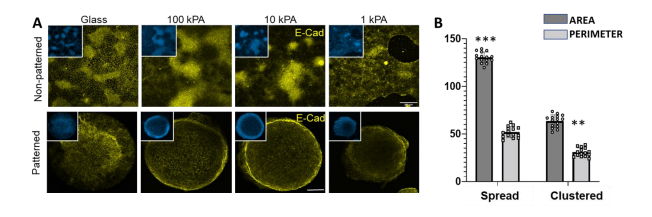


Figure S9: A) E-cad expression pattern on BMP4 treated non–patterned and patterned colonies. B) Nuclear size comparison for spread and clustered cells after 48hr treatment with BMP4 (N=15)

Scale bars: 100µm

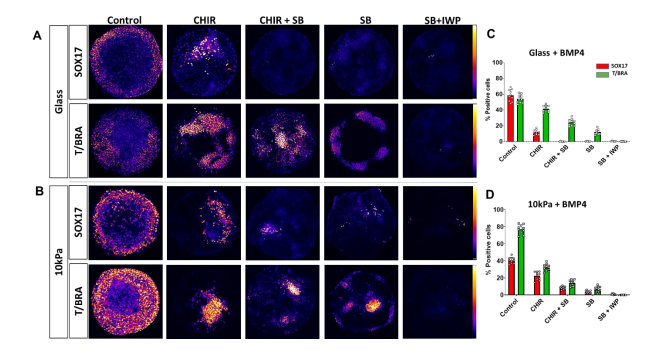


Figure S10: Small molecule inhibition on BMP4 sets: **A)** Heat maps for glass patterns at 48 hours – combinations of small molecule inhibitors for Wnt and Nodal SB = SB431542, IWP2 **B)** Heat maps for 10 kPa patterns at 48 hours – combinations of small molecule inhibitors for Wnt and Nodal **C)** Percent positive quantification for SOX17 and T/BRACHYURY in glass patterns + BMP4 N=10 p<0.001 **D)** Percent positive quantification for SOX17 and T/BRACHYURY in T/BRACHYURY in 10kPa patterns + BMP4 N=10 p<0.05

Scale bars - 100 µM.

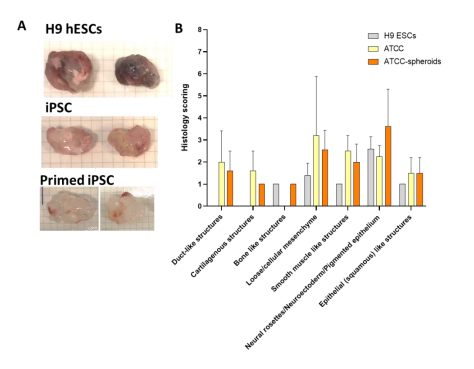


Figure S11: A) Comparative images for excised *in vivo* teratomas from H9 hESCs From ATCC hiPSCs from primed PA spheroids B) histology scoring of germ layer derivatives in all three groups

Table:1 Raw data -Gene list+Av double delta Ct values (2^Λ ΔCt) for +BMP4 gene expression analysis

	Average delta-	lelta Ct Values 2	?^(-Avg.(Delta(Ct))]
Symbol	Control Group	Glass pattern	10kPa Non-Pattern	10kPa Patterns	Pluripotency related
DNMT3B	0.083380	0.040025	0.026851	0.031173	Ectoderm
GDF3	0.006942	0.056342	0.004126	0.008220	Mesoderm
Lefty1	0.000734	0.016431	0.001607	0.002329	Endoderm
Nanog PODXL	0.000020	0.000022 0.051916	0.000015 0.048039	0.000007 0.017083	Ectoderm progenitors
POUSF1	0.126264	0.258880	0.173147	0.135897	Mesoderm Progenitor Endoderm Progenitor
CCDC42	0.000015	0.000016	0.000048	0.000022	EMT and Pathways
FGF5	0.000027	0.000011	0.000007	0.000033	Housekeeping Etc
foxD3	0.000936	0.000591	0.000030	0.000107	
SOX2	0.044362	0.007490	0.000327	0.000939	
SOX1	0.000053	0.000003	0.000003	0.000004	
OTX2	0.046724	0.030134	0.007147	0.017804	
ZIC1	0.000490	0.000088	0.000040	0.000059	
GBX2	0.000050	0.000807	0.000217	0.000493	
NEUROG2 BMP4	0.000009	0.000003	0.000003 0.027341	0.000016 0.023206	
SYNE2	0.003301 0.002741	0.010989 0.002934	0.003152	0.002040	
EMD	0.005133	0.002131	0.002345	0.001970	4
GATA2	0.000172	0.001306	0.006505	0.003623	
HAND1	0.000878	0.003483	0.050469	0.036176	
WDR5	0.005790	0.010967	0.009596	0.009469	1
MIXL1	0.003876	0.345962	0.099864	0.193306]
PDGFRA	0.000187	0.005592	0.004861	0.010295	
RUNX1	0.000023	0.000285	0.000858	0.000750	
T	0.001442	0.042410	0.041309	0.061455	
FOXA1	0.000004	0.000013	0.000005	0.000010	{
HDAC1 GATA6	0.015423	0.020934 0.020511	0.026255 0.014246	0.017377 0.032425	{
HNF4A	0.000055	0.020511	0.014246	0.032425	1
FOXA2	0.000004	0.000625	0.000190	0.000963	1
SOX17	0.000026	0.001798	0.003787	0.008814	1
LMNA	0.001139	0.001806	0.003998	0.002427	j
FABP7	0.000003	0.000011	0.000005	0.000005	
HES5	0.000029	0.000003	0.000006	0.000006]
PROM1	0.014451	0.029616	0.022208	0.013372	
DCX	0.000177	0.000351	0.000263	0.000261	
GAD2	0.000011	0.000012	0.000003	0.000017	
SLC32A1	0.000037	0.000026	0.000003	0.000008	
ENO1 MSLN	0.381987 0.000120	0.629827 0.000026	0.612041 0.000016	0.563835 0.000012	
FOXG1	0.000073	0.000054	0.000023	0.000038	
OLIG2	0.000015	0.000014	0.000006	0.000011	1
NKX2-2	0.000003	0.000015	0.000025	0.000066	
ISL1	0.000980	0.003851	0.036022	0.012290	1
NKX2-5	0.000012	0.000094	0.000220	0.000779	
HAND2	0.00008	0.000061	0.000133	0.000088	
CD79A	0.000028	0.000027	0.000034	0.000029	
CD3E	0.000006	0.00008	0.000004	0.000004	
PTCRA	0.000005	0.000004	0.000008	0.000004	
KRT19 APOH	0.047587	0.066626	0.293498	0.152485	4
DPP4	0.00003	0.000003	0.000003 0.000136	0.000004 0.000224	
MAP3K12	0.001645	0.000631	0.000910	0.000906	
CDH1	0.038705	0.062715	0.130371	0.047686	
CDH2	0.007641	0.038342	0.018475	0.028120	
VTN	0.000053	0.000024	0.000035	0.000031	
TLN1	0.007674	0.011206	0.012123	0.010122	
WNT5A	0.000741	0.005946	0.032960	0.043917	
WNT11	0.000007	0.000005	0.000030	0.000015	
ITGAV	0.005030	0.017435	0.026362	0.011416	{
ITGB5	0.019187	0.075126	0.078140	0.068715	{
PTK2 MAPK12	0.017327	0.022404 0.000916	0.023348 0.001040	0.020481 0.002247	{
MAPK12 MAPK8	0.011537	0.000916	0.018500	0.002247	1
SMAD1	0.004290	0.011979	0.012022	0.008974	1
SMAD2	0.009751	0.014924	0.015479	0.013362	1
DKK1	0.000383	0.029994	0.013634	0.027696]
DKK3	0.003045	0.003386	0.005925	0.003083	
SNAI1	0.000281	0.001145	0.001722	0.003606	
SNAI2	0.000127	0.001293	0.008024	0.020658	
LEFTY2	0.000345	0.006870	0.002637	0.002865	
SFRP1	0.052326	0.001923	0.002485	0.001780	
SFRP2	0.041860	0.089382	0.039669	0.037022	
GSC W/NT3A	0.000095	0.008685 0.000012	0.000654 0.000061	0.001557 0.000096	{
WNT3A GSK3B	0.007835	0.000012	0.013754	0.000096	1
RHOA	0.060488	0.014126	0.062518	0.055504	1
YY1AP1	0.006157	0.006609	0.006532	0.007376	1
TAZ	0.003490	0.001321	0.001933	0.002140	j
МАРКЗ	0.002751	0.002145	0.004202	0.004113]
CER1	0.000343	0.408546	0.054540	0.147443]
LEF1	0.001305	0.008189	0.013166	0.015809	
HDAC3	0.019248	0.024767	0.022489	0.020884	
АСТВ	1.554094	2.481468	2.676260	2.164344	
B2M	0.017275	0.030518	0.052066	0.033520	
GAPDH	0.482533	0.590769	0.613479	0.484756	}
HPRT1	0.005505	0.009302	0.005799	0.006078	}
RPLP0 GDC	1.985474	1.653981	1.598363	1.977917	{
U.S.L.I.	0.00003	0.000006	0.000003	0.000004	1

Table 2: Raw data -Gene list+fold-change and fold regulation values for +BMP4 gene expression
analysis

	Fold Change Va	lues (compar	ing to control
	group-Glass)		
Cumbel	Glass Patterns		10kPa Pattern
Symbol DNMT3B	Fold Change 0.48	Fold Change 0.32	0.37
GDF3	8.12	0.59	1.18
Lefty1	22.40	2.19	3.17
Nanog	1.09	0.73	0.35
PODXL	0.68	0.63	0.22
POU5F1	2.05	1.37	1.08
CCDC42	1.09	3.22	1.50
FGF5	0.39	0.25	1.19
foxD3 OTX2	0.63 0.64	0.03 0.15	0.11 0.38
ZIC1	0.84 0.18	0.15	0.12
GBX2	16.06	4.32	9.82
NEUROG2		0.40	1.90
BMP4	3.33	8.28	7.03
SYNE2	1.07	1.15	0.74
EMD	0.42	0.46	0.38
GATA2	7.59	37.79	21.05
HAND1	3.97	57.48	41.20
WDR5 MIXL1	1.89 89.27	1.66 25.77	1.64 49.88
PDGFRA	29.88	25.98	55.01
RUNX1	12.37	37.28	32.57
T	29.41	28.64	42.61
FOXA1	3.25	1.27	2.46
HDAC1	1.36	1.70	1.13
GATA6	374.51	260.12	592.06
HNF4A	1.56	1.42	2.12
SOX17	69.34	146.05	339.95
LMNA	1.59	3.51	2.13
FABP7 HES5	3.30	1.58	1.57 0.22
PROM1	0.09 2.05	0.19 1.54	0.22
SOX2	0.17	0.01	0.02
DCX	1.99	1.49	1.48
GAD2	1.09	0.31	1.56
SLC32A1	0.68	0.09	0.22
ENO1	1.65	1.60	1.48
MSLN	0.22	0.14	0.10
FOXG1	0.73	0.31	0.52
OLIG2	0.94	0.38	0.73
NKX2-2	4.33	7.26	19.62
HAND2 CD79A	8.10 0.96	17.63 1.21	11.59 1.01
CD3E	1.40	0.70	0.60
PTCRA	0.95	1.75	0.79
KRT19	1.40	6.17	3.20
APOH	0.80	1.01	1.08
DPP4	0.29	0.20	0.33
MAP3K12		0.55	0.55
CDH1	1.62	3.37	1.23
CDH2 VTN	5.02 0.45	2.42 0.65	3.68 0.59
TLN1	1.46	1.58	1.32
WNT5A	8.03	44.49	59.28
WNT11	0.64	4.13	2.11
ITGAV	3.47	5.24	2.27
ITGB5	3.92	4.07	3.58
PTK2	1.29	1.35	1.18
MAPK12	0.62	0.71	1.52
MAPK8 SMAD1	2.01 2.79	1.60 2.80	1.61 2.09
SMAD1	1.53	1.59	1.37
ISL1	3.93	36.76	12.54
NKX2-5	7.65	17.89	63.31
DKK1	78.37	35.62	72.37
DKK3	1.11	1.95	1.01
SNAI1	4.07	6.12	12.82
SNAI2	10.19	63.26	162.86
LEFTY2 SFRP1	19.88 0.04	7.63 0.05	8.29 0.03
SFRP1	2.14	0.95	0.88
GSC	91.66	6.90	16.44
WNT3A	3.54	18.09	28.22
GSK3B	1.80	1.76	1.19
RHOA	1.14	1.03	0.92
YY1AP1	1.07	1.06	1.20
TAZ	0.38	0.55	0.61
KLF4	0.51	0.56	0.44
MAPK3	0.78	1.53	1.50
FOXA2 SOX1	160.08 0.06	48.81 0.06	246.83 0.07
CER1	1191.87	159.11	430.14
LEF1	6.28	10.09	12.12
HDAC3	1.29	1.17	1.09
ACTB	1.60	1.72	1.39
B2M	1.77	3.01	1.94
GAPDH	1.22	1.27	1.00
HPRT1	1.69	1.05	1.10
RPLP0 GDC	0.83 1.70	0.81 1.01	1.00 1.08

	Glass Pattern	ation (comparing to 10kPa NP	10kPa Pattern
Symbol	Fold Regulation	Fold Regulation	Fold Regulation
DNMT3B	-2.08	-3.11	-2.67
GDF3	8.12	-1.68	1.18
Lefty1	22.40	2.19	3.17
Nanog	1.09	-1.37	-2.90
PODXL	-1.47	-1.59	-4.46
POU5F1	2.05	1.37	1.08
CCDC42	1.09	3.22	1.50
FGF5	-2.55	-3.95	1.19
foxD3	-1.58	-31.20	-8.74
OTX2	-1.55	-6.54	-2.62 -8.29
ZIC1 GBX2	-5.59 16.06	-12.28 4.32	9.82
NEUROG2		-2.51	1.90
BMP4	3.33	8.28	7.03
SYNE2	1.07	1.15	-1.34
EMD	-2.41	-2.19	-2.61
GATA2	7.59	37.79	21.05
HAND1	3.97	57.48	41.20
WDR5	1.89	1.66	1.64
MIXL1	89.27	25.77	49.88
PDGFRA	29.88	25.98	55.01
RUNX1	12.37	37.28	32.57
Т	29.41	28.64	42.61
FOXA1	3.25	1.27	2.46
HDAC1	1.36	1.70	1.13
GATA6	374.51	260.12	592.06
HNF4A	1.56	1.42	2.12
SOX17	69.34	146.05	339.95
LMNA	1.59	3.51	2.13
FABP7	3.30	1.58	1.57
HES5	-10.54	-5.15	-4.64
PROM1	2.05	1.54	-1.08
SOX2	-5.92	-135.61	-47.26
DCX	1.99	1.49	1.48
GAD2	1.09 -1.47	-3.22	1.56 -4.64
SLC32A1 ENO1	-1.47 1.65	-10.96 1.60	-4.64 1.48
MSLN	-4.52	-7.34	-10.04
FOXG1	-1.37	-3.23	-1.93
OLIG2	-1.07	-3.23	-1.37
NKX2-2	4.33	7.26	19.62
HAND2	8.10	17.63	11.59
CD79A	-1.05	1.21	1.01
CD3E	1.40	-1.43	-1.66
PTCRA	-1.06	1.75	-1.26
KRT19	1.40	6.17	3.20
АРОН	-1.25	1.01	1.08
DPP4	-3.43	-5.04	-3.07
MAP3K12	-2.61	-1.81	-1.82
CDH1	1.62	3.37	1.23
CDH2	5.02	2.42	3.68
VTN	-2.23	-1.53	-1.70
TLN1	1.46	1.58	1.32
WNT5A	8.03	44.49	59.28
WNT11	-1.56	4.13	2.11
ITGAV	3.47	5.24	2.27
ITGB5	3.92	4.07	3.58
PTK2	1.29	1.35	1.18
MAPK12	-1.61	-1.42	1.52
MAPK8	2.01	1.60	1.61
SMAD1 SMAD2	2.79 1.53	2.80	2.09
ISL1	1.53 3.93	1.59 36.76	1.37 12.54
NKX2-5	7.65	17.89	63.31
DKK1	78.37	35.62	72.37
DKK1 DKK3	1.11	1.95	1.01
SNAI1	4.07	6.12	12.82
SNAI2	10.19	63.26	162.86
LEFTY2	19.88	7.63	8.29
SFRP1	-27.21	-21.06	-29.40
SFRP2	2.14	-1.06	-1.13
GSC	91.66	6.90	16.44
WNT3A	3.54	18.09	28.22
GSK3B	1.80	1.76	1.19
RHOA	1.14	1.03	-1.09
YY1AP1	1.07	1.06	1.20
TAZ	-2.64	-1.81	-1.63
KLF4	-1.96	-1.80	-2.29
MAPK3	-1.28	1.53	1.50
FOXA2	160.08	48.81	246.83
SOX1	-18.13	-15.65	-14.63
CER1	1191.87	159.11	430.14
LEF1	6.28	10.09	12.12
HDAC3	1.29	1.17	1.09
ACTB	1.60	1.72	1.39
B2M	1.77	3.01	1.94
GAPDH	1.22	1.27	1.00
HPRT1	1.69	1.05	1.10
RPLPO	-1.20	-1.24	-1.00

Table 3: Antibody Information:

Pluripotency markers:

Antibody	Cat. No	Produced in	Dilution
name			used
Oct4	Goat Polyclonal Anti-Oct4 antibody (ab27985)	Goat	1:500
Nanog	Nanog Monoclonal Antibody (hNanog.2), eBioscience 14-5768-82	Mouse	1:400

Differentiation markers:

Antibody name	Cat. No	Dilution Used
Sox17	RDSAF1924 R&D Systems Human SOX17 Affinity Purified	1:300
	Goat Polyclonal Ab	
T/Brachyury	ab209665 Rabbit monoclonal [EPR18113] to Brachyury	1:500
Sox2	ab79351 Mouse monoclonal [9-9-3] to SOX2	1:500
E-Cadherin	Mouse monoclonal [M168] to E Cadherin ab76055	1:500
FoxA2	Mouse monoclonal [7E6] to FOXA2 ab60721	1:500
Sox1	SOX1 Recombinant Rabbit Monoclonal Antibody MA5-32447	1:500
Snail	SNAIL Rabbit Monoclonal Antibody (F.31.8) MA5-14801	1:400

2nd antibodies

Antibody	Cat. No	Dilution
name		Used
Anti-goat 647	ab150131 Donkey Anti-Goat IgG H&L (Alexa Fluor® 647)	1:400-
		1:600
Anti-rabbit	Donkey anti-Rabbit IgG (H+L) Highly Cross-Adsorbed	1:400-
488	Secondary Antibody, Alexa Fluor 488 A-21206	1:600
Anti-Mouse	Goat anti-Mouse IgG (H+L) Highly Cross-Adsorbed	1:400-
555	Secondary Antibody, Alexa Fluor 555 A-21424	1:600