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Nodal Signaling has dual roles in fate specification and directed migration during germ layer segregation

by

Zairan Liu

DISSERTATION

Submitted in partial satisfaction of the requirements for the degree of

DOCTOR OF PHILOSOPHY

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Biophysics

in the

GRADUATE DIVISION

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Ву

Zairan Liu

For my family,

for their love and support

ACKNOWLEDGEMENTS

First, I would like to thank my advisor, Orion Weiner. I first met Orion during the interview for iPQB program at UCSF. He was the last interviewer of the day and I had been excited and overwhelmed, nervous and exhausting for a whole day. But during that very 30 minutes of interview, I felt the breathtaking joy of scientific discovery. He told me a story of membrane tension and cell migration. He told it in a way that was like a mystery novel. Piece and piece of evidence unfolds; theories tested, disprove then led to the next. Then the Eureka moment, when it was unveiled, so much excitement fulfilled the young and ignorant me, which led me here for grad school.

Since rotation, Orion has been a great mentor to guide me through identifying my own research interest. After I told him I was interested in systems biology from a multicellular perspective, he came up with a very detailed list of classical papers in cross of these fields for me to read deeply. Then we would hold weekly discussions to identify more interesting topics, such as the cellular system, the signaling dynamics, or the technology. Through this process, I had a much deeper understand of the field and dissection of the interesting parts of biology. This had been such a valuable experience and had sculptured my critical thinking process.

Orion is always supportive of students coming up with their own projects. The research interest in the lab is cell decision making. When I joined the lab in 2014, there were people working on all kinds of systems, ranging from yeast to dicty to T cell to neutrophils to zebrafish to chick embryos. All the systems have been brought in by trainees and

developed independently. When I was thinking of my own thesis project, he helped me do a couple brainstorming sessions and put all the pieces together. I was able to come up with a project on my own instead of taking on someone else's or just doing what he picked for me. Since then, Orion has been very helpful to structure the project and lead me to different professors in the field to chat and learn more about it. I was very gracious for being able to conduct research independently from early on and feeling immensely supported at the same time.

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STATEMENT REGARDING AUTHOR CONTRIBUTIONS

Conceptualization, Z.L. and O.W.; Methodology, Z.L. and S.W.; Formal Analysis, Z.L.; Writing – Original Draft, Z.L.; Writing – Review & Editing, O.W. and S.W.; Funding Acquisition, Z.L., S.W., and O.W.

Nodal signaling has dual roles in fate specification and directed migration during germ layer segregation

Zairan Liu

During gastrulation, endodermal cells actively migrate to the interior of the embryo, but the signals that initiate and coordinate this migration are poorly understood. By transplanting ectopically-induced endodermal cells far from the normal location of endoderm specification, we identified the inputs that drive internalization without the confounding influences of fate specification and global morphogenic movements. We find that Nodal signaling triggers an autocrine circuit for initiating endodermal internalization. Activation of the Nodal receptor directs endodermal specification through sox32 and also induces expression of more Nodal ligands. These ligands act in an autocrine fashion to initiate endodermal cell sorting. Our work defines an "AND" gate consisting of sox32-dependent endodermal specification and Nodal ligand reception controlling endodermal cell sorting to the inner layer of the embryo at the onset of gastrulation.

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CHAPTER ONE: INTRODUCTION

During development, a single cell gives rise to hundreds of different cell types. Through proliferation, differentiation, migration and death, organisms generate complex patterns of cell fates and morphologies in space and time. This process, termed pattern formation, creates order among a collective of cells. Recent studies have suggested a self-organizing framework for symmetry breaking and patterning in embryogenesis(Karsenti 2008; Wennekamp et al. 2013). Self-organization is defined as a process where global order spontaneously arises out of the local interactions among the components of an initially disordered system. This process usually involves feedback loops that confer robustness to the system. Development contains highly regulative capacity revealed by experiments done in 19th century(Davidson 1991), which indicates interdependent factors underlie embryonic pattern formation. But what the factors are and how they interact to confer such an emergent phenomenon remain unclear up till this day.

How different cell fates are determined has been a central question in the field of development. There are mainly two modes of specification: autonomous specification and conditional specification. Autonomous specification means that a developing cell is able to differentiate only with its intrinsic properties without receiving any external signals. In contrast to autonomous specification, conditional specification means that cells rely on interactions with neighboring cells or morphogen, which imparts positional information in a morphogenic field with its local concentration. Lewis Wolpert refined the concept of morphogen in 1960s as a French Flag Model(Wolpert 1969). Different colors of the French Flag represent different states (fate choices) of the cell, which are controlled by

concentrations of the morphogen. Forty years have gone by and people have been mainly focusing on how far the positional information can be transmitted and how precise it can be specified. However, to establish a precise pattern, morphogen gradients alone are not sufficient. Cell movement has been proven to be an essential part of the pattern formation process. Global morphogenesis is critical to establish embryonic patterning, as well as local cell movement to refine initial patterning. As an example of the complex interplay of cell fate specification and cell movement, gastrulation crystalizes both processes and offers a platform to dissect the factors at play.

Gastrulation is central to animal development and involves the specification of three different germ layers (endoderm, mesoderm, and ectoderm) and their segregation to different locations in the embryo (Wolpert 1992). In contrast to the mechanisms underlying cell fate specification, the mechanisms used to drive segregation of the three germ layers are much less well understood. In this work, we focus on endodermal cells, which are initially specified on the surface of the embryo but must segregate to the interior, where they give rise to the gut and associated tissues. Endoderm migration is crucial for the formation of the gut tube and digestive tract across the animal kingdom. The in-folding of surface blastoderm cells to form the endoderm is well-documented in a wide range of species (Stern 2004; Wolpert 1992). However, it has been experimentally difficult to separate the initiation of migration events from cell fate specification. Thus, the molecular logic of the cell internalization, including which signals trigger this migration and how cell fate and migration are related, still remain unclear.

Several models have been proposed for how the germ layers segregate during embryogenesis. Most prominently, the differential adhesion hypothesis proposed that is that differences in intercellular adhesion among the different germ layers drives sorting (Steinberg 1962). However, although differential adhesion and cortex tension have been observed *in vitro*, *in vivo* measurements of tissue surface tension were indistinguishable among the three germ layers. Thus, differential adhesion is unlikely to fully account for the ability of the germ layers to sort in the embryo (Krieg et al. 2008; Maître et al. 2012; Krens et al. 2017).

Previous studies have shown that directed cell migration appears to be the driving force for endoderm segregation *in vivo* for zebrafish (Montero et al. 2005; Krens et al. 2017; Giger and David 2017). At the onset of zebrafish gastrulation, the blastoderm consist of several thousand cells positioned above the yolk cell. Internalization begins on the dorsal side where inward-moving cells form the hypoblast (mesoderm and endoderm) in contrast to the cells remaining on the outside as epiblast (ectoderm) (Warga and Kimmel 1990). A germ ring forms at the boundary of hypoblast and epiblast and the embryonic shield is formed on the dorsal side of the margin. Early dye labeling experiments showed that cells relocate to deeper levels within the germ ring by inverting their order relative to the margin as they internalize (Kimmel and Warga 1987). An involution model was proposed to describe the population flow as a cellular sheet (Lewis 1985). Later, time-lapse tracking showed that individual cells within germ ring transiently move out of the epiblast and relocate into the hypoblast (D'Amico and Cooper 1997; Concha and Adams 1998). More

recent studies have shown such cells extend protrusions inward and exhibit active directed migration (Montero et al. 2005; Krens et al. 2017; Giger and David 2017).

Nodal, as a member of the TGF-β superfamily, is essential for germ layer patterning in zebrafish. Nodal ligands are expressed at the margin and yolk syncytial layer (YSL) during the blastula stage, where it forms a morphogen gradient (Chen and Schier 2001; Dougan et al. 2003). The signaling pathway is activated by Nodal binding to a type II TGF-β receptor, inducing interaction with a EGF-CFC co-receptor, One-eyed-pinhead (Tdgf1), and the type I TGF-β receptor, Acvr1ba (Weng and Stemple 2003; Gritsman et al. 1999; Aoki et al. 2002). Subsequent phosphorylation of the transcription factors Smad2 and Smad3 facilitates the formation the Smad-complexes, which translocate into the nucleus to regulate the expression of target genes (Weng and Stemple 2003; Jia et al. 2008). One of the key downstream targets is the sox32, which plays an essential cell-autonomous role in endoderm formation (Kikuchi et al. 2001). Surprisingly, the signals which initiate and direct endoderm migration are not well understood. From previous studies, it is known that endodermal cells undergo random walk regulated by chemokine signaling downstream of Nodal pathway (Nair and Schilling 2008). Our lab has recently identified that Nodal signaling regulates endodermal cell motility and actin dynamics via Rac1 and Prex1 (Woo et al. 2012). However, it is not known whether endodermal cell fate alone is sufficient to drive migration or whether additional cues in their morphogenic field are required.

Here we utilized an *in vivo* system to study germ layer segregation in zebrafish embryos. In the early zebrafish embryo, an initially mixed mesendodermal population ultimately resolves into distinct mesodermal and endodermal cell layers, but these complex morphogenetic movements occur simultaneously with fate specification (Ho 1992). To disentangle the endodermal specification program from the migration program, we used a constitutively-active version of the Nodal receptor, acvr1ba*, to predispose cells into an endodermal fate (Renucci, Lemarchandel, and Rosa 1996; Aoki et al. 2002; David and Rosa 2001). By transplanting these ectopically-induced endodermal cells into the animal pole of the embryo, we removed them from the endogenous signals that normally orchestrate endodermal development as well as the effects of nearby involuting cells. We found that these ectopically-introduced endodermal cells do not take the normal path of endogenous endoderm migration by internalizing at the germ ring; instead they radially ingress into the inner layer. Nodal signaling is necessary and sufficient to initiate this process, and the ectopic endodermal cells (but not the surrounding cells) need to receive the Nodal ligand in an autocrine fashion to trigger ingression. Our results suggest that Nodal signaling plays dual roles in specifying endodermal fate and initiating the sorting of these cells to the interior of the embryo. As these migration events are not observed for in vitro culture conditions, this in vivo approach for endodermal sorting should be a powerful system for continued dissection of the logic of germ layer segregation during gastrulation.

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CHAPTER TWO: ACVR1BA* - INDUCED ECTOPIC ENDODERMAL CELLS INGRESS INTO THE INNER LAYER OF THE EMBRYO

To determine the requirements for initiating and directing endoderm migration, we developed a cell transplantation model that allowed us to directly guery endodermal sorting while disentangling the endodermal specification program from the migration program. We generated ectopic endodermal cells by expression of the constitutively activated Nodal receptor acvr1ba*. We then transplanted these cells (Fig. 1A) into the animal pole of a wild-type host embryo, far from the marginal location of endogenous endodermal cells, and determined whether these misplaced ectopic endodermal cells could sort into the correct endodermal layer. (Fig. 1B). First, ectopic endoderm production by acvr1ba* was confirmed by qPCR analysis of sox17 and sox32 expression (Fig. 2A), markers for endodermal cell fate (Kikuchi et al. 2001, 17; Shivdasani 2002). Although Nodal signaling can also induce mesoderm fate (Peyriéras, Strähle, and Rosa 1998; Aoki, Mathieu, et al. 2002), we found that acvr1ba* expression upregulated the mesodermal markers gsc and ntl to a lesser extent that sox17 and sox32 (Fig. 2B), demonstrating that acvr1ba*-expressing cells are biased to an endoderm fate. Next, we found that, after transplantation to the animal pole, these ectopically introduced endodermal cells accumulated in endoderm-derived tissue by preferentially migrating to the correct endodermal layer (Fig. 1B-E). When induced endodermal cells were transplanted together with non-endodermal cells (Fig. 1F), these cell types separated into two layers from an originally mixed population (Fig. 1G). Visualizing the migration path of these cells by time-lapse microscopy showed that induced endodermal cells did not move towards the margin and then involute to form endodermal layer (the normal path of endogenous

endoderm migration); instead, they radially ingressed into the inner layer (**Fig. 1G**), consistent with a recent report (Giger and David 2017). Additional single-cell tracking analysis revealed that the trajectories of transplanted cells did not exhibit random walk or sample both inward and outward directions; instead, the ingression was highly unidirectional (**Fig. 1H**). These data indicate that endodermal cells produced by acvr1ba* expression can initiate ingression if placed ectopically via highly polarized and unidirectional migration.

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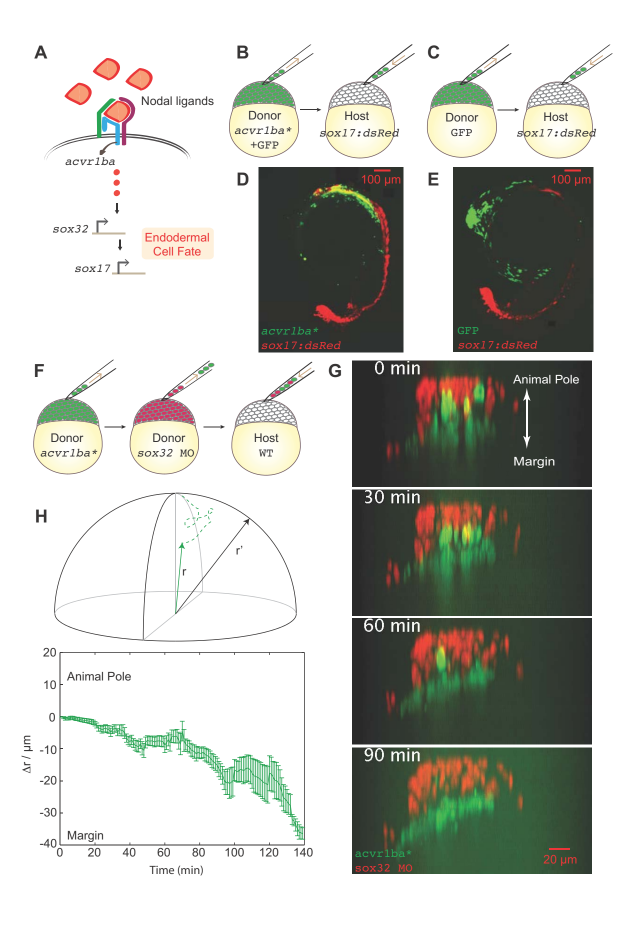


Figure 1:

Constitutively active Nodal receptor (acvr1ba*)-induced ectopic endodermal cells sort into the inner layer of the embryo by ingression.

- (A) Schematic diagram depicting Nodal signaling and specification of endodermal cell fate. Nodal ligands activate the *acvr1ba* receptor and signal to *sox32*, a transcription factor controlling endodermal specification.
- **(B-E)** Schematic diagrams of ectopic endoderm transplant assay (B,C) and representative results (D, E). *acvr1ba**-expressing or control cells were transplanted to the animal pole of *Tg(sox17:dsRed)* host embryos. At 21-somite stage, transplanted *acvr1ba**-expressing cells localized to endoderm-derived tissue, primarily the pharynx, (D) while control transplanted cells localized to non-endodermal tissue, particularly the head (E).
- **(F)** Schematic diagram of the double donor transplant assay. Donor endodermal cells expressing *acvr1ba** (green) were transplanted together with non-endodermal donor cells injected with *sox32* MO (red) to the animal pole of a single wild-type host.
- **(G)** Still images from a time-lapse movie of a WT host containing both *acvr1ba**-expressing (green) and *sox32* MO-containing (red) donor cells. Time lapse microscopy began immediately after transplantation (0 min). Over time, *sox32* MO donor cells remain in the outer layer of the embryo, while *acvr1ba**-expressing donor cells migrate into the inner layer of the embryo. Data was resliced and projected onto the XZ plane, with the animal pole towards the top and the margin towards the bottom.
- **(H)** Single-cell tracking analysis of ingression. Top panel: Cartesian coordinates for transplanted cells were transformed into spherical coordinates. Dashed lines represent cell trajectories. The radial distance, r, was measured as the distance from each cell's position at the end of the time lapse movie to the center of the embryo (solid lines). r' was measured as the distance to the host surface for normalization. Bottom panel: Average relative distance with standard error of *acvr1ba**-expressing cells plotted against time. Relative distance for each time point was calculated by measuring the radial distance of *acvr1ba**-expressing cells to the center of the embryo, subtracted by the distance of host cell expanding during gastrulation.

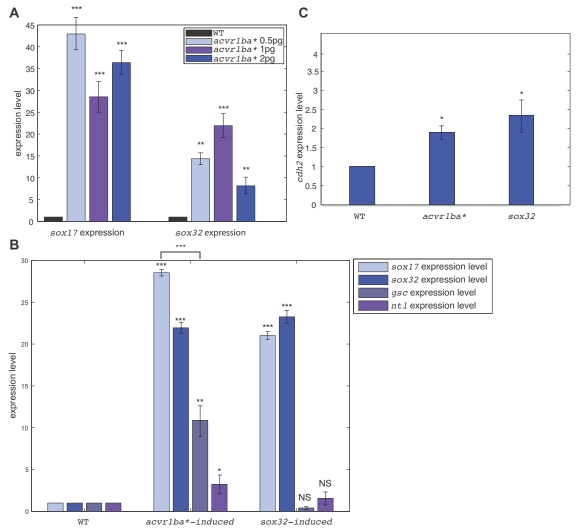


Figure 2: acvr1ba* induces expression of sox17 and sox32.

- **(A)** Expression of sox17 and sox32 endodermal markers was measured by real-time quantitative PCR. Constitutive activation of the Nodal pathway by expression of $acvr1ba^*$ upregulated sox17 and sox32 expression (normalized to uninjected controls). **p<0.01, ***p<0.001.
- **(B)** Expression of *sox17*, *sox32*, *gsc* and *ntl* was measured by real-time quantitative PCR in *acvr1ba**-expressing cells and *sox32*-expressing cells in wildtype background. Both acvrb1a* and sox32 more potently induce endodermal markers (*sox17* and *sox32*) than mesodermal markers (*gsc* and *ntl*). *p<0.05, **p<0.01, ***p<0.001, NS, not significant.
- **(C)** Expression of *cdh2* at 6hpf was measured by real-time quantitative PCR. Both *acvr1ba** and *sox32*-induced endodermal cells have elevated expression comparing to wild type uninjected controls. *p<0.05.

CHAPTER THREE: NODAL LIGAND EXPRESSION IS NECESSARY TO TRIGGER THE SORTING OF ECTOPIC ENDODERMAL CELLS

We next sought to define the molecular logic of ectopic endodermal cell sorting. In addition to expressing the constitutively activated Acvr1ba* receptor, ectopic endodermal cells can also be produced by overexpression of the transcription factor Sox32, a target of Nodal signaling (Dickmeis et al. 2001; Kikuchi et al. 2001; Sakaguchi, Kuroiwa, and Takeda 2001, 32). However, previous work suggested that, unlike Acvr1ba* expression, overexpression of sox32 is not sufficient to drive the sorting process (Kikuchi et al. 2001). We too observed that cells overexpressing sox32 could preferentially segregate to endoderm-derived tissues when placed near the dorsal margin but not when transplanted to the animal pole (Fig. 1A-B, 2).

Notably, this means that these two different means of generating endodermal cells (acvr1ba* vs sox32) are not equivalent in their ability to drive internalization movements when transplanted far from the normal endodermal domain (**Fig. 1C**); only acvr1ba* induced endodermal cells are capable of ingression when placed at the animal pole. These data suggest that Nodal signaling initiates endodermal sorting in addition to specifying endodermal fate. acvr1ba* likely activates additional pathways that are absent when sox32 is overexpressed, thus allowing cells to sort regardless of the location within the embryo. In contrast, cells overexpressing sox32 may require extrinsic factors present at the margin to activate these additional "sorting" pathways, explaining why they can only sort in regions close to the margin (Kikuchi et al. 2001). These observations suggest that

the triggering of sorting involves an "AND" gate consisting of *sox32*-dependent endodermal specification and additional signaling downstream of *acvr1ba** (**Fig. 1D**).

To better understand the differences between these two methods of generating ectopic endoderm, we compared the signaling and transcriptional networks activated by *acvr1ba* and *sox32*. A recent report (Giger and David 2017) suggested that N-cadherin (cdh2) expression triggers endoderm ingression. However, we found that both acvr1ba* and sox32 overexpression induced cdh2 expression (**Fig. 2C** from Chapter Two). This suggests that N-cadherin expression alone does not account for the difference of ingression capability between these two types of endodermal cells.

Nodal signaling through *acvr1ba*, but not *sox32* alone, is capable of inducing expression of Nodal ligands *ndr1* and *ndr2* (Chan et al. 2009; Feldman et al. 1998; Dougan et al. 2003). In wild-type embryos, these ligands are expressed highest near the margin, which could explain the observation that sox32-induced endodermal cells only sort in this location. In contrast, the autocrine production of Nodal ligands downstream of acvr1ba* could enable these cells to sort regardless of location in the embryo. To confirm that ndr1/2 is secreted by acvr1ba*-expressing cells but not by cells expressing sox32- alone, we quantified the ndr1/2 expression profile under all experimental conditions (**Fig. 3**). When transplanted to WT embryos, we saw a 2-3 fold increase of ndr1/2 expression in acvr1ba*-expressing cells compared to the sox32-expressing cells. However, this is likely an underestimate of the difference in nodal ligand expression induced by acvr1ba* or sox32 expression due to the presence of as the maternally deposited ndr1/2 and Nodal

positive feedback loop. To address this complication, we expressed acvr1ba* and sox32 MZ *tdgf1* background embryos, which lack sufficient Nodal signaling (Gritsman et al. 1999) (Fig. 3B). In this background, we found that in acvr1ba*-expressing cells increased ndr1 expression 25-fold higher than sox32-expressing cells. Finally, injection of ndr1/2 mRNA increased the expression of ndr1/2 in sox32-expressing cells to similar levels seen in acvr1ba*-expressing cells. Taken together, these data confirm that acvrb1a*, but not sox32, induces nodal ligand expression (Fig. 3C).

Next, we sought to test whether the excess production of Nodal ligands is the driver of ingression. We first examined the necessity of Nodal ligand expression for endodermal sorting by using morpholinos against *ndr1* and *ndr2* in the *acvr1ba**-induced endodermal cells, which were then transplanted into the animal pole of wild-type embryos (**Fig. 1E**). We verified the functionality of the morpholinos by demonstrating the inhibition of *sox17* expression (**Fig. 4**). At 20 hpf, while cells expressing *acvr1ba** only preferentially localized to endodermal tissues such as the pharynx, the *acvr1ba** cells with *ndr1* and *ndr2* MO knockdown primarily localized in non-endodermal tissue, particularly in the head region (**Fig. 1F**, **H**). Single cell tracking revealed that the *acvr1ba** cells with *ndr1* and *ndr2* MO remain in the ectoderm and move near the surface of the embryo (**Fig. 1I**). However, knockdown of *ndr1* and *ndr2* did not affect the ability of acvr1ba* to induce ectopic endoderm production as demonstrated by *Tg(sox17:GFP)* reporter expression (**Fig. 1J-K**), suggesting endodermal specification is unaffected.

The other essential input into the "AND" gate is sox32-dependent endoderm specification (**Fig. 1D**). To confirm that it is necessary to have sox32-dependent endoderm specification to sort properly, we conducted similar transplantation experiments where sox32 MO was injected into the acvr1ba*-induced endodermal cells as the donor, these cells lost the ability to ingress into the inner layer of the embryo after transplanted to the animal pole of a wild-type host (**Fig. 1G, H**). These results suggest that both autocrine production of Nodal ligands and sox32-dependent endodermal specification are necessary to trigger ectopic endodermal sorting.

To further test if ndr1/2 secretion and autocrine signaling are critical to sorting, we analyzed the internalization dynamics of marginally-transplanted cells in host embryos injected with ndr1/2 MO. After transplantation into the margin of a ndr1/2-depleted host, acvr1ba*-expressing cells localized to the endoderm-derived tissue (green). Cells overexpressing sox32 localized to both endoderm and ectoderm-derived tissue (though statistically less endodermal localization compared to acvr1ba* cells) (**Fig. 5**). Note that for cells overexpressing sox32, there is still global morphogenesis happening at the margin for the host cells. This might account for the increased percentage of internalized endoderm comparing to transplantation to the animal pole. This corroborates with previous literature showing MZ *tdgf1* mutant cells transplanted to the margin of wild-type blastula embryos initially internalize with their neighbors (Carmany-Rampey and Schier 2001).

In zebrafish, the two Nodal ligands Ndr1 and Ndr2 are known for their functional redundancy in inducing mesendoderm fate (Feldman et al. 1998; Erter, Solnica-Krezel, and Wright 1998; Jing et al. 2006). But it is not known whether they behave redundantly to induce ingression. To address this question, we performed transplantation experiments of acvr1ba*-expressing cells with either ndr1 MO or ndr2 MO alone. Our results showed that neither ndr1 MO or ndr2 MO abolished the ingression behavior of acvr1ba*-expressing cells (**Fig. 6**). These data suggest that these nodal ligands act redundantly to support the ingression of acvr1ba*-expressing cells.

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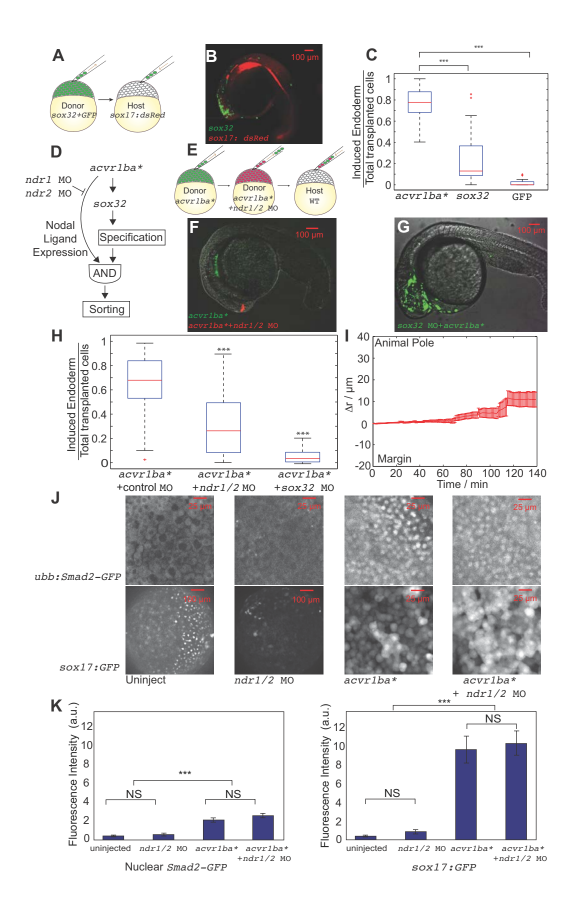


Figure 1:

Nodal ligand expression is necessary to trigger the sorting of ectopic acvr1ba*-induced endodermal cells.

- (A-B) Schematic diagrams depicting sox32-induced ectopic endoderm transplant assay (A) and representative result (B). sox32-overexpressing cells were transplanted to the animal pole of wild-type host embryos. At 21-somite stage, transplanted sox32-overexpressing cells primarily localized to non-endodermal tissues, primarily in the head and skin.
- **(C)** Boxplot quantification of endoderm contribution of transplanted cells at 20 hpf, assessed by co-localization with Tg(sox17:dsRed) expression. On each box, the central mark indicates the median, and the bottom and top edges of the box indicate the 25th and 75th percentiles, respectively. The whiskers extend to the most extreme data points not considered outliers, and the outliers are plotted individually using the '+' symbol. Compared to $acvr1ba^*$ -expressing cells, fewer cells overexpressing sox32 contributed to endodermal tissues. Data is shown as mean \pm SEM of 3 independent transplantation experiments with 26 embryos per condition. Student's t-test was performed. * p<0.05, ** p<0.01, *** p<0.001.
- **(D)** Schematic diagram depicting a potential AND gate for endoderm sorting. Constitutively activate *acvr1ba** upregulates both *sox32* as well as Nodal ligand expression. Only with both inputs do cells successfully sort to the inner layer of the embryo.
- (E) Schematic diagram of the cell transplantation assay to test the necessity of Nodal ligand expression for cell sorting. Donor cells containing ndr1 and ndr2 MOs plus acvr1ba* mRNA (red) were transplanted together with cells overexpressing acvr1ba* only (green) into the animal pole of wild-type host embryo. (F) Representative images showing distribution of transplanted cells at 21-somite stage. Cells expressing acvr1ba* only (green) localize to endoderm-derived tissue, primarily the pharynx. Cells containing acvr1ba* along with ndr1 MO and ndr2 MO (red) localize to non-endodermal tissue, primarily in the head. Lateral view, anterior to the bottom-left.
- **(G)** Representative images showing distribution of transplanted cells at 21-somite stage. Cells expressing *acvr1ba** along with *sox32* MO (green) localize to ectoderm-derived tissue, primarily the head. Lateral view, anterior to the bottom-left.
- **(H)** Boxplot quantification of endoderm contribution of transplanted cells at 20 hpf. *ndr1* and *ndr2* knockdown as well as sox32 knockdown reduced the ability of $acvr1ba^*$ -expressing cells to contribute to endodermal tissue. Data is shown as mean \pm SEM of 3 independent transplantation experiments with 22 embryos per condition. Student's t-test was performed. * p<0.05, ** p<0.01, *** p<0.001.
- (I) Single-cell tracking analysis of ingression. Average relative distance with standard error of acvr1ba*-expressing cells with *ndr1* MO and *ndr2* MO plotted against time. Relative distance was calculated as in Fig. 1H.
- (J) Nodal signaling level assessed by Tg(ubb:Smad2-GFP) and Tg(sox17:GFP). Top panel: Smad2-GFP showed no nuclear localization in the center of animal poles (AP) of uninjected embryos and ndr1/ndr2 morphants. Smad2-GFP showed comparable levels of nuclear localization in $acvr1ba^*$ -injected embryos and $acvr1ba^*$ with and ndr1 and ndr2 MOs-injected embryos. Full images in Fig. S2. Bottom Panel: Sox17:GFP labels wild-type endodermal cells in the uninjected control embryo but few GFP-positive cells are

present in the *ndr1/ndr2* morphants. Animal Pole View. Sox17:GFP shows elevated level of expression in both *acvr1ba** injected embryos and *acvr1ba** with and *ndr1* and *ndr2* MOs injected embryos. Lateral view. Margin depicted by dash line.

(K) Quantification of Nodal signaling level. Nuclear Smad2-GFP and Sox17:GFP fluorescence levels are quantified. Data is shown as mean ± SEM of 3 independent embryos.

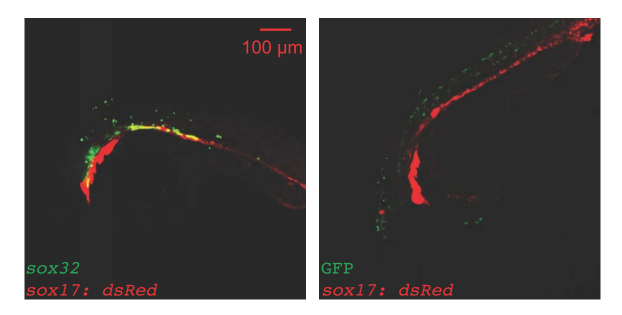


Figure 2: sox32 preferentially segregate to endoderm-derived tissues when placed near the dorsal margin.

Representative images showing distribution of *sox32*-overexpressing cells or GFP-expressing cells that were transplanted to the margin of wild-type host embryos. At 21-somite stage, transplanted *sox32*-overexpressing cells primarily localized to endodermal tissues while GFP-expressing cells localized to mesodermal tissues.

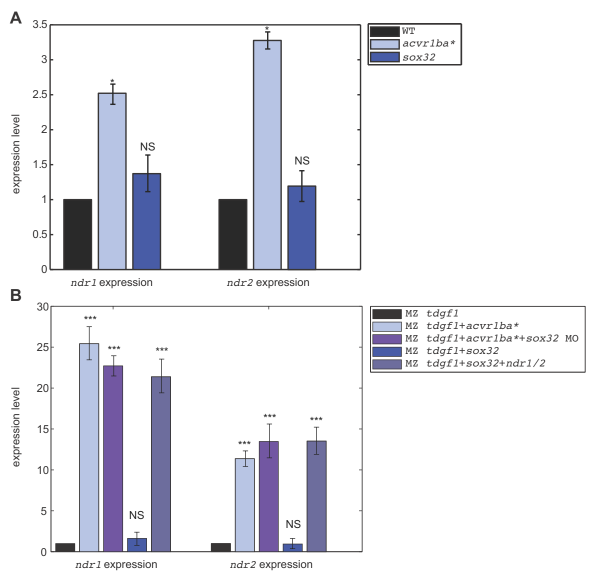


Figure 3: ndr1/2 is upregulated by acvr1ba*, and sox32 is neither necessary or sufficient for this upregulation.

- **(A)** *ndr1/2* expression in *acvr1ba**-expressing cells and *sox32*-overexpressing cells in wildtype background measured by real-time quantitative PCR. *p<0.05, NS, not significant.
- **(B)** *ndr1/2* expression under all experimental conditions in MZ *tdgf1* background, which removes the confounding effects of maternally deposited Ndr1/2 on driving nodal signaling. ***p<0.001, NS, not significant.

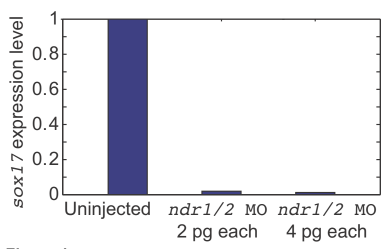


Figure 4: Validation of *ndr1* and *ndr2* morpholinos.

Quantification of *sox17* expression level, demonstrating that the *ndr1* and *ndr2* morphants block the endogenous Nodal gradient and endoderm specification.

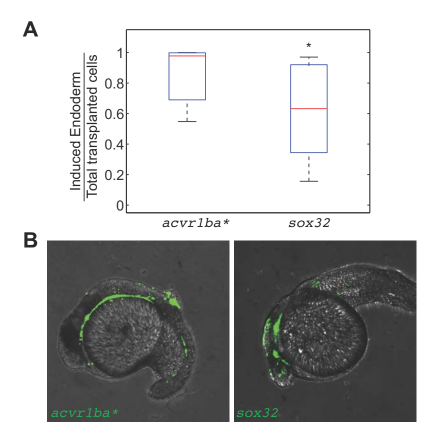


Figure 5: Induced endodermal cells internalize following transplantation to the margin.

- (A) Boxplot quantification of endoderm contribution of transplanted cells at 18 hpf. Data is shown as mean ± SEM of independent transplantation experiments with 14 embryos per condition. Student's t-test was performed. * p<0.05.
- **(B)** Representative image showing distribution of transplanted cells depicted in (A) at 18 hpf. *acvr1ba**-expressing cells localized to the endoderm-derived tissue (green). Cells overexpressing *sox32* localize to both endoderm and ectoderm-derived tissue. Lateral view, anterior to the left.

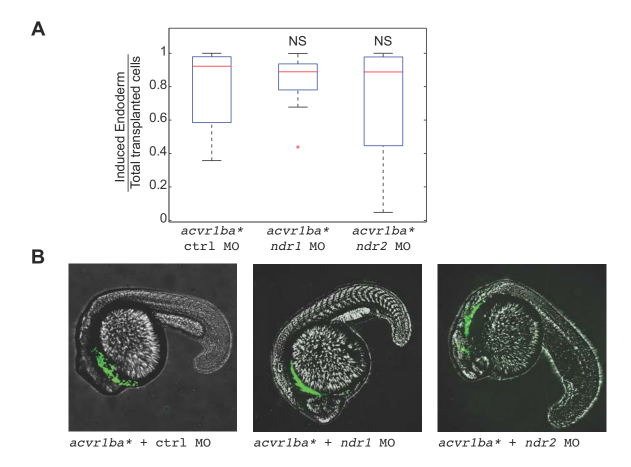


Figure 6:

Ndr1 and Ndr2 act redundantly to support the ability of acvr1ba* cells to internalize.

- **(A)** Boxplot quantification of endoderm contribution of transplanted cells at 20 hpf. Data is shown as mean ± SEM of independent transplantation experiments with 16 embryos per condition. Student's t-test was performed. NS, not significant.
- **(B)** Representative image showing distribution of transplanted cells depicted in (A) at 18 hpf. *acvr1ba**-expressing cells localized to the endoderm-derived tissue (green)in all three conditions, in contrast to the block of internalization when both Ndr1 and Ndr2 MO are combined in acvr1ba* cells (**Fig. 1H**). Lateral view, anterior to the left.

CHAPTER FOUR: NODAL LIGAND EXPRESSION IS SUFFICIENT TO DRIVE INGRESSION OF SOX32-INDUCED ENDODERMAL CELLS

Next, we investigated whether addition of Nodal ligands could trigger ectopic endodermal cell sorting in sox32-induced endodermal cells, which were not able to ingress into the inner layer when they are transplanted to the animal pole (**Fig. 1A**). To test this, we injected donor cells with mRNA for both sox32 and Nodal ligands (ndr1, ndr2) prior to transplantation into wild-type embryos, using $acvr1ba^*$ -induced endodermal cells as a positive control (**Fig. 1B**). When examined at the18-somite stage, sox32-injected cells also expressing Nodal ligands accumulated in endoderm-derived tissue significantly better than cells expressing sox32 alone (**Fig. 1D, 1F-G**). These data indicate that the expression of Nodal ligands is sufficient to confer sorting ability in sox32-induced endodermal cells.

Can any cell expressing Nodal ligands sort to endodermal tissues, or do cells require both endodermal specification and Nodal ligand expression to support sorting? Because Nodal ligands themselves can drive endodermal fate(Chen and Schier 2001; David and Rosa 2001; Dougan et al. 2003), we addressed this question by overexpressing Nodal ligands in conjunction with *sox32* MO, which blocks the transcriptional program that initiates endodermal specification (Sakaguchi, Kuroiwa, and Takeda 2001; Dickmeis et al. 2001) (**Fig. 1C**). We found that Nodal ligands cannot support sorting in the *sox32* MO background (**Fig. 1E, 1G**). These data suggest that Nodal ligands can only trigger sorting in conjunction with *sox32*-dependent endodermal specification. Together, our necessity

and sufficiency experiments demonstrate that Nodal ligands and sox32 constitute an "AND" gate to initiate internalization in the early embryo.

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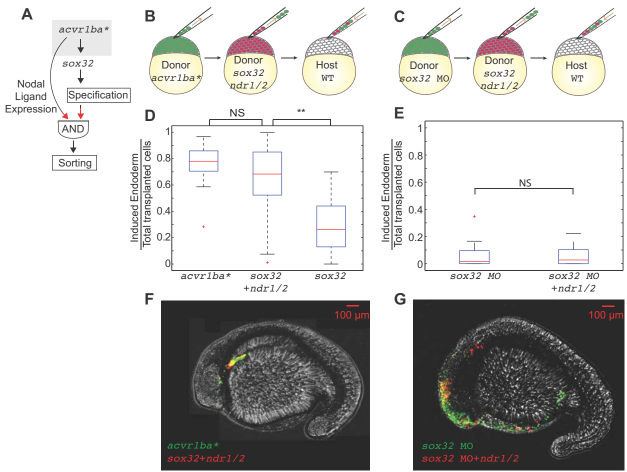


Figure 1: The combination of Nodal ligand expression and endodermal fate is sufficient to trigger ectopic endodermal cell sorting.

- (A) Schematic diagram depicting putative AND gate for endoderm sorting. Red arrows demonstrate the experimental perturbation to test the sufficiency of Nodal ligands to induce ingression.
- **(B-C)** Schematic diagrams of double transplantation assay to test the sufficiency of the AND gate depicted in (A) for endodermal sorting. Cells overexpressing *acvr1ba** (green) were transplanted together with cells overexpressing *sox32*, *ndr1*, and *ndr2* (red) into the animal pole of a wild-type host embryo (B). Cells containing *sox32* MO only were transplanted together with cells containing *sox32* MO as well as *ndr1* and *ndr2* mRNAs were transplanted into the animal pole of a wild-type host embryo (C).
- **(D)** Boxplot quantification of endoderm contribution at 20 hpf of transplanted cells depicted in (B). Cells overexpressing *sox32*, *ndr1*, and *ndr2* contributed to endoderm at a similar rate compared to cells overexpressing *acvr1ba**. Data is shown as mean ± SEM of 3 independent transplantation experiments with 18 embryos per condition. Student's t-test was performed. * p<0.05, ** p<0.01, *** p<0.001.
- **(E)** Boxplot quantification of endoderm contribution at 20 hpf of transplanted cells depicted in (C). Neither cells containing sox32 MO nor cells containing sox32 MO and overexpressing ndr1 and ndr2 contributed to endodermal tissue. In addition, cells expressing $acvr1ba^*$ and sox32 MO did not contribute to endodermal tissue. Data is

shown as mean ± SEM of 2 independent transplantation experiments with 14 embryos per condition. Student's t-test was performed. * p<0.05, ** p<0.01, *** p<0.001.

- **(F)** Representative image showing distribution of transplanted cells depicted in (B) at 18-somite stage. *acvr1ba**-expressing cells localized to the endoderm-derived tissue, primarily the pharynx (green). Cells overexpressing *sox32*, *ndr1*, and *ndr2* also localize to endoderm-derived tissue, primarily the pharynx (red). Lateral view, anterior to the left.
- **(G)** Representative image showing distribution of transplanted cells depicted in (E) at 18-somite stage. Cells expressing sox32 and Nodal ligands (ndr1, ndr1) localize to endodermal tissues similar to cells expressing $acvr1ba^*$. In contrast, sox32 MO-injected cells (green) and cells injected with sox32 MO and ndr1 and ndr2 mRNAs (red) localized to non-endodermal tissue, primarily in the head and skin.
- **(H)** Representative image showing distribution of transplanted cells injected with *sox32* MO and *acvr1ba** at 21-somite stage. Transplanted cells (green) mainly localized to non-endodermal tissue. Lateral view, anterior to the left.

CHAPTER FIVE: SORTING REQUIRES ECTOPIC ENDODERMAL CELLS TO RECEIVE NODAL SIGNALING IN AN AUTOCRINE CIRCUIT

So far, we have shown that Nodal ligands production is necessary and sufficient to trigger the ingression-based cell sorting of ectopic endodermal cells. We next determined which cells are responding to the Nodal ligands to support sorting. The Nodal ligands could either be acting on the same endodermal cells that undergo sorting to form an autocrine circuit or on the surrounding cells in a paracrine circuit, possibly by orchestrating endodermal extrusion by the surrounding ectoderm (Fig. 1A, 1B). We pharmacologically inhibited Nodal signaling by applying the Nodal receptor inhibitor SB505124 to block the autocrine reception of Nodal ligands in acvr1ba*-expressing cells. The Nodal inhibition drug SB505124 interferes with the signaling circuit upstream of acvr1ba, inhibiting endodermal cell fate specification (Fig. 2A-C). Following pharmacological Nodal inhibition, these cells fail to internalize after being transplanted to the animal pole (Fig. 2D). To inhibit only the autocrine reception of the Nodal ligands while maintaining endodermal cell fate, we used the maternal-zygotic (MZ) tdgf1 mutant to block Nodal signal reception; this mutant lacks the EGF-CFC co-receptor essential for the ability to signal downstream of Nodal ligands (Gritsman et al. 1999). When we transplanted MZ tdgf1 donor cells expressing acvr1ba* into wild-type recipient hosts, the donor cells were incapable of ingressing into the inner layer of the host embryo and did not contribute to endodermderived tissue at the 18-somite stage (Fig. 1C, 1F). In contrast, when we injected acvr1ba* into wild-type donor embryos and transplanted these cells into MZ tdgf1 mutant host embryos, these transplanted cells still successfully ingressed into the inner layer, indicating that ectopic endodermal cells retained their ability to sort irrespective of the

Nodal signaling state of the surrounding cells (**Fig. 1D**, **1G**). Together, these results suggest that an autocrine circuit of Nodal ligand reception is required to support sorting of ectopic endodermal cells (**Fig. 1E**). Furthermore, because the MZ *tdgf1* mutant host embryos lack endogenous endoderm (Gritsman et al. 1999; David and Rosa 2001) but still supported ingression of ectopic endodermal cells, these experiments further suggest that signals released by endogenous endodermal cells are not required for ectopic endodermal cell sorting.

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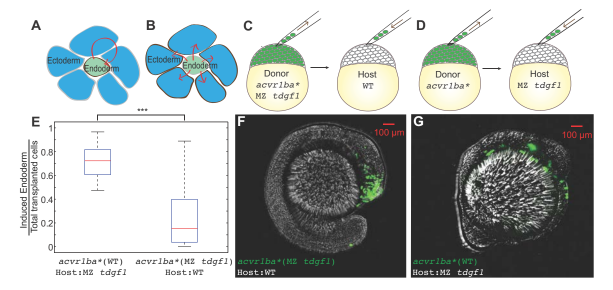


Figure 1: Nodal ligand reception acts cell autonomously to support sorting.

- (A-B) Schematic diagrams depicting autonomous (A) versus non-autonomous (B) Nodal ligand reception (red arrows).
- **(C)** Schematic diagram depicting single donor transplant assay to test cell autonomous Nodal signal reception. *acvr1ba**-expressing cells from MZ *tdgf1* donor embryos were transplanted to the animal pole of a wild-type host embryo.
- **(D)** Schematic diagram depicting single donor transplant assay to test cell non-autonomous Nodal signal reception. *acvr1ba**-expressing cells from wild-type donor embryos were transplanted to the animal pole of a MZ *tdgf1* host embryo.
- **(E)** Boxplot quantification of endoderm contribution at 18-somite stage for all transplanted cells. Wild-type donor cells expressing *acvr1ba** contributed to endodermal tissues while *acvr1ba**-expressing cells from MZ *tdgf1* embryos did not. Data is shown as mean ± SEM of 2 independent transplantation experiments, with 14 embryos per condition. Student's t-test was performed. * p<0.05, ** p<0.01, *** p<0.001.
- **(F)** Representative image showing distribution of MZ *tdgf1* cells expressing *acvr1ba** in a wild-type host. Donor cells (green) localized to ectoderm-derived tissue, primarily the head. Lateral view, anterior to the right.
- **(G)** Representative image showing distribution of wild-type cells expressing *acvr1ba** in a MZ *tdgf1* host. Donor cells (green) localized to endoderm-derived tissue. Lateral view, anterior to the right.

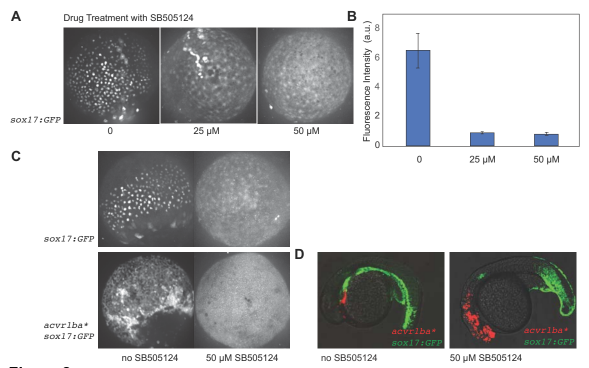


Figure 2:

Nodal signaling inhibitor SB505124 blocks *acvr1ba**-expressing cells from sorting. (A) sox17:GFP expression under 0, 25 μ M or 50 μ M SB505124 treatment at 6 hfp. Animal pole view.

- **(B)** Quantification of sox17:GFP expression in (A). *** p<0.001.
- (C) sox17:GFP expression for embryos with and without injection of acvr1ba* under no drug SB505124 treatment and 50 µM drug SB505124 treatment.
- **(D)** Transplant of *acvr1ba**-expressing cells into *sox17:GFP* background under no drug SB505124 treatment and 50 µM drug SB505124 treatment at 18hfp.

CHAPTER SIX: NODAL LIGANDS INITIATE BUT DO NOT GUIDE THE INGRESSION OF ENDODERMAL CELLS

In our transplant experiments, we observed ectopic endodermal cells moving from the outer layer of the embryo radially to the inner layer, but we never observed cells moving in the opposite direction (i.e., extruded from the embryo). A recent study has shown that during normal gastrulation movements, endodermal cells at the margin extend polarized protrusions toward the yolk syncytial layer and appear to internalize by active migration(Giger and David 2017). Therefore, we investigated whether ectopically placed endodermal cells similarly undergo active, directed migration to enter the interior of the embryo. To visualize actin dynamics in the ectopic endodermal cells during sorting, we expressed GFP-UTRN (Burkel, von Dassow, and Bement 2007), an actin reporter that we have previously used to analyze endodermal actin dynamics in zebrafish (Woo et al. 2012), in acvr1ba*-induced endodermal cells. We divided each single transplanted cell into two sectors, one facing towards the interior of the embryo and the other facing toward the embryo surface and then quantified the accumulation of actin in each sector. We observed a significant accumulation of actin in the interior sector of ectopic endodermal cells as well as actin-based protrusions extending towards the interior of the embryo can be seen as well. Control transplanted cells not expressing acvr1ba* lacked this polarity of actin enrichment and protrusions (Fig. 1A-B). In comparison, acvr1ba*-expressing cells injected with ndr1/2 MOs still produced protrusions but could not direct the protrusions properly to the inner layer, compared to the acvr1ba*-expressing cells (Fig. 2). Together, these data indicate that ectopic endodermal cells polarize their actin protrusions towards the inner layer of the embryo and that ndr1/2 are necessary to direct the actin-enriched protrusions, consistent with sorting based on active migration.

Which spatial cues are ectopic endodermal cells reading to achieve their directional migration? Such cues likely do not arise from the endogenous endodermal cells, as ectopic endoderm can still sort in an MZ tdgf1 host that lacks endogenous endoderm (Fig. 1E from Chapter Five). Might the endogenous Nodal gradient of the host embryo set the direction for ectopic endodermal cell migration? To investigate this hypothesis, we used morpholinos against *ndr1* and *ndr2* to knock down endogenous Nodal ligands (**Fig. 1C**). We then injected these MOs into host embryos and found that acvr1ab*-expressing donor cells maintain their ability to ingress into the inner layer of the embryo after transplantation (Fig. 1D-E). These data suggest that the endogenous Nodal is not necessary to trigger the sorting behavior. Conversely, we saturated the endogenous Nodal gradient by overexpressing *ndr1* and *ndr2* ligands in the host embryo (Fig. 1F-G). As before, the acvr1ba*-induced endodermal cells retained their ability to ingress into the inner layer (Fig. 1H-I). Together, these data suggest that although autocrine Nodal reception is essential for initiating internalization, neither the endogenous endodermal cells nor the endogenous Nodal ligands spatially direct ingression.

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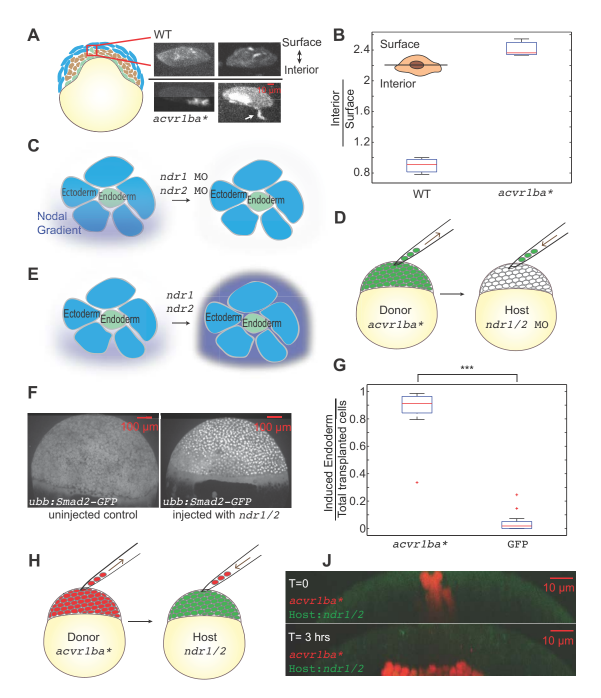


Figure 1: Nodal ligands initiate but do not guide the ingression of endodermal cells.

(A) Actin localization in ectopic endodermal cells. Blue, ectoderm; brown, mesoderm; green, endoderm Donor embryos were injected with GFP-UTRN mRNA to label actin filaments. Cells overexpressing *acvr1ba** or control cells expressing GFP-UTRN only were transplanted to the animal poles of wild-type host embryos. Actin was enriched on the interior side of *acvr1ba**-expressing cells while control cells exhibited uniform actin distribution. Data is resliced and projected to the XZ plane, with the surface of the embryo towards the top and the interior towards the bottom. Arrow shows interior-facing protrusion.

- **(B)** Boxplot of the ratio of interior to surface accumulation of actin. *acvr1ba**-expressing cells exhibited significant interior enrichment of actin compared to control cells. Data is shown as mean ± SEM of 3 independent transplantation experiments, with 58 cells per condition. Student's t-test was performed. * p<0.05, ** p<0.01, *** p<0.001.
- (C-F) Testing whether the endogenous Nodal is required as a directional cue for endoderm ingression through knockdown of endogenous Nodal ligands. (C) *ndr1* and *ndr2* MOs were injected into host embryos to remove the endogenous Nodal gradient. (E) Cells expressing *acvr1ba** were transplanted to the animal pole of a Nodal-depleted host. (F) Boxplot quantification of endoderm contribution at 20 hpf for all transplanted cells. Cells overexpressing *acvr1ba** still contributed to endodermal tissues even in the absence
- Cells overexpressing *acvr1ba** still contributed to endodermal tissues even in the absence of an endogenous Nodal gradient. Data is shown as mean ± SEM of 2 independent transplantation experiments, with 15 embryos per condition. Student's t-test was performed. * p<0.05, ** p<0.01, *** p<0.001.
- (F-I) Saturating the endogenous Nodal gradient to test whether it acts as a directional cue. (G) *ndr1* and *ndr2* mRNAs were injected into host embryos to produce uniform Nodal expression. (H) *Tg(ubb:Smad2-GFP)* shows uniform nuclear translocation in a *ndr1* and *ndr2* injected embryo, suggesting uniform Nodal signaling. (I) Cells expressing *acvr1ba** were transplanted to the animal pole of a Nodal-saturated host. (J) Representative image showing positions of acvr1ba* cells (red) immediately and 3 hours after transplantation in a Nodal-saturated host.

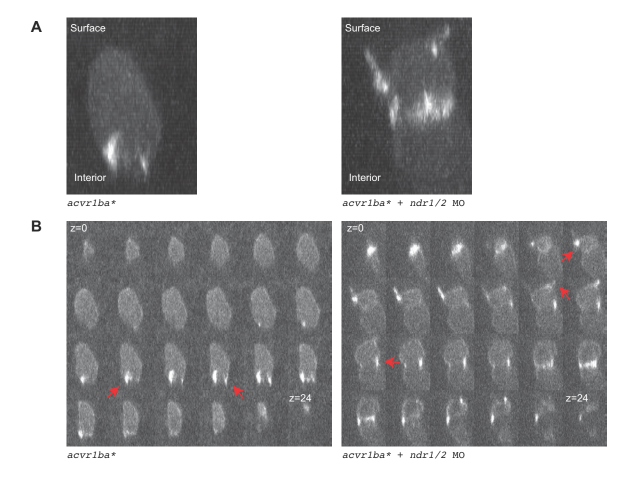


Figure 2: Blocking autocrine production of *ndr1/2* interferes with polarity of actin-based protrusions in *acvr1ba** cells.

- (A) Z projection of individual transplanted cells injected with either acvr1ba* alone or acvr1ba* with ndr1/2 MOs.
- **(B)** Montage of Z stack of cells shown in (A). Red arrows indicate actin enrichment.

CHAPTER SEVEN: SUMMARY AND DISCUSSION

In this work, we investigated the molecular signals that initiate the movement of endodermal cells from the surface to the interior of the embryo during zebrafish development. By leveraging the ability of ectopically-induced endodermal cells to sort to the endogenous endodermal domain, we dissected the molecular logic of sorting without the confounding influences of fate specification and global morphogenetic movements at the margin. Our work shows that an autocrine circuit of Nodal activated by acvr1ba* is both necessary and sufficient to trigger internalization of endodermal cells (**Fig. 1H** from Chapter Three, **1D** from Chapter Four, **1E** from Chapter Five). Neither the endogenous Nodal gradient nor endogenous endodermal cells are required to direct the sorting process. Our work defines an "AND" gate consisting of *sox32*-dependent endodermal specification and Nodal ligand reception that initiates the internalization process (**Fig. 1A** from Chapter Four).

Nodal ligands specify both endodermal cell fate and endodermal sorting

Most of the focus on Nodal signaling during endoderm development has centered around its role in fate specification (Aoki, David, et al. 2002; Hagos and Dougan 2007; Dubrulle et al. 2015). Here we identified an additional role for Nodal as a signaling molecule that regulates endodermal sorting. Endodermal cells that either lack the ability to generate Nodal ligands or to receive Nodal ligands fail to undergo internalization when transplanted to the animal pole (**Fig. 1H** from Chapter Three, **1E** from Chapter Five). Normally both endoderm specification and Nodal ligand reception occur in the same location in the embryo near the margin, and activation of this "AND" gate (endodermal specification +

Nodal ligand reception) could help specify when and where the internalization process occurs. The requirement for both Nodal ligands and endodermal specification could prevent non-endodermal cells that transiently receive Nodal ligands from internalizing. The autocrine nature of this circuit could help control the timing of internalization, which could be triggered when differentiation has proceeded sufficiently to drive this positive feedback loop. This positive feedback loop in which cells that receive Nodal ligand input release more Nodal ligand has previously been implicated in the large-scale self-organization of the Nodal field (Chan et al. 2009), and our work demonstrates an additional role for this feedback loop in coordinating endodermal cell sorting. This circuit could also enable the multicellular coordination of internalization. In chick embryos, single cell ingression can be amplified to induce more of the epiblast to undergo ingression (Voiculescu et al. 2014). Such community effect is Nodal-dependent and underlies the formation of primitive streak.

Nodal ligands are received through the Nodal receptor Acvr1ba and its co-receptor Tdgf1 (the zebrafish homolog of TDGF1/Cripto). The constitutively active Nodal receptor Acvr1ba* has frequently been used to investigate the Nodal signaling pathway (Schier and Shen 2000; Gritsman et al. 1999; Rachel M. Warga and Kane 2003). We find that acvr1ba* requires additional autocrine production and reception of Nodal ligands to support endodermal sorting (**Fig. 1** from Chapter Five). Why might this be? One possibility is that internalization is only triggered above a certain threshold of Nodal signaling. For wild-type cells, this signaling threshold might only be reached at the margin, where Nodal expression is highest, whereas cells expressing acvr1ba*, in which Nodal

signaling is activated beyond wild-type levels, can reach the thresholds needed for internalization even at positions far from the margin. However, in either case (wild-type or acvr1ba*), these high signaling levels are achieved by a Nodal-induced Nodal expression positive feedback loop. In acvr1ba*-induced endodermal cell, Smad2 activation levels are comparable with and without ndr1/2 knockdown (Fig. 1J-K from Chapter Three), suggesting acvr1ba* can initiate Nodal signaling independent of Nodal ligands. In future experiments, this model could be tested by perturbing Smad function to varying degrees in presence of acvr1ba* and assessing the effects on ingression ability. An alternative explanation for the role of a Nodal autocrine circuit could be activation of tdgf1, which may have signaling roles independent of acvr1ba. This model would be consistent with previous literature showing that acvr1ba* can only partially rescue tdgf1 loss-of-function (Rachel M. Warga and Kane 2003). In future work, it will be interesting to further examine the differential signaling engaged by acvr1ba* in the absence and presence of *tdgf1* to identify the *tdgf1*-specific effectors that could participate in endoderm migration.

Directional cues are not limited to Nodal ligands

We dissected the role of Nodal as a trigger for endodermal cell internalization. Through experiments with MZ *tdgf1* as a background for donor and host, we found that ectopic endodermal cells trigger sorting in an autocrine manner. By labeling the actin dynamics, we observed basal enrichment of actin-based protrusions, consistent with other reports suggesting that endodermal cells internalize through active migration(Giger and David 2017). Previous work in hydra also demonstrated the ability of individual endodermal cells

to migrate towards the center of ectodermal aggregates, suggesting that invasion of endodermal cells into ectoderm may represent an ancient morphogenetic behavior (Takaku, Hariyama, and Fujisawa 2005).

But which spatial cues are these cells reading to migrate towards the interior of the embryo? We ruled out endogenous endodermal cells as an attractive positional cue because acvr1ba*-expressing cells can ingress in the MZ tdgf1 background, which lacks endogenous endodermal cells. Moreover, a functioning Nodal gradient does not exist in the MZ tdgf1 host embryos, suggesting it is not providing positional cue either. Because neither knocking down the endogenous Nodal gradient nor flooding the embryo with uniform Nodal gradient block involution, our experiments further ruled out the endogenous Nodal gradient as essential positional cues for ingression. The intersection of endodermal specification and Nodal ligand reception could unlock the ability of these cells to read other extracellular cues that are polarized from the outside to the inside of the embryo such as soluble ligands, ECM components, and mechanical cues (Piccolo 2013; Brunet et al. 2013). Apela (also known as Toddler and Elabela) functions as a motogen and enhances the movement of mesodermal and endodermal cells through Apelin receptor signaling, and Nodal is known to activate Apelin receptor expression (Pauli et al. 2014). However, we found that morpholino-directed knockdown of Apelin receptor a and b did not affect the ability of acvr1ba*-induced cells to ingress into the interior of the embryo (Fig. 1), suggesting that Apela is unlikely to be the spatial cue. Alternatively, the cells could be responding to intrinsic polarity cues such as an oriented apical-basal polarity followed by apical constriction. Consistent with this idea, Xenopus bottle cells

and *C.elegans* endodermal progenitor cells have apical-basal polarity and activate apical constriction to initiate gastrulation movements (Nance and Priess 2002). Cleary, additional work is needed to resolve this question.

Ingression functions as a pattern-refinement mechanism

This work aims to understand the molecular cues that initiate endodermal internalization and germ layer sorting. In addition to laying the foundation for coordinated cell movement at the primary site of endodermal cell internalization during normal development, single-cell ingression may also function as a backup plan to ensure that endodermal cells that are specified late or otherwise miss involution can still find a path into the inner layer. Given that this sorting behavior is based on an autocrine circuit, endodermal cells can still ingress even if they are no longer adjacent to the margin, and this could increase the precision of the first step of endoderm morphogenesis.

From previous work on zebrafish morphogenesis, it is known that dorsal endodermal cells migrate highly asynchronously, which could lead to challenges in germ layer segregation (Keller et al. 2008). Cell sorting is thought to enable systems with initially noisy fate specification to generate robust final patterns. One extreme example is Dictyostelium, whose initial differentiation decision into prestalk or prespore cell is random, and differential migration is responsible for the final pattern (Dormann, Vasiev, and Weijer 2000). Similarly, during neural tube formation in zebrafish, heterogeneous sonic hedgehog responsivity is sharpened by neural progenitor cells sorting into discrete domains (Xiong et al. 2013). If migration were random, it would be expected to blur the

boundaries between different germ layers for cells responding to a source of morphogen such as Nodal. In contrast, by linking directed migration to cell fate specification and signaling, this movement may instead improve the precision of the overall process. By establishing the necessary and sufficient triggers for endodermal sorting *in vivo*, our approach should be useful for continuing to define the logic of endodermal sorting during zebrafish gastrulation.

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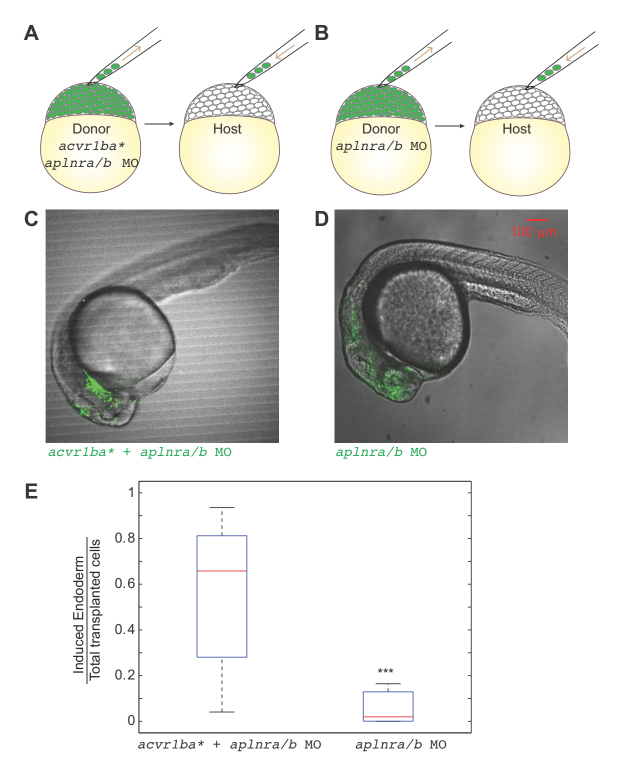


Figure 1:

Apelin receptor signaling is not essential for ectopic endoderm ingression.

(A-B) Schematic diagrams depicting single donor transplant assay to test the

(A-B) Schematic diagrams depicting single donor transplant assay to test the role of apelin receptor signaling. (A) *acvr1ba**-expressing cells with *aplnra* and *aplnrb* MOs were transplanted to the animal pole of a wild-type host embryo. (B) Cells with *aplnra* and *aplnrb* MOs alone were transplanted to the animal pole of a wild-type host embryo.

- **(C-D)** Representative images showing distribution of induced endodermal cells in a wild-type host. Donor cells in (A) (green) mainly localized to endoderm-derived tissue (C), while donor cells in (B) mainly localized to ectoderm-derived tissue (D). Lateral view, anterior to the right.
- **(E)** Boxplot quantification of endoderm contribution at 21 hpf of transplanted cells depicted in (A-B). *acvr1ba**-expressing cells with *aplnra* and *aplnrb* MOs contributed to endoderm significantly more than cells with *aplnra* and *aplnrb* MOs alone. Data is shown as mean ± SEM of 3 independent transplantation experiments with 18 embryos per condition. Student's t-test was performed. *** p<0.001.

CHAPTER EIGHT: ZEBRAFISH TECHNIQUES AND METHODS

Zebrafish strains and embryos maintenance

Zebrafish maintenance was carried out under standard laboratory conditions in the

zebrafish facility at Smith Cardiovascular Research Institute. Embryos were grown at 28-

31°C in egg water and staged as described previously (Kimmel et al. 1995). The following

wild-type (WT), mutant and transgenic lines were used: (WT) AB/TL; (mutant) tdgf1tz57/+

(a generous gift from Lilianna Solnica-Krezel lab in Washington University in St. Louis);

(transgenic) $Tg(sox17:GFP)^{s870}$, $Tg(sox17:DsRed)^{s903}Tg(h2afva:h2afva-mCherry)^{tud7}$,

Tg(ubb: GFP-Smad2) sfc16. Tg(sox17:GFP)s870 and Tg(sox17:DsRed)s903 have been

previously described (Chung and Stainier 2008; Mizoguchi et al. 2008).

Tg(h2afva:h2afva-mCherry)^{tud7} has been previously described (Knopf et al. 2011). To

construct $Tg(ubb:GFP-Smad2)^{sfc16}$, transgene plasmid mTol2-ubiq:GFP-Smad2 was

created by separate PCR amplification of the ubiquitin promoter and GFP ORF and then

cloned into pmTol2-ef1a:Venus-Smad2 (gift from Steve Harvey) cut with EcoRV and Agel

to remove the ef1a promoter and Venus ORF. $Tg(ubb:GFP-Smad2)^{sfc16}$ was created by

injecting 20 pg of the transgene plasmid DNA along with 100 pg of Tol2 transposase

mRNA at the one cell stage. Injected embryos were then sorted by fluorescence on d0,

raised to adulthood, and then screened for founders by outcrossing to wild-type.

Genotype analysis

Maternal zygotic tdgf1 mutants.

57

To create maternal zygotic *tdgf1* mutants, *tdgf1*^{tz57/+} parents were in-crossed, and all embryos were injected with *tdgf1* mRNA so that homozygous embryos could survive. Genotyping was performed according to established protocols (Hashimoto et al. 2000 and Pogoda et al. 2000).

RNA expression constructs and morpholinos

Capped messenger RNA was synthesized using the mMESSAGE mMACHINE kit (Ambion). The following expression plasmids were used in this study: acvr1ba* in pCS2 (pCS2-acvr1ba*-tBFP), full-length zebrafish sox32 in pCS2 (pCS2-sox32; Chung and Stainier, 2008), ndr1 and ndr2 in pCS2 independently and GFP-UTRN in pCS2. The sox32 MO was designed to target the translation initiation site and was used at 2 ng (5'-CCTCCTCAGTGTTTATTTCGCTCAT-3'). ndr1 MO (5'-ATGTCAAATCAAGGTAATAATCCAC-3') and ndr2 MO (5'-GCGACTCCCGAGCGTGTGCATGATG-3') were used at 4ng. MOs targeting aplnra (5'-CGGTGTATTCCGGCGTTGGCTCCAT-3') aplnrb (5'and CAGAGAAGTTGTTTGTCATGTGCTC-3') were injected at the one-cell stage at 1 ng or 0.5 ng, respectively.

mRNA, morpholino and dye injection

mRNA, morpholino and dye injections were performed with a micromanipulator connected to Picospritzer III. Drop size was regulated by the duration and pressure of the pulse. mRNA of appropriate concentration for different genes was injected into the yolk of the embryo at 1-cell stage. To obtain induced endodermal cells, 0.5pg *acvr1ba** mRNA

or 100pg sox32 mRNA were injected into the embryo. To study the effect of Nodal ligands on ingression, 4pg *ndr1* and *ndr2* mRNA were injected into the embryo. 200 pg GFP-UTRN was injected into the embryo to visualize actin dynamics. Morpholinos were briefly incubated at 65 °C to prevent precipitation and then injected into the yolk before the first cell division. 4ng *ndr1*, *ndr2* MO (Feldman and Stemple 2001; Karlen and Rebagliati 2001) or 2ng sox32 MO (Sakaguchi, Kuroiwa, and Takeda 2001) or 1ng *aplnra* MO, 0.5ng *aplnrb* MO (Scott et al. 2007; Paskaradevan and Scott 2012; Pauli et al. 2014)was injected into the embryo to inhibit the translation of the corresponding genes. Dyes including Dextran-FITC or Dextran-tetra-methyl-rhodamin-dextran (TMR-dextran) or Dextran-Alexa Fluor 680 (LifeTechnologies) were injected at 1ng at the one-cell stage to label whole cells while 1 ng Histone H1 - Alexa Fluor 488 conjugate was injected into the embryo at 1-cell stage to label the nucleus.

Real-time quantitative PCR

For Nodal-activated conditions, wild-type embryos were injected at the one-cell stage with 2 pg *acvr1ba** mRNA or *mCherry* mRNA as a control. Expression of *sox17*, a known Nodal target gene, as well as *sox32*, were used to confirm Nodal activation. Expression of *cdh2* was measured under different Nodal-activated conditions. At shield stage, total RNA was extracted using the RNAqueous-Micro Kit, and 1 ng was used for reverse transcription with the SuperScript VILO cDNA Synthesis Kit (Invitrogen). The quantitative PCR reaction mixture contained 2 µl of 10-fold diluted cDNA, 12.5 µl SYBR green PCR master mix (Applied Biosystems), 714 nM of each primer, and nuclease-free water for a total volume of 25 µL in 48-well plates (Ilumina). Reactions were performed in the Eco

Real-Time PCR System (Illumina, Inc.) as follows: initial activation at 95°C for 10 min followed by 40 cycles of 30 s at 95°C, 30 s at 60°C, and 30 s at 68°C. Once the PCR was completed, a melt-curve analysis was performed to determine reaction specificity. Samples were run in triplicate. The housekeeping gene *ef1a* was used as a reference. Refer to Tabel S1. List of Oligonucleotides for the primers used in this study.

Transplantation

Donor and host embryos were dechorionated with forceps under a dissection stereomicroscope and transferred into a glass plate with 0.3x Ringer's Buffer. 25~50 cells were taken from a dechorionated donor embryo(s) at sphere stage (4 hpf) and transplanted into the animal pole of a dechorionated host at the same stage using a bevelled borosilicate needle with a 35 µm inner diameter attached to a syringe system. In single donor transplantation experiments, the donor embryo was injected with mRNAs and/or morpholinos described in the main text and wild-type or MZ *tdgf1* embryos were used as hosts. In double donor transplantation experiments, the endoderm donor embryo was injected with 2pg *acvr1ba** mRNA, control ectoderm donor embryo was injected with 2ng *sox32* morpholino, and wild-type embryos were used as the host. Dextran dyes were used to differentiate donor versus host cells. H1 - Alexa Fluor 488 conjugate was used label the nucleus for single cell tracking. After transplantation, embryos were either immediately mounted for microscopy or maintained in 0.3x Ringer's Buffer at 28-31°C for further analysis.

Nodal inhibitor SB505124 treatment

For drug treatment, embryos from one dish were removed at the desired stage and split into glass dishes containing the drug in 5 ml embryo medium, at a density of 25 embryos/dish. For SB505124, the lowest dose that produced the sqt; cyc phenotype ranged from 30–50 µM, depending on the age of the drug (Hagos and Dougan 2007). Desired concentration is diluted from 10mM stock. For transplants, drug treatment is initiated after the transplantation is finished at 4hpf.

Time-lapse Confocal Microscopy

Dechorionated embryos, immediately after transplantation, were embedded in 1% low-melting agarose within glass-bottom Petri dishes, with animal pole mounted towards the glass bottom. For tracking, transplanted embryos were imaged with a 20x/0.75 NA Plan Fluor multi-immersion objective with water as the immersion media. For actin dynamics visualization, a 40x/1.15 NA water immersion objective was used. 10x/0.45 NA Plan Apo λ objective was used for imaging 24 hpf or 18-somite stage embryos. A high-speed widefield Nikon spinning disk confocal microscope was used for all imaging. This microscope is equipped with an Andor Borealis CSU-W1 unit, an Andor DU-888 EMCCD camera, and a stage-top incubator unit from OkoLab. Andor 4-line laser launch (100 mW at 405, 561, and 640 nm; 150 mW at 488 nm) was used for excitation. Micro-Manager Open Source Microscopy Software Version 2.0 Beta was used to control the microscope. Image stacks of 70-150 μ m with 1-2 μ m (1 μ m for timelapse and 2 μ m for end-point scanning) z stack were recorded in continuous mode, resulting in an image sampling rate of 2-4 min. Embryos were kept at 28.5°C throughout imaging.

Image Processing

Tracking with Gaussian Mixture Models (TGMM) software for automated large-scale segmentation and tracking of fluorescently labeled cell nuclei from the Keller Lab was adapted for single cell tracking of the transplanted cells (Amat et al. 2014). Timelapse datasets with Z-stacks were rendered into 3D tracks and filtered by track length. A sphere was used for modeling the zebrafish embryo, and Cartesian coordinates were transformed into spherical coordinates to determine the radial distance travelled by the transplanted cells.

Quantification and Statistical Analysis

Quantification of percentage of transplanted cell that localized to endodermal vs. non-endodermal domains was performed by analyzing images with Fiji. Z-stack images were converted to maximum intensity projections and thresholded by Renyi entropy. Particles were analyzed with Fiji and size of ROIs were measured. For image reslicing, Z-stack images were resliced to achieve 1 x 1 x 1 voxel size, then converted to maximum intensity projections to generate an XZ projection. Statistical data analysis was performed using Student's t-test in Matlab.

Table 1. List of Oligonucleotides

Oligonucleotide Name	Sequence
ef1a_forward	5'-CAAGAAGAGTAGTACCGCTAGCAT-3'
ef1a_reverse	5'-CACGGTGACAACATGCTGGAG-3'
sox17_forward	5'-CACAATGCGGAGCTGAGTAA-3'

sox17_reverse	5'-GCCTCCTCAACGAATGGAC-3'
sox32_forward	5'-CGGACCTGGAGAACACTGAC-3'
sox32_reverse	5'-GCATGTACGGACGCTTATCTG-3'
cdh2_forward	5'-CATCCCGGAGACATAGGAGA-3'
cdh2_reverse	5'-GCCCTCGTAGTCAAACACCA-3'
Oep5	5'-GAGATGGAGATGTTCTAATG-3'
Oep3m	5'-GAACAGTTGACTCGTCAC-3'
Oep3w	5'-GAACAGTTGACTCGTCAT-3'
Sox32 MO	5'-GCATCCGGTCGACATACATGCTGTT-3'
Sqt MO	5'-ATGTCAAATCAAGGTAATAATCCAC-3'
Cyc MO	5'-GCGACTCCCGAGCGTGTGCATGATG-3'
Aplnr a MO	5'-CGGTGTATTCCGGCGTTGGCTCCAT-3'
Aplnr b MO	5'-CAGAGAAGTTGTTTGTCATGTGCTC-3'
Control MO	5'-CCTCTTAACCTCAGTTACAATTTATA-3'

Table 2. Key Resource Table

Reagent or Resource	Source	ldentifier	
Chemicals, Peptides, and Recombinant Proteins			
Dextran, Alexa Fluor™	Invitrogen	Cat#D22914	
647			

Dextran,	Invitrogen	Cat#D1868
Tetramethylrhodamine		
Dextran, Fluorescein	Invitrogen	Cat#D1821
Dextran, Alexa Fluor™	Invitrogen	Cat#D34680
680		
Histone H1 From Calf	Invitrogen	Cat#H13188
Thymus, Alexa Fluor™		
488 Conjugate		
Critical Commercial Assays		
mMESSAGE mMACHINE	Ambion	Cat#AM1340
SP6 Transcription Kit		
SuperScript VILO cDNA	Invitrogen	Cat#11754050
Synthesis Kit		
SYBR green PCR master	Applied	Cat#4309155
mix	Biosciences	
Experimental Models: Organisms/Strains		
Zebrafish: AB/TL	This study	ZFIN: ZDB-GENO-960809-7
Zebrafish: EKW	This study	ZFIN: ZDB-GENO-031202-1

Zebrafish: Tg(sox17:GFP)	This study	ZFIN: ZDB-GENO-061228-1
Zebrafish:	This study	ZFIN: ZDB-GENO-080812-1
Tg(sox17:DsRed)		
Zebrafish:	This study	ZFIN: ZDB-GENO-100923-1
Tg(h2afva:h2afva-		
mCherry)		
Zebrafish: Tg(ubb:GFP-	This study	N/A
Smad2)		
Zebrafish: tdgf1 ^{tz57/+}	Lilianna	ZFIN: ZDB-GENO-080708-1
	Solnica-	
	Krezel lab	
Zebrafish: tdgf1tz57/tz57	This study	ZFIN: ZDB-GENO-980202-989
Oligonucleotides		
List of oligonucleotides	See Table	N/A
	S1	
Recombinant DNA	1	
pCS2-acbr1ba*	This study	N/A
pCS2-acbr1ba*-p2a-tBFP	This study	N/A

pCS2-sox32	This study	N/A	
pCS2-sox32-p2a-tBFP	This study	N/A	
pCS2-ndr1	This study	N/A	
pCS2-ndr1-GFP	This study	N/A	
pCS2-ndr2	This study	N/A	
pCS2-ndr2-tBFP	This study	N/A	
pCS2-GFP-UTRN	This study	N/A	
pCS2-GFP	This study	N/A	
pCS2-h2a-mCherry	This study	N/A	
pCS2-tdgf1	This study	N/A	
pmTol2-ef1a:Venus-	Steve Harvey	N/A	
Smad2			
Software and Algorithms			
Fiji	NIH	https://fiji.sc	
Matlab2013a	MathWorks	http://mathworks.com	
	Inc.		

ТӨММ	Philipp Keller	https://www.janelia.org/lab/keller-
	lab	lab/software/fast-accurate-reconstruction-
		cell-lineages-large-scale-fluorescence

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