RESEARCH ARTICLE

Interstitial fluid osmolarity modulates the action of differential tissue surface tension in progenitor cell segregation during gastrulation

S. F. Gabriel Krens1, Jim H. Veldhuis2, Vanessa Barone1, Daniel Čapek1, Jean-Léon Maître1,* G. Wayne Brodland2 and Carl-Philipp Heisenberg1,‡

ABSTRACT

The segregation of different cell types into distinct tissues is a fundamental process in metazoan development. Differences in cell adhesion and cortex tension are commonly thought to drive cell sorting by regulating tissue surface tension (TST). However, the role that differential TST plays in cell segregation within the developing embryo is as yet unclear. Here, we have analyzed the role of differential TST for germ layer progenitor cell segregation during zebrafish gastrulation. Contrary to previous observations that differential TST drives germ layer progenitor cell segregation in vitro, we show that germ layers display indistinguishable TST within the gastrulating embryo, arguing against differential TST driving germ layer progenitor cell segregation in vivo. We further show that the osmolarity of the interstitial fluid (IF) is an important factor that influences germ layer TST in vivo, and that lower osmolarity of the IF compared with standard cell culture medium can explain why germ layers display differential TST in culture but not in vivo. Finally, we show that directed migration of mesendoderm progenitors is required for germ layer progenitor cell segregation and germ layer formation.

KEY WORDS: Tissue surface tension, Cell internalization, Gastrulation, Zebrafish

INTRODUCTION

During gastrulation, the germ layer progenitor cell types—ectoderm, mesoderm and endoderm—segregate into distinct germ layers with ectoderm positioned on the outside of the embryo and mesoderm and endoderm on its inside (Stern, 2004). In zebrafish embryos, progenitor cell segregation is initiated by progenitor cells that have been induced to become mesoderm or endoderm (epiblast; Montero et al., 2005; Rohde and Heisenberg, 2007; Solnica-Krezel and Sepich, 2011; Warga and Kimmel, 1990). The molecular, cellular and biophysical mechanisms that underlie cell segregation and tissue self-organization have been studied for decades (Borghi and Nelson, 2009). Differences in cell adhesion and cortical tension, which together determine tissue surface tension (TST), are generally thought to constitute crucial determinants that drive cell sorting and tissue layering in development (Foty and Steinberg, 2013; Krens and Heisenberg, 2011). In zebrafish and *Xenopus* gastrulation, differential TST between the forming germ layers has been postulated to trigger progenitor cell segregation and germ layer positioning (Krieg et al., 2008; Maître et al., 2012; Schötz et al., 2008). However, evidence that supports this view has so far nearly exclusively come from experiments performed on cells and tissues in culture. Moreover, studies in *Xenopus* embryos have suggested that cadherin-dependent differential TST causes cell sorting in vitro, but not in the embryo (Ninomiya et al., 2012). The main difficulty in determining the contribution of differential TST to cell sorting in vivo has been the lack of techniques for determining TST within the physiological environment where these processes naturally occur.

Here, we introduce CellFIT-3D, a 3D force inference method (Brodland et al., 2010, 2014) that allows us to analyze TST within the zebrafish gastrula. Combining this tool with live cell imaging and genetic perturbation, we provide evidence that directed cell migration rather than differential TST drives progenitor cell segregation in vivo, and that osmolarity of the surrounding fluid is an important factor influencing germ layer TST.

RESULTS

To analyze the potential contribution of TST to progenitor cell segregation during gastrulation, we developed a new version of video force microscopy, CellFIT-3D, that is capable of analyzing interfacial tensions in cells from three-dimensional (3D) confocal stacks (Brodland et al., 2010, 2014). First, we validated our CellFIT-3D method by analyzing TST during cell segregation in heterotypic aggregates of ectoderm and mesoderm progenitor cells in vitro (Fig. 1A; Movie 1), previously shown to be driven by differential TST (Krieg et al., 2008; Maître et al., 2012; Schötz et al., 2008). For our analysis, we considered five different types of interfaces: two homotypic cell-cell interfaces (ectoderm-ectoderm, mesoderm-mesoderm), one heterotypic cell-cell interface (ectoderm-mesoderm) and two cell-fluid interfaces (ectoderm-medium, mesoderm-medium) (Fig. 1B). Consistent with biophysical measurements (Krieg et al., 2008; Maître et al., 2012; Schötz et al., 2008), our CellFIT-3D analysis revealed a higher ratio of cell-medium to homotypic cell-cell interfacial tensions in ectoderm compared with mesoderm cells (Fig. 1C), indicative of ectoderm displaying higher TST than mesoderm. This confirms previous findings of stronger actin and myosin II localization at cell-medium
interfacial tensions in ectoderm compared with mesoderm progenitors (Krieg et al., 2008; Maître et al., 2012; Fig. S1), and is consistent with the assumption that differential TST between ectoderm and mesoderm drives progenitor cell segregation in vitro (Schötz et al., 2008). It further supports the notion that CellFIT-3D is a reliable method with which to determine germ layer TST and analyze the specific contribution of differential TST to germ layer progenitor cell sorting.

For analyzing TST between ectoderm and mesoderm cells during cell segregation in vivo, we applied our CellFIT-3D method to confocal time-lapse movies of anterior axial mesendoderm (prechordal plate, ppl) cell internalization within the dorsal germ ring margin at the onset of gastrulation [5-6 h post fertilization (hpf)] Fig. 1E; Movie 2]. We chose to analyze ppl progenitor cells, because they are easy to identify in the developing embryo and show features common to mesendoderm progenitor cell internalization...
During gastrulation (Montero et al., 2005), we considered the ratio of progenitor cell-fluid (interstitial fluid; IF) to homotypic cell-cell interfacial tensions as a read-out for germ layer TST (Maître et al., 2012). Surprisingly, upon analyzing more than 450 manually digitized angle sets of 119 cell contacts using CellFIT-3D (Fig. 1F,F′,F″), we found that, different from the situation in culture (Krieg et al., 2008; Maître et al., 2012; Schötz et al., 2008), TST of ectoderm and mesoderm were largely indistinguishable in vivo (Fig. 1G). To further validate this observation, we also analyzed TST during internalization of ppl progenitors that were transplanted directly below the surface of the dorsal germ ring of pre-gastrula stage (40% epiboly; 5 hpf) MZoop mutant embryos lacking endogenous mesendoderm cells (Fig. 1H; Movie 3; Gritsman et al., 1999). In contrast to the situation of endogenous ppl cell internalization, where unambiguously locating heterotypic interfaces between mesoderm and ectoderm progenitors was impossible, this transplantation assay also allowed us to identify clearly and analyze these heterotypic interfaces. Similar to the endogenous situation, we found indistinguishable TST between ectoderm and mesoderm upon analysis of about 200 angle sets obtained from 60 cell contacts (Fig. 1J). Together, these analyses suggest that, unlike the situation in vitro (Krieg et al., 2008; Maître et al., 2012; Schötz et al., 2008), ectoderm and mesoderm display indistinguishable TST during mesoderm internalization at the onset of gastrulation. It further points to the possibility that while differential TST is sufficient to drive progenitor cell segregation in vivo, it might not have such a function within the embryo.

To further test this possibility, we asked to what extent the relative interfacial tension values obtained by CellFIT-3D during progenitor cell segregation in vitro versus in vivo can trigger progenitor cell segregation in silico. To this end, we performed simulations of TST-driven cell segregation using Finite Element (FE)-based forward modeling (Brodland, 2004; Brodland et al., 2007). We started our simulations with a configuration of randomly intermixed ectoderm and mesoderm cells forming a coherent cluster that is surrounded by a liquid medium, equivalent to the actual situation of progenitor cell sorting in vitro (Krieg et al., 2008; Maître et al., 2012; Schötz et al., 2008). When using the relative interfacial tension values obtained from the in vitro cell segregation experiments, ectoderm and mesoderm cells were efficiently segregating into a configuration where mesoderm surrounded ectoderm (Fig. 1D; Movie 4, left). By contrast, when the relative interfacial tension values found in vivo were used, no progenitor cell segregation was observed (Fig. 1K; Movie 4, right). These findings support our assumption that differential TST is sufficient to drive progenitor cell segregation in vitro but not in vivo.

Our analysis raises two main questions: (1) why are cell interfacial tensions different in the embryo compared to the situation in culture; and (2) what mechanism(s) – if not differential TST – drive progenitor cell segregation in vivo? In addressing the first question, we reasoned that differences between the physiological environment in vivo and cell culture conditions in vitro might be responsible. To identify those differences, we searched for factors that might vary between the situation in vivo and in vitro, and have the potential to affect cell interfacial tensions. There is increasing evidence that osmolarity of the surrounding medium plays an important role in determining hydrostatic cell pressure and, consequently, cell cortex tension, a crucial cell property that influences cell interfacial tensions (Salbreux et al., 2012; Stewart et al., 2011). We thus speculated that IF in vivo might have a different osmolarity than cell culture medium used in vitro, and that this difference might be responsible for the observed discrepancy between TST in vivo versus in vitro. To address this hypothesis, we first sought to determine osmolarity of the IF in vivo at the onset of gastrulation. To this end, we made use of a nanoliter osmometer (Otago osmometers; Braslavsky and Drori, 2013) that allows measuring the osmolarity of small fluid quantities (∼10 nL). As the total amount of IF per embryo is very small (∼15 nL) and distributed between progenitor cells throughout the gastrula (Fig. 2F,G), we were unable to extract sufficient amounts of IF directly from embryos. Instead, we made use of blastoderm explants (animal caps) excised from sphere-stage embryos (4 hpf; Krens et al., 2011), which formed a clearly recognizable accumulation of IF at the explant interior when kept in culture for ∼2-3 h (Fig. 2A). We then extracted IF from multiple of those explants by micropipette aspiration (Fig. 2B,C) and analyzed the osmolarity of the extracted IF using our nanoliter osmometer (Fig. 2D). Strikingly, we found that the osmolarity of the IF was considerably lower (250.3±47.4 mOsm/L) than the osmolarity of the cell culture medium typically used to study progenitor cell sorting in vitro (∼300 mOsm/L; Fig. 2E).

To determine whether this difference in osmolarity between IF and culture medium might be responsible for the observed differences in germ layer TST between the situations in vivo (IF) versus in vitro (cell culture medium), we used our 3D-CellFIT method to analyze TST of germ layer explants cultured in the presence of culture medium with osmolarity ranging from 126 to 300 mOsm/L. Consistent with previous observations (Krieg et al., 2008; Schötz et al., 2008), we found that in 300 mOsm/L culture medium, ectoderm explants displayed higher TST than mesoderm (Fig. 3A′), and that this differences caused mesoderm to envelop ectoderm when placing these explants adjacent to each other (Fig. 3A). By contrast, when explants were cultured in medium with an osmolarity similar or lower to that of the IF (250-126 mOsm/L; Fig. 3C-E), mesoderm and ectoderm explants displayed indistinguishable TST (Fig. 3C′-E′) and, consequently, there was no envelopment observed when these tissues were brought into contact with each other (Fig. 3C-E). Interestingly, ectoderm and mesoderm tissues still displayed differential TST and, consequently, mesoderm enveloped ectoderm in the presence of culture medium with an osmolarity intermediate between the osmolarity of standard culture medium and IF (275 mOsm/L; Fig. 3B′,B′). This suggests that medium/IF osmolarity must be at least as low as 250 mOsm/L for differential TST between ectoderm and mesoderm to vanish. This conclusion was further supported by simulations of explant envelopment with the interfacial tension values obtained from our 3D-CellFIT analyses using FE-based forward modeling, producing envelopment behaviors similar to the ones observed in the experiments (Fig. 3F,G; Movies 5 and 6), confirming that osmolarity-induced changes in explant envelopment were indeed due to associated changes in explant TST. Additionally, to test the validity of our 3D-CellFIT based findings on interfacial tension values in the presence of culture medium with different osmolarity, we directly measured cell-medium interfacial tensions (cortical tensions) of individual ectoderm and mesoderm progenitors in the presence of high (300 mOsm/L) versus low (190 mOsm/L) osmolarity culture medium using single cell force spectroscopy (Krieg et al., 2008). Consistent with our 3D-CellFIT data, cell-medium interfacial tension was higher in ectoderm compared with mesoderm progenitors in the presence of culture medium with 300 mOsm/L, whereas no such difference was detectable anymore when medium osmolarity was lowered to 190 mOsm/L (Fig. S1). Collectively, these findings support our initial assumption that osmolarity affects TST, and that differences in the osmolarity...
between IF and cell culture medium can explain the discrepancy in the measured TST in vivo versus in vitro.

Finally, we asked whether the observed effect of culture medium osmolarity on germ layer explant TST and, consequently, their envelopment behavior in culture, was indeed mediated by changes in medium osmolarity rather than alterations in the concentration of specific culture medium ingredients. To this end, we tested whether increasing culture medium osmolarity from 250 to 300 mOsm/L by adding the non-ionic osmolyte mannitol, a sugar frequently used to manipulate culture medium osmolarity (Enyedi et al., 2013), would have the same effect on explant envelopment behavior as observed when placing these explants directly into 300 mOsm/L culture medium. We found that the addition of mannitol to the culture medium induced germ layer explant envelopment to a similar degree to that observed when placing heterotypical explants directly into 300 mOsm/L culture medium. Our findings suggest that changes in medium osmolarity, rather than the concentration of specific culture medium ingredients, affect germ layer TST.

Our findings so far suggest that germ layer progenitor cells do not display differential TST in vivo, and, consequently, that differential TST is unlikely to drive progenitor cell segregation during gastrulation. To investigate which mechanism(s) – if not differential TST – then drive progenitor cell segregation within the embryo, we performed multi-photon time-lapse imaging of endogenous ppl progenitors internalizing at the dorsal germ ring at the onset of gastrulation (5-6 hpf; Fig. 4A-A″,E; Movie 7). We found that individual ppl progenitor cells marked by their expression of GFP in Tg(gsc:eGFP-Hsa.HRAS) embryos moved from the outside to the inside of the germ ring margin, a behavior characteristic of progenitor cell ingestion (Montero et al., 2005). Moreover, internalizing ppl progenitor cells, but not non-internalizing ectoderm cells, displayed dynamic actin cortical patterns typically associated with migrating cells, such as preferentially localizing actin to their protrusive front-end (Fig. 4G,G″; Fig. S2,3). Myosin II localization, by contrast, did not show any preferential localization to the leading or trailing edges of internalizing ppl cells (Fig. 4G″), and exposing cultured ppl cells to the myosin II inhibitor blebbistatin did not interfere with protrusion formation in these cells (Fig. S3). Together, this points to the possibility that ppl progenitors segregate from non-internalizing ectoderm progenitors by undergoing directed cell migration. To further test this possibility, we asked whether interfering with the migratory capacity of ppl progenitors would disrupt progenitor cell ingestion and thus segregation within the germ ring. To interfere with ppl cell migration, we expressed a dominant-negative version of Rac (DN-Rac), previously shown to reduce cell protrusion formation and migration (Hall, 1998; Ridley et al., 1992), either uniformly within the gastrulating embryo (Fig. 4B″,F) or specifically within transplanted ppl progenitor cells (Fig. 4H″,K). Strikingly, we found that in both of these cases, ppl progenitor cells failed to undergo internalization (Fig. 4B-D,F; Movie 8). Moreover, ppl progenitors overexpressing DN-Rac did not show any preferential localization of actin to their front ends in vivo (Fig. 4H″,J), and displayed reduced protrusion formation in vitro when cultured on fibronectin-coated substrates (Figs S2 and 3; Movie 9). Importantly, overexpression of DN-Rac did not affect differential TST-driven envelopment of ectoderm by mesoderm tissue in culture (Fig. S4), suggesting that DN-Rac does not strongly interfere with the differential TST that these tissues display. Collectively, these findings suggest that directed ppl progenitor cell migration plays a crucial role in progenitor cell internalization and segregation during gastrulation.

DISCUSSION

Previous studies have shown that germ layer progenitor cell segregation in culture is driven by differences in TST among the forming germ layers, with ectoderm displaying higher TST than mesoderm and endoderm (Krieg et al., 2008; Maitre et al., 2012; Schötz et al., 2008). Here, we show that this difference in germ layer TST crucially depends on the osmolarity of the surrounding fluid interface, and that within the gastrulating embryo under physiological osmolarity levels, this difference in TST diminishes. This argues against TST playing an instructive role in germ layer progenitor cell segregation during zebrafish gastrulation.

Osmolarity has previously been shown to affect cell interfacial tensions by altering hydrostatic cell pressure that in turn is balanced by cortex tension (Lang et al., 1998; Salbreux et al., 2012; Stewart,...

---

**Fig. 2. Measurements of interstitial fluid osmolarity.** (A) Schematic illustration of animal cap explant preparation, and IF labelling and extraction procedures. (B,C) Overlaid images of an animal cap explant (gray scale) with IF-filled lumen (blue) and an extracted IF droplet (blue) at the tip of a glass capillary used for IF extraction (C). Scale bar: 200 μm. (D) Images of a droplet of IF floating in immersion oil placed within the sample-holder hole of a nanoliter osmometer in a frozen state (left) and directly after melting (right). The displayed formula was used to calculate the IF osmolarity, with O$_s$ denoting osmolarity (Osm/L) and T$_m$ being the temperature of freeze point depression. Scale bar: 50 μm. (E) Box and whisker plot of the measured IF osmolarity (n=8 measurements). Black line indicates the median value; whiskers show the spread of the data. (F,G) 3D renderings of a lateral fluorescence image of the dorsal germ ring in a Tg(-4gsc:eGFP-Hsa.HRAS) embryo at the onset of gastrulation (6 hpf), expressing GFP in internalizing ppl progenitors (green), utrophin-mCherry to label the actin cell cortex of all cells (red) and dextran-Cascade Blue to label the IF (blue; F). (G) The same image showing dextran-Cascade Blue labeling of IF only. Scale bar: 20 μm.
et al., 2011). So far, studies on the interplay between medium osmolarity and hydrostatic cell pressure have mostly focused on cell responses to changes in medium osmolarity on timescales of seconds to minutes. By contrast, progenitor cell segregation both in vitro and in vivo occurs over a period of minutes to hours, and thus we recorded the response of progenitor cells to changes in medium/IF osmolarity on comparably long timescales. Consequently, the response of ectoderm and mesendoderm progenitors to changes in medium/IF osmolarity in our analysis describes the specific ability of those cell types in maintaining fluid homeostasis rather than their immediate response to changes in hydrostatic pressure. How, over such comparably long timescales, medium and/or IF osmolarity affects progenitor cell interfacial tensions is not yet clear, but the ability of progenitor cells to undergo regulated volume increase or decrease in response to osmotic swelling or shrinkage, and associated changes in the ionic composition of the cell cytoplasm are likely involved. A systematic analysis of how medium and/or IF osmolarity affects progenitor cell interfacial tensions, and how the

Fig. 3. Modulation of progenitor cell interfacial tensions by medium osmolarity. (A–E) Representative single plane confocal images of tissue aggregates consisting of ectoderm or mesoderm progenitor cells expressing Lyn-Venus at the plasma membrane (green) of all cells and cultured for 5 h in the presence of medium with 300 (A), 275 (B), 250 (C), 190 (D) or 126 (E) mOsm/L osmolarity. Ectoderm aggregates were additionally labeled with cytoplasmic dextran-Alexa648 (red) (see also Movie 3). Scale bar: 50 μm. (A′–E′) Relative interfacial tensions ($\gamma_{rel}$) obtained by 3D-CellFIT for enveloping tissues cultured for 5 h in the presence of medium with 300 (A′), 275 (B′), 250 (C′), 190 (D′) or 250 (E′) mOsm/L osmolarity with e (ectoderm), m (mesoderm) and cm (culture medium). Error bars indicate standard deviations. (F,G) Stable configurations of finite element simulations of tissue envelopment (10,000 simulation iterations) in heterotypic tissue aggregates consisting of ectoderm (red) or mesoderm (green) cells surrounded by culture medium (blue), using 3D-CellFIT-obtained tension distributions shown in A (300 mOsm/L) with $\gamma_{e-e}=1.00$, $\gamma_{m-m}=1.62$, $\gamma_{e-m}=2.41$, $\gamma_{e-cm}=2.94$, $\gamma_{m-cm}=1.58$; and in C (250 mOsm/L) with $\gamma_{e-e}=1.00$, $\gamma_{m-m}=1.16$, $\gamma_{e-m}=1.32$, $\gamma_{e-cm}=1.52$, $\gamma_{m-cm}=1.41$. (H–J) Tissue envelopment of ectoderm (red) and mesoderm (green) progenitor cell aggregates cultured for 5 h in the presence of ~300 mOsm/L culture medium ($n=20$ engulfment assays; H), ~250 mOsm/L osmolarity culture medium ($n=24$ engulfment assays; I) or culture medium containing mannitol to restore osmolarity from 250 mOsm/L to 300 mOsm/L (250+50 M; $n=21$ engulfment assays, J). (K) Degree of envelopment was quantified by calculating the relative ectoderm surface occupation taking the heterotypical cell aggregate size into account. Error bars are standard deviations; *$P<0.05$. 

1802
acquisition of different cell fates by those progenitor cells modulates their response to IF and/or medium osmolarity will be needed to further explore how osmolarity functions in gastrulation movements.

We also show that instead of differential TST driving germ layer progenitor cell segregation, directed migration of mesendoderm cells from the outside to the inside of the germ ring margin is required for mesendoderm cell internalization during gastrulation. Why mesoderm cells polarize and migrate from the outside to the inside of the germ ring is still unclear, but one possibility is that the blastoderm displays an overall polarity along the radial axis of the embryo, and that this tissue polarity then triggers mesoderm polarization and internalization. Supporting this assumption are previous findings that progenitor cells show a preferential localization of their microtubule organizing centers (MTOCs) into the dorsal germ ring margin of a wild-type host embryo at shield stage (6 hpf). Host embryos were injected with dextran-rhodamine to label the IF (blue). Scale bar: 20 μm.

Fig. 4. Mesoderm cell internalization relies on directed mesoderm cell migration. (A,A′,B,B′) Bright-field (A,B) and fluorescence images (A′,B′) of wild-type (A,A′) and DN-Rac-expressing (B,B′) Tg(−4gsc:eGFP-Hsa.HRAS) embryos, expressing GFP in internalizing ppl progenitors (green) at the onset of gastrulation (5.5 hpf). Scale bars: 100 μm. (A′,B′) Orthogonal view of a confocal image stack of a wild-type (A′) and DN-Rac-expressing (B′) Tg(−4gsc:eGFP-Hsa.HRAS); Tg[Actin:Utrophin-mCherry] embryo at the onset of gastrulation (6 hpf), expressing GFP (green) in internalizing ppl progenitors and Uetrophin-mCherry (red) marking actin in all cells. Transgenic embryos were also injected with dextran-rhodamine to label the IF (blue). Dashed line delineates the position of the EVL and dotted lines demarcate the deep cell mass. While arrows indicate the direction of mesoderm progenitor cell movement. Scale bar: 20 μm. (C) The fraction of DN-Rac-expressing and wild-type control embryos displaying defective mesoderm cell internalization (nWT=124 and nDN-Rac=96 embryos from three independent experiments). Error bars are standard deviations. (D) Box and whisker plot showing the spread of internalized mesoderm progenitors from the EVL in wild-type and DN-Rac-expressing embryos (nWT and nDN-Rac=264 cells from three embryos each; ***P<0.01). Black line indicates the median value; whiskers show the spread of the data. (E,F) Orthogonal views from a confocal image stack of internalizing ppl progenitors within the dorsal germ ring margin of wild-type (E) and DN-Rac-expressing (F) Tg(−4gsc:eGFP-Hsa.HRAS) embryo, expressing GFP in ppl progenitors (green) at shield stage (6.5 hpf). Transgenic embryos were also injected with Lym-TagBFP mRNA to outline the membrane of all cells (red) and dextran-rhodamine to label the IF (blue). Cell tracks delineate the movements of exemplary mesoderm (green spheres) and ectoderm cells (red spheres) during internalization (5.7-6.5 hpf). Track speed calibration bar indicates movements of exemplary mesoderm (green spheres) and ectoderm cells (red spheres) during internalization (5.7-6.5 hpf). Track speed calibration bar indicates...
result of this polarized IF distribution, mesendoderm progenitors might preferentially be in contact with IF closer to the germ ring outside, where more of it can be found, and this polarized IF interface might in turn trigger radial mesendoderm polarization. To test this assumption, techniques need to be developed that would allow direct analysis and manipulation of IF distribution within the developing embryo.

The role of differential TST in early development is still debated. Our CellFIT-3D-based analysis of cell interfacial tensions within the gastrulating embryo provides the first direct evidence that differential TST is not sufficient to explain germ layer progenitor cell segregation during zebrafish gastrulation. This does not argue against differential TST playing other important roles in early development, but clearly shows that complex morphogenetic processes, such as the formation and positioning of the different germ layers during gastrulation, depend on the interplay between different processes, including directed cell migration and polarization.

**MATERIALS AND METHODS**

**Zebrafish handling**

Zebrafish maintenance was carried out as described previously (Westerfield, 1993). Embryos were grown at 28-31°C in Danieau’s embryo medium and staged as described previously (Kimbel et al., 1995). The following wild-type (WT), mutant and transgenic lines were used: (WT) TL; (mutant) maternal zygotic (MZ) oep (Gritsman et al., 1999); (transgenic) Tg(dharma: eGFP) (Ryu et al., 2001), Tg(-4gsc:eGFP-Hsa.HRAS), Tg(bact:Ras-eGFP) (Cooper et al., 2005), Tg(bact:myl12.1-eGFP) and Tg(bact:myl12.1-mCherry) (Maitre et al., 2012); and Tg(bact:Lif6cAct-eGFP) and Tg(actb1:mCherry;utrCH) (Behnert et al., 2012).

**Embryo microinjections**

Embryo microinjections consisting of one germ layer progenitor cell type only were obtained by injection of one-cell stage embryos with either 100 pg lefty1 mRNA (ectoderm) or 100 pg ndr2/cyclops mRNA plus 2 ng casanova morpholino (case MO; mesoderm) (Krieg et al., 2008). To visualize the plasma membrane and filamentous actin, 100 pg lyn-TagBFp or lyn-Venus (plasma membrane), and 50-100 pg LifeAct-eGFP (F-actin; Behnert et al., 2012) were injected at the one-cell stage. To inhibit cell protrusion formation, one-cell stage embryos were injected with 400 pg DN-Rac mRNA. To distinguish between ectoderm and mesoderm populations in cell sorting and tissue envelopment experiments, embryos were additionally injected with Dextran-FITC or Dextran-tetra-methyl-rhodamin-dextran (TMR-dextran, LifeTechnologies) at the one-cell stage to label the cell cytoplasm. To visualize interstitial fluid (IF), 0.5-1.0 NL of 0.1% (w/v) TMR-dextran or Dextran-Cascade Blue (LifeTechnologies) were injected between deep cells of sphere-stage embryos (4 hpf) or tissue explants.

**Cell transplantsations**

Donor and host embryos were dechorionated with forceps and transferred into an agarose plate containing Danieau’s embryo medium. For the exogenous mesoderm internalization assay, two to five cells were taken from a mesoderm-induced donor embryo using a bevelled borosilicate needle with a 20 μm inner diameter attached to a syringe system, and transplanted directly below the surface cells close to the dorsal blastoderm margin of a dome stage (4.5 hpf) host embryo. For clonal analysis of actin and myosin II substrates for 60-90 min prior to imaging. Imaging was performed on an inverted microscope (Axio Observer Z1 Zeiss) equipped with an automated TIRF/Epi-fluorescence system (Visitron Systems), with 488 nm and 561 nm laser lines. Images were acquired using a 20× objective (Zeiss) and an EMCCD camera (Evolve, Photometrix) with frame rates of 2 min and exposure times of 20-500 ms. Protrusion analysis was performed by segmenting out the seeded cells, from the images and comparing the perimeter length of the segmented area to the perimeter of an ellipsoid that was fitted to have the same area, and the same longest axes. An ellipsoid was chosen to compensate for unequal spreading or tissue stretching.

**IF osmolarity measurement**

Donor and host embryos were dechorionated with forceps and transferred into an agarose plate containing Danieau’s embryo medium. Animal poles were cut from the embryo at high-to-sphere stage (3-4 hpf), left to round up for 0.5 h at room temperature (22-25°C), and then cultured in Danieaus embryo medium at 28.5°C (Krens et al., 2011). After 2-4 h, when a clear fluid-filled cavity was formed at the explant interior, the IF was extracted and pooled from two to four explants per measurement using a bevelled borosilicate needle with a 10 μm inner diameter attached to a syringe system. For osmolarity rescue experiments, diluted medium osmolarity. For osmolarity rescue experiments, diluted medium was supplemented with mannitol (Sigma-Aldrich).

**In vitro cell protrusion assay**

Glass-bottom dishes (MatTek) were coated with fibronectin by adding 50 μL of 200 μg/ml bovine fibronectin (Sigma), air dried at room temperature and overlaid with 50 mg/ml BSA (Invitrogen) for 10 min. Mesoderm (prechordal plate) progenitor cells were isolated by first removing the animal poles of mesoderm-induced embryos (see above), and cutting them into smaller pieces with a watchmaker forceps. Cells-clusters were seeded at 1×DMEM/F12 medium (Invitrogen) and left to adhere on fibronectin-coated glass substrates for 60-90 min prior to imaging. Imaging was performed on an inverted microscope (Axio Observer Z1 Zeiss) equipped with an automated TIRF/Epi-fluorescence system (Visitron Systems), with 488 nm and 561 nm laser lines. Images were acquired using a 20× objective (Zeiss) and an EMCCD camera (Evolve, Photometrix) with frame rates of 2 min and exposure times of 20-500 ms. Protrusion analysis was performed by segmenting out the seeded cells, from the images and comparing the perimeter length of the segmented area to the perimeter of an ellipsoid that was fitted to have the same area, and the same longest axes. An ellipsoid was chosen to compensate for unequal spreading or tissue stretching.

**Embryo imaging**

Embryos were mounted in 0.7% low melting point agarose in Danieaus embryo medium. To record high-resolution time-lapse movies of cells and tissues deep within the embryo, a TriM Scope multi-photon microscope (LaVision BioTec) was used equipped with a multi-photon laser (Chameleon from Coherent) set to 830 nm and an OPO laser set to 1100 nm for exciting mCherry-labeled proteins. Image stacks of 70-150 μm were made from a polydimethylsiloxane (PDMS) negative obtained from www.microtissues.com according to the supplier’s guidelines. The different germ layer progenitor cell types were isolated and mixed by first removing the embryo animal poles of both mesoderm- and ectoderm-induced embryos (see above). For tissue envelopment assays, the animal poles of sphere-stage embryos (4 hpf) were cut into four equally sized pieces that were left to round up for 1 h at room temperature (22-25°C). One animal pole tissue piece from a MZoep mutant embryo (ectoderm) was then co-cultured with a similarly sized piece from a mesoderm-induced embryo in a micro-well, and their envelopment behavior was recorded for at least 5 h at 28.5°C. For cell-sorting experiments, the same number of animal poles from ectoderm- and mesoderm-induced embryos were pooled in a tube and dissociated by gently tapping the tube. The resulting heterotypic ectoderm-mesoderm cell mixture was then seeded on micro-wells, and their sorting was recorded for at least 5 h in 3D over time at 28.5°C by acquisition of 4 μm spaced z stacks of the aggregate in two or three channels every 5 min, using a Leica SPS confocal microscope equipped with a Leica 25×0.95NA dipping lens. All experiments were performed in CO2-independent DMEM/F12 medium, or water dilutions of it to lower medium osmolality. For osmolarity rescue experiments, diluted medium was supplemented with mannitol (Sigma-Aldrich).
Image analysis
Quantification of signal intensity and cell size was performed by analyzing images with Fiji (Schindelin et al., 2012). Intensity plots were generated by normalization to the average fluorescence intensity value and by scaling the front to the rear perimeter length. At least eight independent measurements were averaged and their mean values were displayed as polar plots with standard deviation. The ratios between the cell-front and cell-rear intensities were calculated using ImageJ (National Institutes of Health). Quantification of relative fluid occupation was plotted as a percentage of fluid occupation over the total distance from the epiblast surface by normalizing to the 256 gray values of the 8-bit image. The obtained values were averaged over 10 μm bins from the epiblast surface (0 μm) to its inside (50 μm). Cell tracking, orthogonal views and 3D-renderings were generated using Imaris version 7.3 (Bitplane). The distance to the enveloping layer (EVL) was calculated from the 3D coordinates of manually generated cell tracks for mesoderm, ectoderm and EVL cells. Statistical data analysis was performed using the GraphPad Prism 5 software.

Single cell force spectroscopy
Cell cortex tension measurements were performed as described previously (Krieg et al., 2008) with the following adjustments: the animal poles were transferred to 1×DMEM/F12 (240 mOsm/L). Five animal poles each were then transferred to either 1×DMEM/F12 (300 mOsm/L) or 0.63×DMEM/F12 (190 mOsm/L). The animal poles were mechanically dissociated and individual cells were seeded on a glass substrate. Cells were probed with colloidal force probes, which were prepared by attaching a glass bead (5 μm diameter, Kisker Biotech) to a cantilever (Vecco MLCT). To prevent non-specific adhesion of the cantilever bead to the cells, the modified cantilevers were incubated with heat-inactivated fetal calf serum for at least 1 h at room temperature (FCS, Invitrogen) prior to the measurements. Force-distance curves were acquired using 500 pN contact force and 1 μm s⁻¹ approach/retract velocity and indentation (8) was calculated from the tip displacement. Up to three curves with at least 10 s waiting time between successive curves were taken per cell to prevent any history effect. The liquid droplet model was applied to extract the cell-cortex tension as described previously (Krieg et al., 2008), with the following adjustments: to determine cell-cortex tension we used a force versus indentation line-fit between a 200 nm and 300 nm indentation range.

CellFIT-3D analysis
To obtain estimates of the relative edge tensions, the angles at triple junctions, such as those between a mesoderm cell, ectoderm cell and the medium, were digitized using custom software as described in the supplementary Materials and Methods. Angles along particular edges were digitized in multiple images within the stack in order to obtain the true angles of the cell membranes with the edge. Force-balance equations were written for each digitized triple junction, and least-squares solutions were found for all such equations. The solutions to these equations provided the relative strengths of the tensions along each edge type.

Finite element simulations
The simulations were carried out using the finite element formulation described previously (Brodland and Chen, 2000), which assumes that cell-cell and cell-medium interfaces carry net interface-specific tensions and the cytoplasm and other contents of the cells generate an effective viscosity that can be described by an orthogonal system of dashpots. In cases where cortical tensions varied within a particular cell, the tension applicable to any particular edge was based on the location of its midpoint. All calculations were carried out using an updated Lagrangian approach and the simulations were run until motion stopped. For all simulations, effective viscosity in ectoderm and mesoderm cells was assumed to be the same, and variation of this parameter did not change the principal outcome of cell segregation.

Acknowledgements
We thank D. Wagner for providing the pky zebrafish line, D. Gilmour, E. Raz and T. Kudoh for providing plasmids, and E. Paluch, D. Siekhaus and M. Sixt for reading earlier versions of the manuscript. We are grateful to all members of the Heisenberg and Sixt groups for fruitful discussions, to M. Sikora, J. Guzman and S. Pallarés for help with the analysis of the AFM data, measuring trans-epithelial resistance and measuring IF osmolarity, respectively, and the zebrafish, light microscopy and mechanical workshop facilities at IST Austria and MPI-CBG for assistance.

Competing interests
The authors declare no competing or financial interests.

Author contributions

Funding
This research received funds from the Natural Sciences and Engineering Research Council of Canada (NSERC) Discovery Grant (RGPIN-36514 to G.W.B.) and from the Austrian Science Foundation (FWF) (P305-B20 to C.-P.H.). Deposited in PMC for immediate release.

Supplementary information
Supplementary information available online at http://dev.biologists.org/lookup/doi/10.1242/dev.144964

References


**Supplementary Information**

**Supplementary Methods - CellFIT-3D**

CellFIT provides a general-purpose mathematical formalism for calculating the forces that produce specific observed cell and tissue motions and geometries (Brodland et al., 2010; Brodland et al., 2014). In the past it was applied to 2D scenarios, but here for the first time, it is applied to a 3D system. This dimensional expansion was possible in part because the cell deformation rates in the present system were sufficiently low that the complex viscous calculations associated with deformation of the cytoplasm could be avoided, and because the Young angles are measured right at the triple junctions, so that they are independent of pressure effects.

In general, confocal images are not normal to the triple junctions visible in them, and so that accurate Young angles could be obtained, a software tool was built. The user identifies any one particular triple junction in successive images with the tool and graphically identifies the in-plane angles of the membranes visible at those junctions. The software then fits a spline through the three to five manifestations of a particular junction and based on it calculates normal planes through the junction points in each image. The digitized vectors $v_1$, $v_2$ and $v_3$ associated with each triple junction are then projected onto their respective normal planes to become $V_1$, $V_2$ and $V_3$ and standard equilibrium equations

$$\sum_{k=1}^{3} V_k \gamma_{i-j} = 0$$

where $\gamma_{i-j}$ is the interfacial tension along vector $V_k$ were constructed. In this case, standard indicial conventions are not followed but rather the hyphenated subscripts on the $\gamma$ indicate the cell or liquid components that form that interface and the subscript on the vector indicates its arbitrary sequence number. The equations are constructed and solved on their respective rotated planes.

Solution produces a set of two or three ratios. For example, at a triple junction with three cell types (A, B and C) or two heterotypic cells and interstitial fluid
(IF), this process returns the ratio of the forces between those three edges, e.g., $\gamma_A:\gamma_B:\gamma_C$ or $\gamma_A:\gamma_B:\gamma_{IF}$. When two edges at a heterotypic triple junction (say A-A-B) involve the same edge combination (A-B in this case), a single average ratio $(\gamma_A, \gamma_B)$ is returned. Typically, the ratios from each imaging plane along a single triple junction are quite similar to each other, but data from the ends of each triple junction tend to be less consistent with the others and any outliers are manually deleted before an average ratio for each triple junction is calculated.

These ratios represent equilibrium equations for each node, and they can be collected together into a single matrix $J$ where each row corresponds to the ratios obtained from a single triple junction and their column placements correspond to their interface type. In a system that has ratio information from four different triplets, $J$ might look like this:

<table>
<thead>
<tr>
<th></th>
<th>$\gamma_{e-e}$</th>
<th>$\gamma_{m-m}$</th>
<th>$\gamma_{e-m}$</th>
<th>$\gamma_{e-if}$</th>
<th>$\gamma_{m-if}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Triplet 1</td>
<td>2</td>
<td>6</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triplet 2</td>
<td>1</td>
<td>1.5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triplet 3</td>
<td>1</td>
<td>0.5</td>
<td>0.25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triplet 4</td>
<td>1</td>
<td></td>
<td>1.5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The first row would arise from a triple junction equilibrium equation whose solution gives the ratio $\gamma_{ee}:\gamma_{em} = 2:6$. Blanks in the tabular representation of the $J$ matrix and the other matrices described here correspond to null entries.

The next step in CellFIT-3D is to find interfacial tensions $\gamma_i$ that best satisfy these potentially incompatible equations. Since the equations in the present analysis involve non-negative ratios exclusively, the approach of Kwiesielewicz (1996) is used to solve them (Kwiesielewicz, 1996). In this approach a system of simultaneous linear equations is constructed and solved. The example $J$ matrix from above would produce the following so-called judgment matrix $R$:
Details of the matrix construction process are discussed in Kwiesielewicz, and in terms of the language used there, the ratios from each triple junction (each row of J) correspond to a separate decision maker (Kwiesielewicz, 1996).

Based entirely on the structure and numerical values of R, a set of equations is then constructed:

\[ A \mathbf{x} = \mathbf{b} \]
Although details of this process are given in Kwiesielewicz (Kwiesielewicz, 1996), we note that A is a symmetric tally matrix whose off-diagonal elements are equal to minus the number of non-zero elements in each of the corresponding vertical rectangles of R. The diagonal elements are equal to minus the sum of the off-diagonal elements in that row (or column) of A. The elements of the b matrix are equal to the natural logarithms of the product of all of the numbers in the rectangles in the corresponding row (the elements in the shaded diagonal rectangles can be ignored as they are all ones, and they make no contribution to the product).

In general, the matrix A will not be directly invertible and its generalized inverse must be used in the solution process. Solution of the sample system gives:

\[
\mathbf{x} = \begin{bmatrix}
0.382 \\
-0.31 \\
-0.72 \\
-0.02 \\
0.67
\end{bmatrix}
\]

and the following gammas:

\[
\gamma = \begin{bmatrix}
0.683 \\
1.365 \\
2.048 \\
1.024 \\
0.512
\end{bmatrix}
\]
where $\gamma_m = e^{(x_m)}$.

The result of this solution procedure is a set of ratios $\gamma_{ee} : \gamma_{mm} : \gamma_{em} : \gamma_{ei} : \gamma_{mi}$ (in this case, 0.68: 1.37: 2.05: 1.02: 0.51) that best satisfy the observed triple junction angles in a (logarithmic) least-squares sense.

To estimate the standard deviations of the interfacial tensions, the individual rows of $J$ were scaled so as to minimize the mean percentage error between the individual entries in that row and the least-squares solution. Standard deviations for individual $\gamma_{ij}$ values were then calculated as the standard deviation of the numbers in each of the columns of the newly scaled $J$ matrix.
Supplementary Figures

Fig. S1 - Medium osmolarity modulates progenitor cell cortical tensions.

(A) Schematic illustration of a Single Cell force Spectroscopy (SCFC) experiment. A passivated colloidal force probe (bead; diameter = 5 μm) is moved towards a germ layer progenitor cell at 1 μm s⁻¹ (i) and the cell surface is deformed by the bead (ii), while the indentation force is being recorded. (B) Box-whisker plot of cell-cortex tension obtained by SCFC. The median is depicted as a line and mean as a cross; asterisks above brackets indicate p-values for the indicated combination with * = 0.05, ** = 0.001, *** < 0.0001.
Fig. S2 – Subcellular localization of actin and myosin II in ectoderm and mesoderm progenitors *in vitro* and *in vivo*.

(A-D) Single confocal image plane of an exemplary Tg(βActin:myl12l-eGFP);Tg(βActin:Utrophin-mCherry) mesoderm (ppl) (A,C) and ectoderm (B,D) progenitor cell *in vivo* (A,B) with the surrounding IF labelled by Dextran-Cascade Blue® (blue), and *in vitro* (C,D) with adjacent cells labelled with Dextran-Alexa647 (white). Scale bars, 10 µm. (A’-D’) Schematic illustration of the cells shown in (A-D) with their leading (front) and trailing edges (back) and cell-medium (out) and cell-cell (in) interfaces indicated. (A’’-D’’) Normalized fluorescence intensity values on the periphery of mesoderm (ppl) (A’’,A’’,C’’,C’’) and ectoderm (B’’,B’’,D’’,D’’) cells from Tg(βActin:myl12l-eGFP);Tg(βActin:Utrophin-mCherry) embryos either *in vitro* (A’’,A’’,B’’,B’’) or within embryos at shield stage *in vivo* (6 hpf; C’’,C’’,D’’,D’’) shown as polar plots for actin (Utrophin-mCherry; A’’-D’’) and myosin II (myl12l-eGFP; A’’-D’’) with the front and rear areas (*in vivo*) and ‘out’ and ‘in’ interfaces (*in vitro*) annotated. Standard deviation is indicated as perpendicular lines to the mean. n (ppl *in vivo*) = 21 cells (5 embryos), n (ectoderm *in vivo*) = 8 cells (5 embryos), n (mesoderm *in vitro*) = 18 cells (3 cell aggregates), n (ectoderm *in vitro*) =18 cells (3 cell aggregates). (E,F) Box-whisker plot of intensity ratios of actin (red) and myosin II (green), from the intensity profiles shown in (A’’-D’’, A’’-D’’), of the front versus the rear of ppl and ectoderm cells *in vivo* (E), and medium-facing outside and aggregate-facing inside of induced mesoderm and ectoderm progenitor cells.
on the surface of a heterotypical cell aggregate during cell sorting in vitro (F). Black line indicates the median value; whiskers show the spread of the data; dashed line indicates a value of 1 (equal intensity front versus rear); * p < 0.05.

Fig. S3 - Protrusion formation in mesoderm (ppl) and ectoderm progenitor cells in vitro.

(A,A’,B,B’) Bright-field (A,B) and fluorescence images (A’,B’) of WT (A,A’) and DN-Rac expressing (B,B’) Tg(βActin:Utrophin-mCherry) ppl progenitor cells plated on fibronectin-coated substrates for 1.5 h to monitor their protrusion formation in vitro. Scale bars, 10 μm. (C) Bar graph showing the percentage of protrusion forming ppl cells at the cell aggregate periphery. n (WT) = 14 and n (DN-Rac) = 15 experiments. Error bars are standard deviations, *** p < 0.01.

(D-I) Bright-field images of WT ppl (D,E,H,I), DN-Rac expressing ppl (F) and ectoderm (G) progenitor cell clusters plated on fibronectin-coated substrates and cultured in 300 mOsm/L (D,F,G,H,I) or 250 mOsm/L (E), supplemented with either Blebbistatin - (H) or Blebbistatin + (I) to monitor their protrusion formation in vitro for 2 h. Scale bars, 20 μm. (J) Line graph showing the quantification of protrusive activity by plotting the calculated difference
between the plated cell cluster perimeter (red lines in D-I) of the segmented cells clusters (blue area in D-I) and a fitted ellipsoid with the same area (red lines in D-I) over time. n (ppl, 300 mOsm/L) = 8, (ppl, 250 mOsm/L) = 4 , n (ppl + DN-Rac) = 8 , n (ectoderm) = 4 experiments. (K) Line graph showing the same quantification approach as in (J) for a separate experimental data set of ppl progenitor cell cluster cultured in 300 mOsm/L supplemented with blebbistatin isoforms, n (ppl) = 7 , n (ppl + Bleb-) = 6 , n (ppl + Bleb+) = 6 experiments.

Fig. S4 – Envelopment of ectoderm and mesoderm cell aggregates overexpressing DN-rac in vitro.

(A) Representative single plane confocal images of tissue aggregates consisting of ectoderm or mesoderm progenitor cells cultured for 5 h in the presence of medium with 300 (A) and 250 mOsm/L osmolarity. Both ectoderm and mesoderm cells express DN-Rac, and Lyn-Venus at the plasma membrane (green). Ectoderm aggregates were additionally labeled with cytoplasmic Dextran-Alexa648 (red). Scale bar, 50 μm. (B) Degree of engulfment was quantified by calculating the relative ectoderm surface occupation taking the heterotypical cell aggregate size into account. Error bars are standard deviations.
Fig. S5 – Interstitial fluid accumulation within the blastoderm.

(A) Schematic illustration of the imaged region within the embryo used for interstitial fluid (IF) quantification. (B,C) Binary converted single plane confocal images of the epiblast with its surface pointing to the right and its IF labelled with Dextran-Rhodamin in embryos at 30 % (4.5 hpf, B) and 60 % (6.5 hpf, C) epiboly stages. Dashed lines (cyan) indicate 5 x 10 µm bins used for IF quantification. Light-to-dark colouring indicates the distance of the binned region from the EVL in µm. Scale bars, 20 µm. (D) Bar-graph presenting the average IF-to-cell space filling ratios per 10 µm bin from the epiblast surface (0 µm, light blue) to its inside (50 µm, dark blue) in embryos from sphere (4 hpf) to shield stage (6 hpf). Error bars, standard deviation within the analysed bin.
Supplementary Movies

Movie 1 - Cell sorting in heterotypic cultures of ectoderm and mesoderm progenitors.
Confocal image series of cell sorting in heterotypic aggregates consisting of ectoderm and mesoderm progenitor cells cultured for 4.5 h. All cells express Lyn-VenusGFP to outline the plasma membrane (green). Ectoderm progenitor cells were additionally labelled with cytoplasmic Dextran-Alexa648 (red). Frame rate, 6 fps; ~ 3min/frame. Scale bar, 50 μm.
**Movie 2 - Endogenous mesoderm cell internalization.**

Confocal image series of endogenous ppl progenitor cells (green) internalizing at the dorsal germ ring margin in a Tg(-4gsc:eGFP-Hsa.HRAS) embryo injected with LynTagBFP mRNA to label the membrane of all cells (red) and fluorescent Dextran (blue) to label the interstitial fluid. Frame rate, ~2 fps; ~2.5 min/frame. Scale bar, 20 μm.
Movie 3 – Internalization of transplanted mesoderm cells.
Confocal image series of internalizing Tg(βActin:Ras-eGFP) mesoderm cells (green) transplanted directly below the EVL at the dorsal germ ring of a Tg(dharma:eGFP);MZoep host embryo, expressing LynTagBFP in all cells to outline the plasma membrane (red) and injected with fluorescent Dextan (blue) to label the interstitial fluid. The transplanted donor cells migrate from the surface to the inside of the germ ring in the host embryo. Frame rate, ~ 2 fps; ~ 4 min/frame. Scale bar, 20 μm.
Movie 4 - Finite Element simulations of progenitor cell sorting using relative interfacial tension distributions determined *in vitro* and *in vivo*.

The movie shows two Finite Element (FE) simulations with identical starting configuration of heterotypic aggregates composed of ectoderm (red) and mesoderm (green) cells surrounded by culture medium (blue). In these finite element-based simulations, the cells in the heterotypic aggregate shown on the left side of the movie were assigned the relative tensions measured by CellFIT-3D in *in vitro* aggregates under standard culture conditions ($\gamma_{e-e} = 1.00$, $\gamma_{m-m} = 1.31$, $\gamma_{e-m} = 1.66$, $\gamma_{e-cm} = 2.65$, $\gamma_{m-cm} = 1.20$; as in Fig. 1C). The model cells (ectoderm cells are shown in red and mesoderm cells in green) displayed a clear sorting behaviour similar to their experimental counterparts. If instead the interfaces are assigned the tensions that CellFIT-3D shows to act in *in vivo* ($\gamma_{e-e} = 1.00$, $\gamma_{m-m} = 1.28$, $\gamma_{e-m} = 1.25$, $\gamma_{e-cm} = 0.78$, $\gamma_{m-cm} = 0.63$; Fig. 1J), cells do not sort (right part of the movie). The simulations of the modelled cell aggregates have the same initial configuration and are time synchronized (5000 iterations). Frame rate, $\sim$ 30 fps.
Movie 5 - Finite Element simulations of ectoderm-mesoderm tissue envelopment using relative interfacial tension ratios obtained from those tissues in the presence of medium with 300 mOsm/L.

When ectoderm and mesoderm aggregates are co-cultured in 300 mOsm/L medium for 5 h, the CellFIT-3D analyses obtained relative interfacial tensions distribution are $\gamma_{e-e} = 1.00$, $\gamma_{m-m} = 1.62$, $\gamma_{e-m} = 2.41$, $\gamma_{e-if} = 2.94$, $\gamma_{m-if} = 1.58$ (see also Fig. 4B). When these relative tensions act in two homotypic model aggregates brought into contact (leftmost part of the movie), the mesoderm cell aggregate partially envelops the ectoderm cell aggregate. Frame rate, ~ 30 fps.
Movie 6 - Finite Element simulations of ectoderm-mesoderm tissue envelopment using relative interfacial tension ratios obtained from those tissues in the presence of medium with 250 mOsm/L.

When ectoderm and mesoderm aggregates are co-cultured in 250 mOsm/L medium for 5 h, the CellFIT-3D analyses obtained relative interfacial tensions distribution are $\gamma_{e-e} = 1.00$, $\gamma_{m-m} = 1.16$, $\gamma_{e-m} = 1.32$, $\gamma_{e-if} = 1.52$, $\gamma_{m-if} = 1.41$ (see also Fig. 4E). When these relative tensions act in two homotypic model aggregates brought into contact, the mesoderm cell aggregate does not envelope the ectoderm cell aggregate. Frame rate, ~ 30 fps.
Movie 7 – Tracking of endogenous mesoderm cell internalization.

Confocal image series of ppl progenitor cell (green) internalization within the dorsal germ ring margin of a Tg(-4gsc:eGFP-Hsa.HRAS) embryo injected with Lyn-TagBFP mRNA to outline the plasma membrane of all cells (red) and Dextran-Rhodamin to label the interstitial fluid (blue). Cell tracks delineate the movements of exemplary ppl (green spheres) and ectoderm (red spheres) progenitor cells during internalization (5 – 6.5 hpf). Frame rate, ~ 2 fps; ~ 2.5 min/frame. Scale bar, 20 μm.
**Movie 8 - Internalization of DN-Rac expressing mesoderm cells.**

Confocal image series of ppl progenitor cell (green) internalization in a Tg(-4gsc:eGFP-Hsa.HRAS) embryo (5 – 6.5 hpf), overexpressing DN-Rac to inhibit progenitor cell protrusion formation, Lyn-TagBFP to outline the plasma membrane (red), and Dextran-Rhodamin to label the interstitial fluid (blue). Frame rate, ~ 2 fps; 3 min/frame. Scale bar, 20 μm.
Movie 9 - Cell protrusion formation of DN-Rac expressing mesoderm cells.

Image series of induced ppl progenitor cell aggregates isolated from a Tg(bAct:utro-mCherry) embryo and seeded on a fibronectin-coated substrate for 1.5 h. Upper panels display bright-field images and lower panels show fluorescent Utrophin-mCherry to label actin. Left panels are WT ppl cells and right panels are DN-Rac expressing ppl cells. Frame rate, ~ 6 fps; 2 min/frame. Scale bar, 10 μm.
Supplementary References

