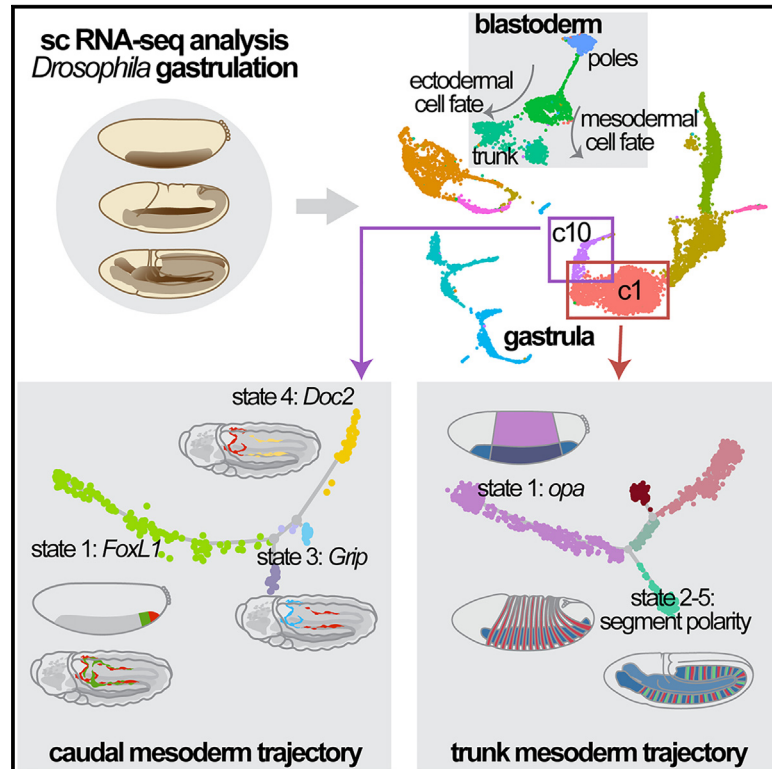


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Single-cell transcriptomics illuminates regulatory steps driving anterior-posterior patterning of *Drosophila* embryonic mesoderm

Graphical abstract



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In brief

Sun et al. used single-cell RNA-seq to investigate how overall gene expression progresses to accommodate cellular changes and coordinate the morphogenetic movements associated with *Drosophila* gastrulation. Identifying regulatory steps involved in patterning the mesoderm, this study provides insights into the mechanisms regulating EMT and collective cell migration during morphogenesis.

Highlights

- Single-cell transcriptomic analysis of *Drosophila* gastrulation using live cells
- Cells spanning 90 min of development were profiled to illuminate temporal transitions
- Spatial information captured includes differences in trunk vs. poles and germ layers
- Mesoderm patterning along the anterior-posterior axis and progenitors uncovered



Resource

Single-cell transcriptomics illuminates regulatory steps driving anterior-posterior patterning of *Drosophila* embryonic mesoderm

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SUMMARY

Single-cell technologies promise to uncover how transcriptional programs orchestrate complex processes during embryogenesis. Here, we apply a combination of single-cell technology and genetic analysis to investigate the dynamic transcriptional changes associated with *Drosophila* embryo morphogenesis at gastrulation. Our dataset encompassing the blastoderm-to-gastrula transition provides a comprehensive single-cell map of gene expression across cell lineages validated by genetic analysis. Subclustering and trajectory analyses revealed a surprising stepwise progression in patterning to transition zygotic gene expression and specify germ layers as well as uncovered an early role for ecdysone signaling in epithelial-to-mesenchymal transition in the mesoderm. We also show multipotent progenitors arise prior to gastrulation by analyzing the transcription trajectory of caudal mesoderm cells, including a derivative that ultimately incorporates into visceral muscles of the midgut and hindgut. This study provides a rich resource of gastrulation and elucidates spatially regulated temporal transitions of transcription states during the process.

INTRODUCTION

Quantitatively capturing the transcription state of an embryonic cell is essential for uncovering its identity, while tracking the dynamic transcriptional changes within the lineage is imperative for understanding the differentiation trajectory as well as for deciphering the gene regulatory networks associated with cell-fate specification.¹ We here apply single-cell sequencing technology in conjunction with conventional genetic tools and methods to investigate the molecular mechanisms driving the *Drosophila* embryo developmental progression from blastula to gastrula. By focusing on the mesoderm, we uncovered the transcription states that account for progenitor cell specification and the regulation that underlies epithelial-to-mesenchymal transition (EMT)^{2,3} during gastrulation.

Gastrulation constitutes a crucial stage during development when metazoan animals acquire cellular diversity and set up basic body plans.^{4–6} Initialized with totipotency, embryonic cells are specialized, forming three germ layers, rearranged through complex morphogenetic movements, and eventually shaped into a body architecture with various rudimentary organs. Transcription changes very rapidly prior to gastrulation during early embryogenesis. The initial 13 rounds of nuclear division in early embryogenesis take place in a syncytial embryo without cytokinesis, and all gap phases (i.e., G1 and G2) are bypassed.⁷ Maternal factors are degraded, and zygotic transcription takes

over gradually through two waves of zygotic gene activation (ZGA).^{8–10} The major wave of gene expression associated with ZGA coincides with the 14th nuclear cycle (nc14), during which time cell membranes grow between the blastoderm nuclei and cellularization takes place.^{11–13} Spatially localized gene transcription in the ventral domain (e.g., the expression of *snail* [*sna*] and *twist* [*twi*]) associated with the presumptive mesoderm at mid-nc14 is the first sign of lineage diversification during embryogenesis. By the end of nc14, mesoderm cells are specified downstream of Toll signaling at the ventral blastoderm, and invagination of those cells leads to mesoderm formation. Developmental time is the major variable that distinguishes gene expression programs of cell populations prior to the emergence of germ layers. However, once cell movements begin during gastrulation, it is generally thought that gene expression programs do not exhibit widespread changes until the cells reach their destination and initiate differentiation programs. Whether this is indeed the case remains unknown, and the interplay of transcriptional programs guiding these complex cellular transitions is also poorly explored. We hypothesized that the single-cell approach might be able to capture the dynamic transcriptional changes in migrating cells as well as elucidate the gene regulatory landscape during this important transition phase.

Gastrulation starts at stage (st) 6, following the accomplishment of dorsal-ventral (DV) patterning at st 5, as *sna* and *twi* genes function together to control the multifaceted cellular



changes that initiate and drive presumptive mesoderm internalization, ultimately resulting in a tube-like structure forming at the ventral side of the gastrula at st 7.^{14–16} Subsequently, at st 8, the tube disassembles as mesoderm cells in the trunk lose apico-basal polarity, detach from each other, enter into the first mitotic cycle after nc14, and through an EMT acquire a migratory ability supporting their movement in the dorsal direction.^{17,18} Concomitantly, these cells move in the posterior direction together with the ectoderm during germband extension (GBE) to double the length of the embryonic trunk along the anterior-posterior (AP) axis.⁵ The coupling between EMT and GBE, two pronounced morphogenetic cellular movements at gastrulation, signifies the cross-regulation between the DV- and AP-axis patterning programs, respectively. Single-cell technology provides us with a unique opportunity to investigate the genetic links connecting EMT with GBE¹⁹ with high spatiotemporal resolution by tracking gene expression trends in trunk mesoderm cells.

Several developmental timer genes, or timing factors, have been shown to control embryonic AP patterning at gastrulation,^{20,21} but their role in supporting patterning and specification of progenitor cells within the mesoderm has not been characterized. These include *odd paired* (*opa*), which is broadly expressed in the trunk region at nc14 and acts as a pioneer factor to modulate chromatin accessibility and thereby regulate the spatiotemporal dynamics of zygotic gene expression.^{22,23} Among *Opa* targets are the pair-rule and segment polarity genes,^{20,22,24} which function in a cascade of segmentation genes to cooperatively pattern the *Drosophila* embryo trunk along the AP axis. The posterior pole, or “tail,” of the embryo is patterned after the trunk under the regulation of the terminal system by the gene *tailless* (*tl*). This process requires *opa* and at least two additional timing factors, *caudal* (*cad*) and *Dichaete* (*D*).^{21,25} The caudal mesoderm contains multipotent progenitors for muscle cells and Malpighian tubules (MTs), the *Drosophila* kidney equivalent.^{26–28} In contrast to mesoderm cells in the trunk (which adopt different cell fates in response to inductive signals originating in the ectoderm), caudal mesoderm cells are migratory precursor cells that remain undifferentiated until they are incorporated into organs. While much focus has been on AP patterning of the ectoderm, the mechanism for subdivisions within the mesoderm remains unclear, including how the posterior founder populations and segmentation gene expression are regulated. Gene expression information at a single-cell resolution also holds the potential to uncover the transcription signatures within a trajectory of cell states that lead to the lineage commitment of posteriorly located progenitor cells.

RESULTS

Single-cell transcriptomic profiling and lineage identification of *Drosophila* embryos during gastrulation

To investigate transcriptional changes during gastrulation, we collected embryos from 3 to 5 hours after egg laying, representing st 5–10, and processed them for single-cell RNA sequencing (scRNA-seq) analysis (Figure 1A; see STAR Methods). The uniform manifold approximation and projection (UMAP) dimension reduction with 20 principal components (PCs) resulted in 12 unique cell clusters, c1–c12 (numbered based on cell population

size ranking of the cluster, Figures 1B and S1B), which can be identified by the associated marker genes (Figure 1C and Table S1). Cells sampled using this approach covered the expected developmental stages (blastoderm and gastrula) and cell lineages, including mesoderm, endoderm and ectoderm anlage, primordia, and a sampling of differentiating cell types. Gene Ontology (GO) enrichment analysis of marker genes associated with each cluster supports the view that cells found in clusters have specialized functions (Figure S1A).

Aiming to investigate the transcription programs specifically within the mesoderm, we used a lineage tracer consisting of *UAS-mCD8GFP*²⁹ reporter and *twi(2xPE)-GAL4*³⁰ driver to follow *GFP* reporter gene expression in the embryo starting at st 5 and compared it with the expression of gene *twi* in mesoderm lineages (Figure 1D). Levels of *twi* expression are highest in undifferentiated mesoderm c1 and c10 (Figure 1H). *mCD8GFP* is also enriched in c1 and c10; however, it is more strongly expressed in c3 and c8, representing more differentiated trunk mesoderm and specialized head mesoderm cell lineages (Figure 1H). These results further confirm the identities of the mesoderm clusters (i.e., *twi>mCD8GFP* largely follows the transcriptional activity of endogenous gene *twi*) but with a delay imposed by indirect expression through use of *GAL4/UAS* (i.e., c3 is older than c1).

Our data provide a comprehensive single-cell map to track gene expression of different cell lineages during gastrulation. For example, the head mesoderm cluster (c8) marked by *glial cells missing* (*gcm*), a zinc-finger transcription factor required for plasmatocyte differentiation, is well separated from the rest of the mesoderm lineages (i.e., c1, 3, 4, 10, 12) (Figure 1E compared with 1D, 1H). This is consistent with its head-specific expression as shown by *in situ* hybridization from st5 to st8 (Figure 1F) as well as its expression pattern predicted by the *Drosophila* Virtual Expression eXplorer (DVEX), an online tool published with a previous single-cell transcriptomic study of embryos at st 6³¹ (Figure S1C). Our analyses also demonstrate a high degree of cell-type coverage within the embryo (i.e., median ~15,000 unique molecular identifiers [UMIs] per cell for our wild-type #1 [WT#1] sample), and deeper sequencing resulted in a single-cell transcriptome dataset containing high-information content that will illuminate even low-abundance transcripts. For example, with fewer than 10,000 cells covering multiple developmental stages, we were able to detect rare cell types of mesoderm origin such as crystal cells, which constitute only ~5% of the population³² in comparison with *gcm*-marked plasmatocytes, which represent 90%–95% of the *Drosophila* blood cells. Despite their low abundance, we can identify the crystal cell marker *Prophenoloxidase 1* (*PPO1*) within c8 of our dataset (Figures 1G and 1H).

The marker genes were identified using Seurat differential expression tests for this single-cell transcriptome dataset (Figure 1C; Table S1) and were used to assign an identity to each of the 12 cell clusters based on previous studies and gene annotations at the Berkeley *Drosophila* Genome Project (BDGP) database and FlyBase (see STAR Methods). This dataset (live WT#1) is the focus of our study. In repeat experiments (live WT#2 and fixed WT#3), also with integrated dataset, cell clusters with similar lineage identities (Figures S2D and S2F) were recovered (see STAR Methods). As *Drosophila* single cells were isolated from a staged

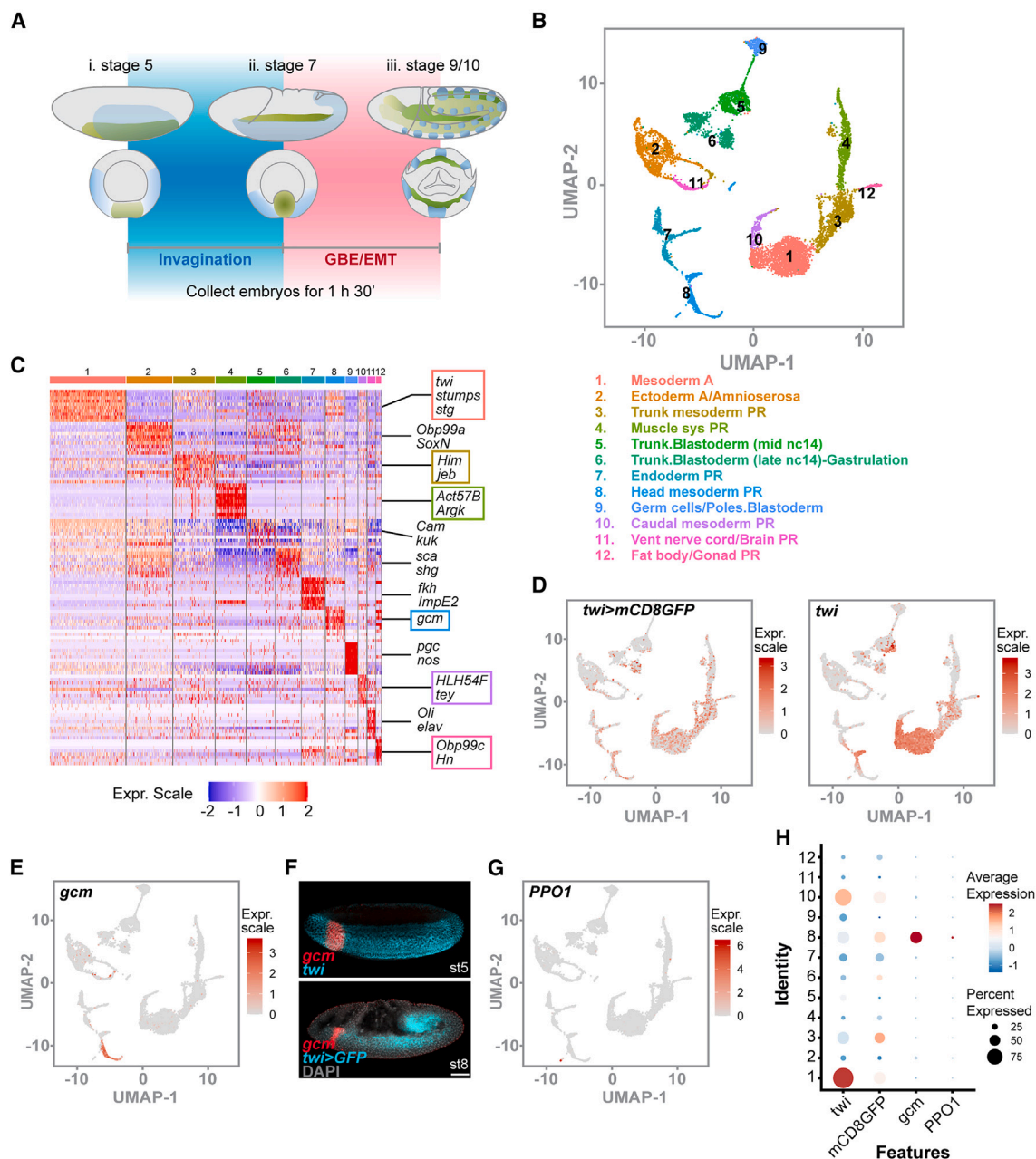


Figure 1. scRNA-seq analysis of *Drosophila* embryos at gastrulation highlights distinct mesoderm cell lineages

(A) Diagram illustrates timing of embryo collection and expected stages at which cells were isolated (see [STAR Methods](#)).

(B) The UMAP dimensionality reduction analysis revealed 12 cell clusters that were annotated based on their associated marker genes. Annotation follows Berkeley Drosophila Genome Project (BDGP) database guidelines: A, anlage (i.e., rudimentary part/organ); PR, primordium (i.e., organ in earliest recognizable state).

(C) Heatmap of the top 10 marker genes expressed in each cluster with examples labeled on the right. Mesoderm lineage markers are indicated within the boxes (see also [Table S1](#)).

(D) Expression of *GFP*, from *twi(2xPE)GAL4*-driven UAS-*mCD8GFP*, and *twi* genes. Red color intensity showing the expression levels or the number of reads at a global scale (normalized by the entire cell population).

(E–G) Detection of head mesoderm cell lineages. UMAP plot and *in situ* hybridization (red, F) of the head mesoderm marker *gcm*. *twi* (cyan, top) and *twi>GFP* (cyan, bottom) transcripts were labeled simultaneously with *gcm* at st 6 and 8, respectively, to indicate mesoderm lineage. DAPI (gray) was used for counterstaining. Scale bar, 50 μ m. (G) Presence of *PPO1*-positive crystal cells in the head mesoderm cluster.

(H) Dotplot of *twi*, *mCD8GFP* driven by *twi(2xPE)GAL4*, *gcm*, and *PPO1* expression in all clusters. Expression levels and percentage of cells in the cluster (y axis) that express indicated genes (x axis) are shown. See also [Figure S1](#) and [Table S1](#).

embryo collection allowing approximately 90-min difference in age, we expect the clusters to segregate based on gene expression programs that differ with respect to both developmental timing as well as spatial registry within the embryo. We therefore examined marker genes using *in situ* hybridization to confirm the categorization (i.e., Figure 1B) and to provide additional insights into the spatiotemporal dynamics in transcription.

To further support data reproducibility, we integrated our dataset (WT#1) with a published transcriptome dataset from stage 12 embryos³³ (Figures S1D and S1E) and compared the mesoderm cell lineages contributed from these two sources. We found that the markers for more differentiated mesoderm cell clusters representing somatic, visceral muscle (s.c2, s.c10), fat body (s.c5), and hemocytes (s.c.12) are highly consistent, whereas the clusters associated with younger mesoderm cells, as expected, predominantly stem from our data focused on gastrulation (s.c4 and s.c17) (see Figure S1F).

Establishing domains and germ layers in the blastoderm embryos

Marker genes were used to map cell clusters to germ layers and domains. To start, we focused on the cells relating to the blastoderm embryo, which encompasses nc14 within st 5. Nc14 is the longest nc, ~60 min in length at 22°C.⁹ During early- to mid-nc14, the embryos exhibit an increase in zygotic transcription while maternal factors continue being degraded, and cellularization is completed by the end of nc14. Previous studies have examined gene expression dynamics at nc14 and characterized expression changes for select marker genes with high temporal resolution.^{9,34} For example, by mid-nc14, maternally deposited transcripts of gene *deadhead* (*dhd*) are depleted, *heartless* (*htl*) and *grainyhead* (*grh*) are switched on, and *sna* and *twi* levels peak, while expression of *Cadherin-N* (*CadN*), activated at early nc14, continues to increase until gastrulation (Figures 2C and 2D).^{9,35,36} Transcripts of early zygotic gene *nullo* and *bottleneck* (*bnk*) both play an essential part in cellularization and are completely eliminated from the embryo by late nc14.^{37,38} Interestingly, *bnk* is only retained in the poles at mid-nc14 (Figure 2C).³⁸ We used this prior knowledge of dynamic gene expression trends to relate clusters c5, c6, and c9 to cells in the blastoderm embryo (Figure 2A). c9 comprises cells that express *dhd*, *bnk*, and *nullo*, likely derived from embryos at mid-nc14 (youngest possible stage based on our collection timing); c6 clearly contains older cells—likely of late nc14 to stage 6—due to higher levels of *CadN* and lack of *dhd*, *bnk*, and *nullo* (compare c9 vs. c6; Figures 2B and S2A). c5 appears to be a transitional state at which all above-mentioned genes are detectable but at low levels (Figure 2B). Therefore, based on their expression profiles, c9 and c5 cells might be at a similar stage and both derived from mid-nc14 embryos, whereas c6 cells are “older” and c5 → c6 represents a temporal progression in transcriptional states from cellularization to the beginning of gastrulation.

In addition, several markers for c6 are spatially localized to regions that give rise to either ectoderm or mesoderm, suggesting that the difference in expression between c9/c5 vs. c6 relates to specification of the three germ layers. *Short gastrulation* (*sog*), a marker for the ectoderm, is expressed in lateral regions

(Figures 2G and 2I); cells expressing *sog* in the UMAP plot are represented on the left side of c6 and are most abundant in c2 that constitutes ectoderm/amnioserosa (Figure 2J *sog*). In contrast, *htl*, like *twi*, is expressed in ventral presumptive mesoderm cells (Figure 2I); *htl*-expressing cells are represented on the right side of c6 and most abundantly present in mesodermal lineages including c1, c3, c8, and c10 (Figure 2J *htl*; see also Figure 1D *twi*). Lastly, *DNasell* is an example of a terminally expressed gene (i.e., anterior and posterior poles; Figure 2I); cells expressing *DNasell* are present in the center of c6 and predominantly in c7, enriched for genes associated with endoderm development (Figure 2J *DNasell*). The distinct localization of ectodermal and mesodermal markers in c6 (e.g., Figure 2J *sog*/ectoderm to the left vs. *htl*/mesoderm to the right of UMAP plots) suggests that germ layers arise at c6 during late nc14. This is also supported by trajectory analysis of the transcriptome of cells comprising c5, c6, and c9, which demonstrates that three transcriptional states emerge: state 1 is associated with maternal and pole cell markers (*dhd*, *pgc*), state 2 with ectodermal markers (*Lac*, *grh*), and state 3 with mesodermal ones (*llp4*, *twi*) (Figures 2E, 2F, and S2B; Table S3.1).

To further validate our conclusions regarding germ-layer specification timing, we subjected the c5 and c6 clusters to subclustering in an unbiased manner and identified five and three subclusters, respectively (Figures S2K and S2L; see STAR Methods). The three subclusters associated with c6 each relate to one of the three germ layers: ectoderm (c6 subcluster1: c6.1, marked by *Lac*, *grh*), mesoderm (c6.2, marked by *CadN*, *htl*), or endoderm (c6.3, marked by *DNasell*, *Pdp1*) (Figures 2K and 2L; see also Figure S2M and Table S3.4). This result confirms germ layers are established in late nc14. In contrast, only one of the five subclusters associated with c5, c5.5, is associated with germ-layer-specific markers corresponding to the presumptive mesoderm (Figures S2J–S2L; Table S3.3), supporting the idea that c5 represents mostly uncommitted younger cells compared with c6, likely from mid-nc14 embryos.

As c9 is associated with marker genes that relate to pole cells (germline) as well as terminal patterning (Figures 2B and S2A; Table S1), we used subclustering to provide additional insights into its identity. This subclustering produced three classes (Figure S2H; Table S3.2). c9.1 expresses high levels of *dhd*, maternal transcripts known to be mostly degraded by mid-nc14 but retained in pole cells^{39,40}; therefore, these are likely pole cells (Figures S2G and S2I). c9.2 is marked by expression of *tailless* (*tll*), *ribbon* (*rib*), and *zerknüllt* (*zen*), all known to be expressed in the anterior and posterior poles at mid-nc14⁴¹ (Figures S2G and S2H). c9.3 appears to be anterior biased as it is enriched for anterior ectoderm marker CG42342 as well as *Psc* and CG45782, both genes known to be expressed in the future head structure at the embryonic anterior (Figures S2G and S2I).^{42–45} The fact that genes expressed at the poles of the embryo (both anterior and posterior; Table S1) are enriched in c9 cells while c5 is marked by *homothorax* (*hth*), only expressed in the trunk blastoderm, with little to no caudal genes expression (e.g., *tll* and *cad*) (Figures S2G–S2L; Table S1), suggests that the distinction between c5 and c9 likely reflects the spatial differences of cells in the embryo (trunk vs. poles) translated into differences in their transcriptional states.

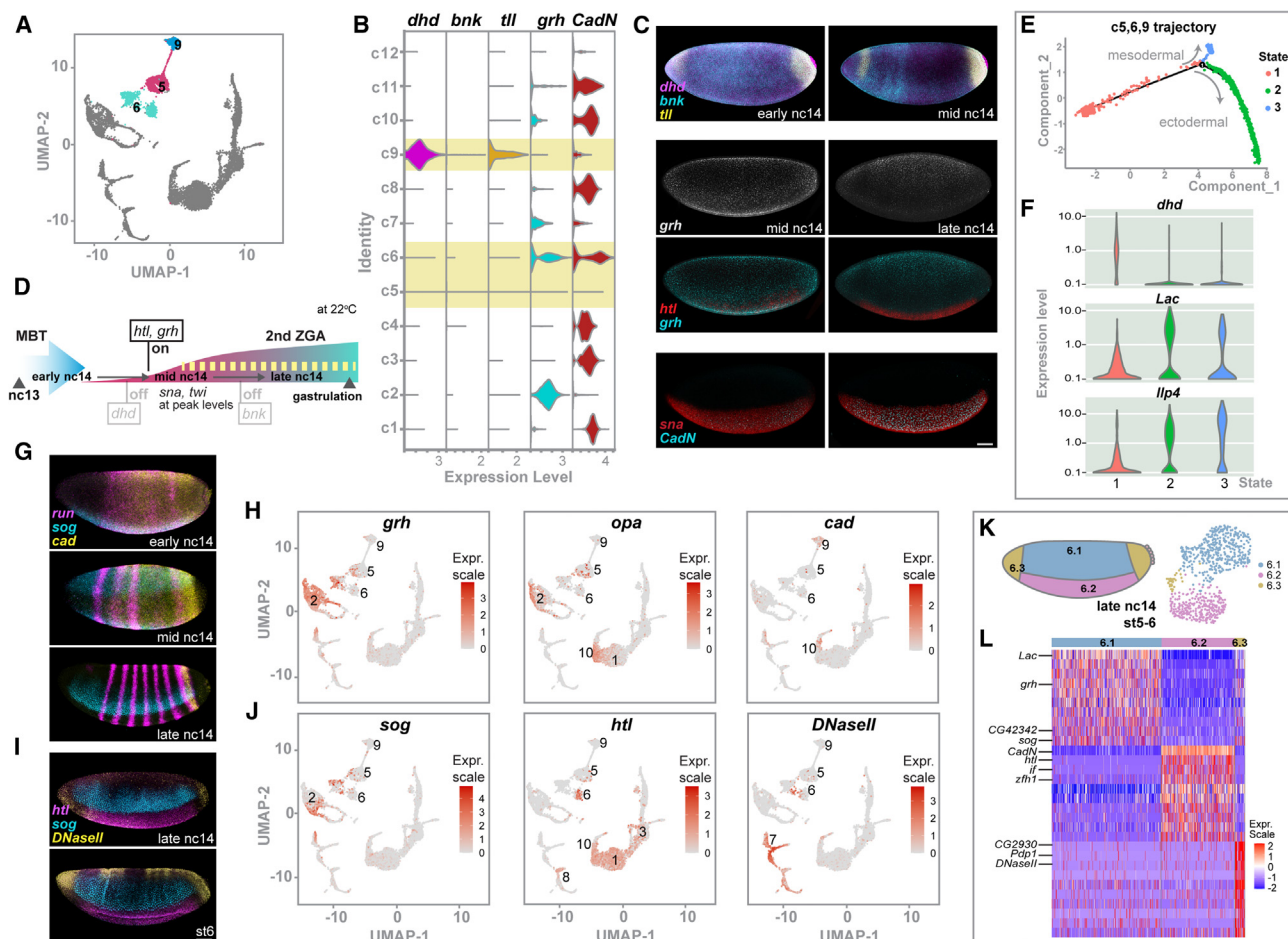


Figure 2. Profiling transcription in the blastoderm at single-cell resolution reveals distinctive transcriptional programs among ectoderm, endoderm, and mesoderm lineages at nc14

(A–D) Blastoderm embryos contribute to three clusters in the UMAP plot: c9, c5, c6 (A). Violin plots (B) and *in situ* hybridization (C) showing representative gene expression in the blastoderm embryos. Schematic representation of transcription changes during nc14 at 22°C; yellow dashed line indicates the expected age of embryos in the collection that by 195 min should be at or older than mid-nc14 (D).

(E and F) Trajectory analysis of cells specifically from c9, c5, and c6 generates 1–3 cell states (E) that are enriched for *dhd*, *Lac*, and *Ilp4*, respectively (F).

(G–J) *In situ* hybridization showing representative marker gene expression in different AP domains or germ layers of the blastoderm embryo during nc14: trunk (*run*) vs. posterior poles (*cad*) (G); ectoderm (*sog*) vs. mesoderm (*htl*) vs. endoderm (*DNaseII*) (I). UMAPs of selected marker genes highlighting timing factors (H) and germ layer-specific genes (J). Red color intensity indicates the relative expression levels at a global scale.

(K and L) UMAP plot (K) and heatmap (L) of c6 cells that are subdivided into three subclusters (c6.1, c6.2, and c6.3). Diagram illustrates a cellularized blastoderm embryo with color-coded germ layers that are labeled with their associated subcluster. Scale bars, 50 μ m. See also Figure S2 and Tables S3–S5.

From our subclustering analysis, we identified additional steps in global gene expression related to activities of timing factors that contribute to embryonic patterning in the blastoderm. These factors, including *opa*, *cad*, and *grh*, function cooperatively to control gene expression at specific developmental time points.^{22,23,25} It is thought that the initial specification of anterior-trunk-posterior programs along the AP axis relates to the localized expression of *opa* (trunk) and *cad* (posterior)^{21,23,25} (Figures 2G and 2H). *grh*,⁴⁶ expressed in both the trunk and poles, is associated with cells in the ectoderm and endoderm anlagen but excluded from mesoderm cells, suggesting a role for it in patterning along the DV axis (Figures 2C and 2H).^{47,48} This single-cell approach provides a platform for systematic identification of genes co-expressed with such timing factors that also contribute to supporting

a particular event in cellular differentiation and morphogenesis. The association of timing factors *grh*, *opa*, and *cad* with c5 and/or c9 (mid-nc14) while germ-layer emergence is associated with c6 (late nc14) supports the view that these clusters relate to transcriptional changes driving germ layer formation through the integration of the two patterning systems (AP and DV) (c6; Figures 2K, 2L, and S2M).

Transcription factors responsible for ectoderm patterning are expressed in the mesoderm at stage 7 and play a role to repress endoderm fate in the trunk region

As our goal was to follow the gene expression program of the mesoderm germ layer during gastrulation, we next examined

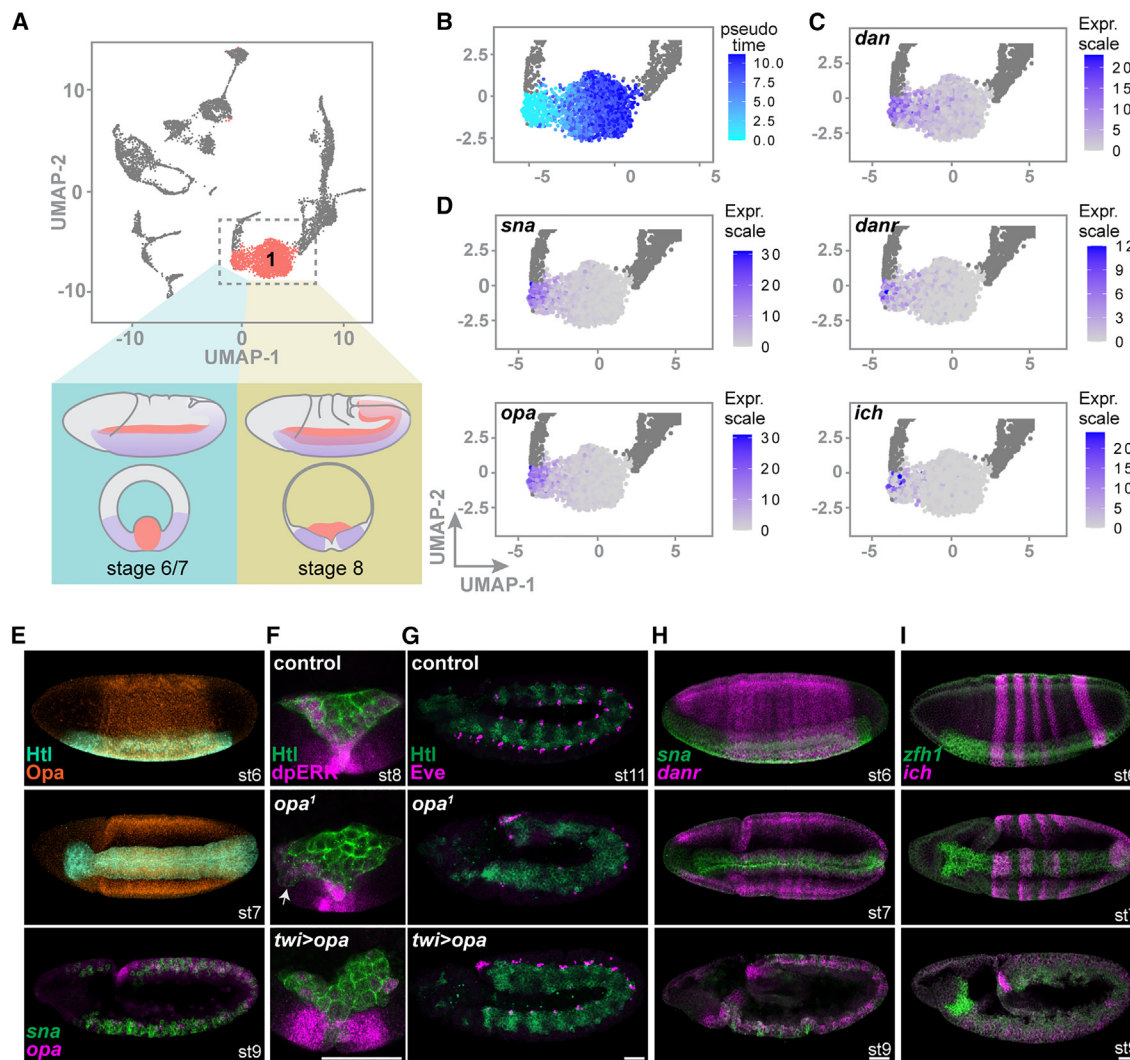


Figure 3. Transcription factors responsible for ectoderm patterning are expressed in the mesoderm as EMT is initiated and play a role to repress endoderm fate in the trunk region

(A) UMAP highlighting trunk mesoderm cells of c1 during gastrulation. Diagram depicting the morphological changes in both whole-mount and cross-sectioned views.

(B) UMAP of c1 in pseudotime with darker shade of blue indicating later developmental time points.

(C and D) Cropped UMAP plots of featured gene expression within c1; cells in view but associated with clusters other than c1 shown in dark gray. Purple intensity indicates the relative expression levels.

(E–G) Opa's role in the mesoderm. Expression of Opa protein (orange; E) and opa transcripts (purple, E) from st 6–9. Mesoderm is labeled by an anti-Htl antibody (green; E–G) or riboprobes to sna (green; E, bottom). Diphosphorylated ERK (dpERK) antibody staining (purple) indicates active receptor tyrosine kinase (RTK) signaling (purple, F). Loss of opa or overexpression of opa in the mesoderm results in asymmetry in mesoderm EMT (transverse plane, F; arrow indicates ectopic protrusions associated with opa⁷ mutant) as well as a loss of pericardial cells (anti-Eve, purple) at st 11 (G).

(H and I) Expression of danr and ich (purple) in the mesoderm (labeled with riboprobes to sna and Zn finger homeodomain 1 (zfh1) in green) detected using riboprobes. Stages are labeled at the bottom right. Scale bars, 50 μ m. See also Figure S3.

the most prominent *twi*-positive cluster, c1 (Figure 3A). This cluster contains trunk mesoderm cells that relate to st 6–8, when mesoderm cells either are invaginating to form a tube (st 6–7) or going through an EMT (st 8). The presence of *sna* and *opa* transcripts serves as a reference for staging cells in c1. The expression of both genes is initiated prior to gastrulation and continues until st 7. By st 8, however, their transcription levels have decreased significantly in the mesoderm and proteins are

no longer detectable by antibody staining (e.g., Figures 3D, 3E, and S3A).⁴⁹ Trajectory analysis (see STAR Methods) confirmed that cells that express *sna* and *opa* are positioned on one end (left side) of the pseudotime trajectory (Figure 3D compared with 3B; see also Figure 4A) and, in contrast, factors known to be expressed later are positioned on the other end (right side). St 8 (~50% of embryos in the collection) cells are expected to be the major component of c1, which is enriched for the GO

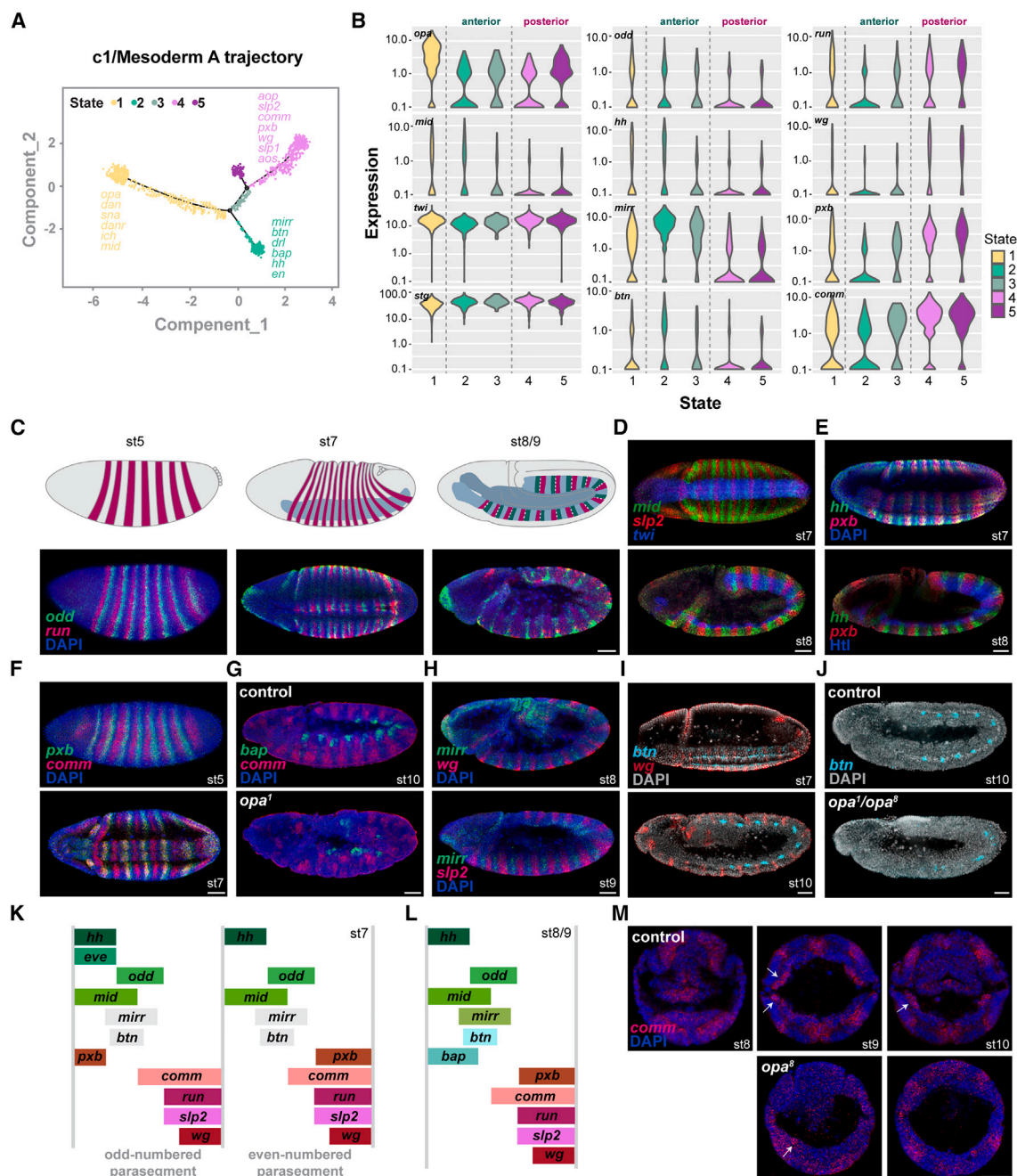


Figure 4. Subclustering and trajectory analysis of cluster 1 mesoderm cells suggests that the transcription program establishing segment polarity follows EMT

(A and B) Monocle trajectory analysis reveals five distinct cellular states (c1.1–c1.5) within c1 cells with marker genes shown (A). Violin plots of selected marker gene expression in cells at different states (B).

(C) Mesoderm-centered view of segmentation showing *odd/run* pair-rule gene expression during the process. Mesoderm is drawn in dark blue, gene expression in a segmented pattern is labeled in fuchsia from st5 to st7, while, at st8/9, the anterior parasegment expression is labeled in green (e.g., *hh*) and the posterior is labeled in fuchsia (e.g., *wg*). White dashed lines indicate the posterior/anterior boundary of the parasegments at st8/9.

(D–J) Selected marker genes of cellular state 2/3 (anterior parasegment) vs. state 4/5 (posterior parasegment) confirmed by *in situ* hybridization. Genes marking the anterior parasegment are in green while genes expressed in the posterior parasegment are in red. In *opa¹* mutants, *comm* and *bap*, normally expressed in complementary domains, are detected in irregular patterns (G), while *btn* expression is greatly reduced (J).

(K and L) Diagram depicting the segmented expression of selected marker genes from cellular state 2–5 (including *mid*) at st7 (L) and st8/9 (M).

(M) Cross sections of st8–10 embryos (transverse plane) showing localized expression of *comm* in the dorsal migrating mesoderm cells (arrows) in control vs. *opa⁸* mutants. Probe sets for *in situ* hybridization and stage of the embryos are indicated. Scale bars, 50 μ m. See also Figure S4 and Table S6.

terms relating to cell cycle (Figures 1C and S1A). Therefore, c1, containing cells from st 6/7 to st 8, represents mesoderm cells undergoing EMT and the expression of *opa*, only at the earlier stage, points to a regulatory role for it at gastrulation.

Such a role for *opa* in the trunk mesoderm (e.g., st 7–10) had not been demonstrated previously. It has been shown that *opa* is required in the mesoderm at a later stage during the specification of visceral mesoderm (st 11 and beyond).⁵⁰ We found that, in *opa*¹ mutants, EMT is aberrant (Figures 3F and S3C). Mesoderm patterning is also affected, as almost all pericardial cell precursors marked by Even-skipped (Eve) fail to be specified (Figures 3G and S3D). *opa* mutants do also exhibit ectodermal patterning defects^{20,24} that could indirectly affect the mesoderm (e.g., misplaced Pyramus [Pyr] and reduced number of *slp1* stripes; Figure S3B). Disrupting Opa function by RNAi using the *twi(2xPE)GAL4* does not effectively knock down its expression or affect mesoderm patterning as only minor defects in EMT are observed (Figure S3E). Therefore, we cannot unequivocally conclude an autonomous requirement for Opa in supporting mesodermal patterning. However, ectopic expression of Opa in the mesoderm interferes with these same processes (Figure 3G), suggesting that precise control of Opa activity within the mesoderm proper is important to support normal mesoderm EMT and cell differentiation.

Within *opa* and *sna* marked cell state 1 (of the five states identified within c1 by trajectory analysis; see Figure 4A), we were surprised to find additional transcription factors with unknown functions in mesoderm development. These include *distal antenna-related* (*danr*) and *ichor* (*ich*), detected in the trunk but excluded from the posterior and largely also from the anterior domains (Figures 3C, 3H, and 3I). As state 1 represents the younger st 6–7 mesoderm cells (discussed above; compare Figures 3C–3B), these factors may relate to a trunk-specific gene expression program that acts at gastrulation. Furthermore, they exhibit similar expression trends to *opa* when visualized by UMAP (Figure S3F; compare with Figure 2H, *opa*). As such, these genes may be co-regulated and involved in common functions and developmental processes with *opa*,^{51,52} acting to support gene expression in the embryonic trunk. The coordinated expression of this cohort of trunk-enriched transcription factors immediately prior to EMT is suggestive of a regulatory role for these genes in mesoderm cell organization and behavior at EMT (see section “discussion”).

Mesoderm cell transcription programs establish segment polarity following EMT

The segmented body plan in insects is built upon the modular organization of similar units, called parasegments, that are arranged serially along the AP axis.⁵³ In *Drosophila*, segmentation has been well studied in the ectoderm. Briefly, following the initial formation of the AP axis that takes place before cellularization, three classes of segmentation genes are activated sequentially in the blastoderm. First, expression of the gap genes in broad bands leads to the activation of pair-rule genes in a series of seven transverse stripes along the AP axis (Figure 4C). The combination of both stimulating and repressive inputs from the pair-rule gene products then establishes the expression patterns of the segment polarity genes.⁵⁴ These genes, such as *engrailed*

(*en*) and *wingless* (*wg*), play a conserved role in defining the boundary between each parasegment, which falls at the anterior edge of the *en* domain or the posterior edge of the *wg* domain^{25,55} (Figure 4C).

As discussed above, ordering c1 cells along a developmental trajectory identifies five distinct cell states, with the *opa*-marked cell state 1 (c1.1) representing cells at the earlier stage on one end and cell states 2–5 (st 8) with two additional branches on the other (Figure 4A). Consistent with *opa*'s role in regulating the segmentation gene network, markers of cell states 2–5 arise later in development within the trunk region and include various pair-rule and segment polarity genes (Figures 4A, 4B, S4A, and S4B; Table S6). Violin plots for selected markers indicate that cell states 2 and 3 (c1.2 and c1.3), associated with high expression of *en* and *hedgehog* (*hh*), comprise cells located at anterior parasegments. In comparison, cell states 4 and 5 (c1.4 and c1.5), associated with high expression of *wg* and pair-rule genes *sloppy paired 1/2* (*slp1/2*), comprise cells located at posterior parasegments (Figure 4B *hh*, *wg*; Figure S4B: *en*, *slp2*).

Several lesser-studied segmentally expressed genes were also uncovered. For instance, T-box transcription factor *midline* (*mid*) is localized anterior to *slp2* in 14 transverse stripes at st 7 (Figure 4D) and reportedly plays a role in maintaining segment polarity in the ectoderm by repressing *wg*.⁵⁶ *mid* is a marker of c1.1, whereas *slp2* is a marker for c1.2 and c1.3 (Figures 4B, S4A, and S4B). At later stages, *mid* has been found to contribute to somatic muscle morphogenesis and heart development,^{57,58} consistent with a potential function in mesoderm patterning. In addition, we found that *pxb*, which encodes a transmembrane protein and was identified as an attenuator of Hh signaling in imaginal discs,⁵⁹ is expressed in a highly dynamic pattern within the mesoderm from st 7 to st 8. At st 7, transcripts of *pxb* overlap with *hh* in the anterior odd-numbered parasegments but are excluded from the anterior region of even-numbered parasegments (Figures 4E and 4K). During st 8, the seven *pxb* stripes double to form 14 stripes and shift to the posterior region of the parasegments (Figures 4E and 4L). The doubling of *pxb* stripes in the mesoderm follows the ectoderm in time and is in register with the ectodermal stripes spatially. We also detected the *mirror* (*mirr*) gene, initially discovered as a regulator that defines the DV boundary in the eye,⁶⁰ in 14 stripes positioned anterior to *wg* and *slp2* from st 8 (Figures 4H and S4C). *mirr* encodes one of the three Iroquois homeobox transcription factors that contribute to the differentiation of cardiac progenitors (Figure S4D).⁶¹ Finally, we also observe *commisuresless* (*comm*), well documented for its role in axon guidance in the nervous system^{62,63} and recently shown to regulate ectodermal patterning during germband elongation (GBE) through enriching Myosin II at the compartmental boundaries,⁶⁴ already transitioned to single-segment periodicity in expression (i.e., 14 stripes) by st 7 and localizing to the posterior parasegments (Figures 4F and 4K). Intriguingly, *comm* expression is restricted to the dorsal mesoderm in WT embryos (arrows, Figure 4M), in line with its ectodermal expression domain, which also present in stripes from st 9 (Figure 4G). It is surprising that the most prominent transcription program associated with mesoderm cells at st 8, when EMT ensues, contains expression of segment polarity genes with demonstrated roles in the ectoderm.

Embryos with mutations in *mirr* or *comm* exhibit abnormalities in pericardial cell distribution, a phenotype typically associated with mesoderm patterning defects (Figure S4D). In addition, *opa* mutants exhibit disorganized and reduced expression of these genes; for example, *comm*, in the mesoderm and ectoderm (Figure 4G)⁶⁵ and *bap* as well as *btn* in the mesoderm (Figures 4G and 4J). It is possible that the observed mesoderm patterning defects in *opa* mutant embryos (Figures 3F, 3G, and S3B–S3E)^{50,66} relate to misregulation of segment polarity genes. Therefore, these data point to a regulatory program supported by Opa's action in the trunk that actively establishes segmented gene expression patterns in mesoderm cells following EMT, possibly to facilitate proper cell migration.

We found that two mesoderm-specific segment polarity genes, *bagpipe* (*bap*) and *buttonless* (*btn*), are expressed exclusively in the mesoderm in segmental patterns. *bap*, known to support a visceral muscle fate later during differentiation,^{65,67} is restricted to the dorsal anterior parasegments (Figures 4G, S4B, and S4C) prior to the completion of mesoderm migration.^{66,67} Transcripts of *btn* are first detectable in the invaginated tube at st 7 (Figures 4I and S4E) with some low-level expression in the procephalic and ventral ectoderm. A segmented pattern for *btn*, arranged periodically at each parasegment boundary anterior to *wg* stripes, does not become apparent until st 9 (Figures 4I and S4C). *btn* has been shown to be required for the differentiation of dorsal median (DM) cells, which are 20 cells located along the midline that contribute to axon guidance of transverse and median nerves.⁶⁸ These results suggest that the later segmentally expressed genes likely function to promote differentiation of specific mesoderm lineages.

Differential gene expression analysis reveals ecdysone signaling acts in the mesoderm as EMT ensues

Trunk mesoderm cells of c1 are at a critical transitional stage: these cells are already specified but have not yet received inductive signals for differentiation. Division and migration are the most prominent cellular changes associated with these cells. In contrast, c3 likely consists of cells of st 9 mesoderm as it is marked by genes implicated in muscle differentiation (e.g., *Him*, *meso18E*, *jeb*) and Notch signaling (e.g., *E(spl)* complex)^{69,70} (Figures 5A and S5H–S5J; Table S1).

We found that genes relating to ecdysone signaling are enriched in cells comprising c1 (i.e., st 6–8) but downregulated in c3 (Figure S5C; Table S1). c1 markers include Halloween genes *phantom* (*phm*, *Cyp306a1*),^{71,72} *dissembled* (*dib*, *Cyp302a1*),⁷³ *shroud* (*sro*),⁷⁴ and *Niemann-Pick type C-2a* (*Npc2a*), a homolog of human NPC2 linked to Niemann-Pick C type 2 neurodegenerative disease⁷⁵ (Figures 5B–5D and S5A–S5C), all of which encode enzymes in the biosynthesis pathway of ecdysone. The expression levels of these genes correlate with titers of ecdysone, which in its active form 20-hydroxyecdysone (20E) affects transcription by binding to nuclear receptor EcR.^{76,77} Ecdysteroidogenic transcription factors are known targets of ecdysone signaling and often act in a positive feedback loop to ramp up the production of ecdysone through upregulation of *phm* and *dib*.⁷⁸ Indeed, we found that expression of a dominant negative EcR receptor construct (*EcR.B2.DN*) blocks *phm* transcription by st 8 (Figure S5D).

Therefore, a role for ecdysone signaling in supporting gastrulation during early embryogenesis was investigated. Previous studies have shown that it supports border cell migration and follicle cell differentiation during oogenesis as well as dorsal closure, head involution, and tracheal and midgut morphogenesis during mid- to late-embryonic development.^{79–84} Although a *lacZ* reporter fused to the hormone binding domain (E region) of EcR was shown to respond to exogenous 20E and drives expression in the visceral mesoderm at the end of gastrulation (i.e., stage 10),^{79,85,86} a functional role for ecdysone at this stage has not been determined. The requirement of EcR for oogenesis prohibits use of a null allele to analyze its zygotic function, as inactivation of EcR in the ovary results in a multitude of defects in oogenesis, including some embryos produced with aberrant egg chamber polarity.⁸² However, expression of the dominant negative allele (*EcR.B2.DN*) using different genetic cross strategies can work to downregulate signaling in a temporal manner.⁸⁷ Through this approach, we found that maternal loading of *EcR.B2.DN* (using the *MTD-GAL4* to support expression in the ovary of first, F1, generation) disrupts AP patterning in the second (F2) generation (Figures 5E and S5E–S5G), in line with EcR's appreciated role during oogenesis.^{81,82} We then observed that early zygotic expression of *EcR.B2.DN* in F1 embryos results in abnormal mesoderm EMT at st 8 (Figure 5F) without affecting AP patterning (Figure 5E). These results suggest ecdysone signaling is important for embryonic development, specifically, to support cell movement and division at EMT during gastrulation (see section “discussion”). We further hypothesize that dysregulated ecdysone signaling might partly account for the EMT and patterning defects in *opa* mutants (Figures 3F and S3C). This is supported by the finding that *phm* expression is lost in trunk mesoderm in *opa*⁸ mutants (Figure 5G), which would be expected to result in reduced ecdysone signaling. Together, our data suggest that EcR acts downstream of Opa to, in part, support the transcriptional changes necessary for EMT of trunk mesoderm cells.

Lineage commitment dynamics for cells arising from the posterior trunk mesoderm

We next turned from the trunk mesoderm to investigate the caudal mesoderm, a less-well-characterized cell population. In the blastoderm embryo, only ~40 cells located posterior to the trunk mesoderm comprise the caudal mesoderm, which encodes a basic helix-loop-helix (bHLH) transcription factor, *HLH54F*.²⁶ *HLH54F* expression is associated with cluster 10 in the UMAP (c10; Figures 6A and S6A, *HLH54F*; Table S1) and by *in situ* hybridization in st 5/6 embryos, and it is expressed in a region posterior to the *htl* expression domain in the trunk mesoderm (Figures 6D and S6B, *HLH54F*). Consistent with their generally non-overlapping patterns in the embryo, *htl*- and *HLH54F*-expressing cells predominantly occupy distinct spaces in the UMAP of c10 (Figures 6C and 6D). This result also suggests that c10 contains at least two progenitor populations: one enriched for *HLH54F* not expressing *htl*, while the other, inversely, is enriched for *htl* not expressing *HLH54F*. To provide insight into the identities of cells in this cluster and gene expression changes associated with the caudal mesoderm, we conducted a Monocle trajectory analysis of c10 that uncovered five cellular states (Figures 6B and S6C; Table S7).

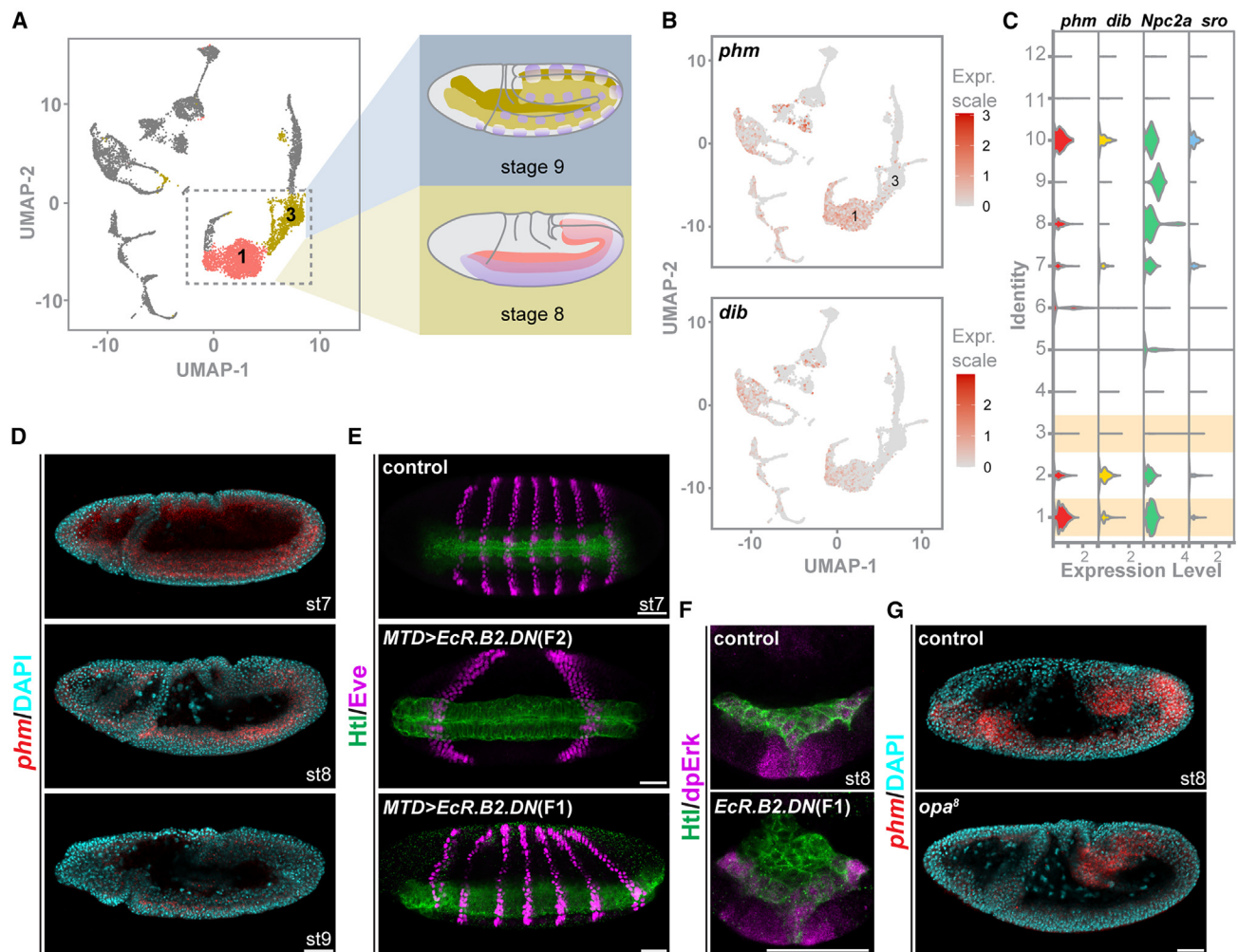


Figure 5. Genes involved in ecdysteroid hormone biosynthesis are upregulated in the mesoderm during EMT

(A) Cells from embryos of st6–8 and st9 comprise c1 and c3, respectively. Therefore, comparison of gene expression trends between these two clusters reveals transcription changes during and after mesoderm epithelial-to-mesenchymal transition (EMT).

(B and C) Expression of selected genes involved in ecdysteroid biosynthesis pathways as shown in UMAP (B) or violin (C) plots. Red intensity indicates the relative expression levels among all clusters. Mesoderm cell clusters are highlighted.

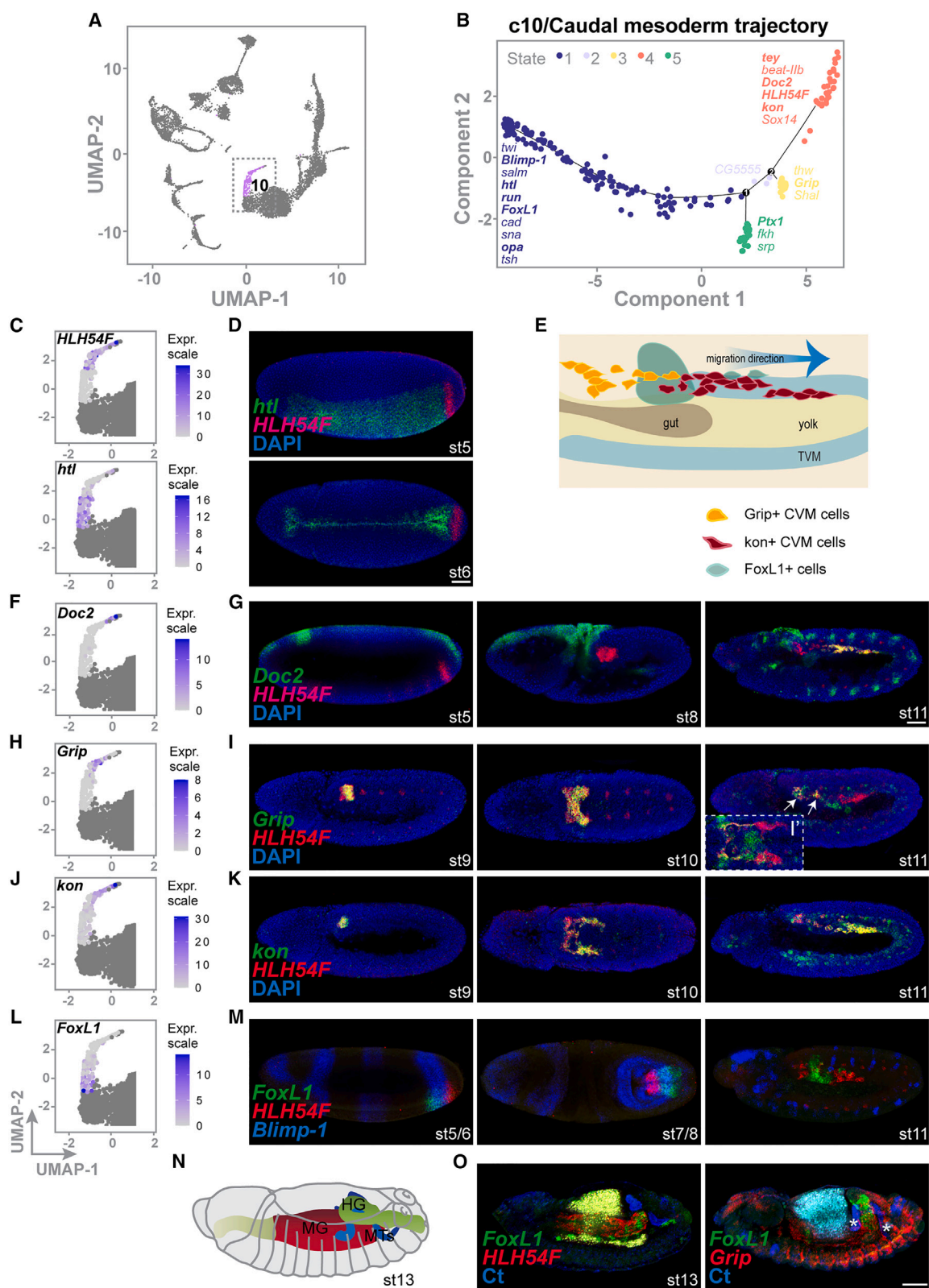
(D) *In situ* hybridization with *phm* riboprobe confirms its expression in the mesoderm is no longer detected post EMT (st9).

(E and F) Disrupting ecdysone signaling results in patterning defects in the second offspring generation F2 (E) or a mesoderm-spreading phenotype in the first offspring generation F1 (F).

(G) *phm* is downregulated within the mesoderm in *opa*⁸ mutants, suggesting a role for *opa* in regulating ecdysteroid signaling in this tissue. Scale bar, 50 μ m. See also Figure S5.

Among these five cellular states associated with c10, two cell states (c10.4 and c10.5) could be assigned to known cell types based on marker gene expression. c10.4 likely represents caudal visceral mesoderm (CVM) cells, a migratory population that expresses *HLH54F*. Cells in this subcluster also express other well-characterized CVM markers including *kon-tiki* (*kon*), *Dorsocross2* (*Doc2*), and *teyrha-meyrha* (*tey*) (Figures 6B, 6F, 6G, 6J, 6K, and S6F, *tey*). The presumptive CVM cells are specified by the end of nc14 (st 5) as a progenitor population.²⁶ Following invagination, these cells become internalized while also rapidly moving posteriorly during GBE. By st 9, they are situated in a pocket between the ectoderm and hindgut/posterior

midgut, where they begin migrating out on top of the trunk visceral muscle (TVM) cells to eventually give rise to the longitudinal muscles of the midgut.^{26,88,89} Intriguingly, *Doc2* expression is located at the very tip of the UMAP for c10 (Figure 6F). By *in situ* hybridization, *Doc2* is not expressed within the CVM until st 10, as demonstrated by colocalization with *HLH54F* (Figure 6G), suggesting that cells located at the tip of the c10 UMAP are older. On the other hand, cells of c10.5 are enriched with transcription factors expressed in the posterior endoderm and hindgut, e.g., *Ptx1* (Figures S6C and S6D, *Ptx1*), *fkh*, *Kr*, and *srp*, suggesting that c10.5 represents a cell type born at the mesoderm/endoderm border. MTs arise from exactly



(legend on next page)

this location in ring-shaped ectodermal primordia shared with the hindgut.^{28,90} MT cells can be recognized from st 10 by the expression of *Cut* (Ct), a homeobox-containing transcription factor functioning to control the evagination of MT primordia at the hindgut-midgut junction.^{91,92} Several markers for c10.5 are implicated in the development of MTs,^{93–95} making it likely that c10.5 contains precursor cells for MTs.

Cells in states 2 and 3 (c10.2 and c10.3; Figure 6B; Table S7) are most closely related to CVM cells according to the trajectory analysis. However, none of the top markers for c10.2 and c10.3 have previously characterized functions in the caudal mesoderm. Because c10.2 only contains one gene, we focused our efforts on c10.3 by further investigating the expression of its associated markers. For instance, *Glutamate receptor-interacting protein* (*Grip*) (Figures 6H, S6A, S6C, and S6D, *Grip*), encoding a scaffold protein known to regulate muscle attachment,^{96,97} was found initially co-expressed with *HLH54F* at st 9 (Figure 6I). However, by st 11, the expression patterns of *Grip* and *HLH54F* have diverged: the majority of the *HLH54F*-positive cells at the front of the migratory cohort have turned off *Grip* and instead initiate expression of *Doc2* (Figures 6I and 6I', and compare with 6G). In contrast to *Grip*, c10.4 marker *kon* also co-localizes with *HLH54F* at st 8–9 (Figures 6J, 6K, and S6D) but transitions to an opposite pattern to *Grip* at st 11: stronger in the front but diminishing at the back. *Doc2*-, *kon*-, and *Grip*-positive cells become migratory, with *Doc2*- and *kon*-coexpressing cells located at an anterior position and *Grip*-positive cells lagging behind (Figure 6J). Therefore, c10.3 likely represents a younger CVM precursor population (st 8–9) relative to c10.4, which constitutes the older CVM cell population from st 10. Upon the commitment of the *HLH54F*-expressing cells toward a muscle cell fate at st 11, the continuing expression of *Grip* defines cells that lag behind, marking a new migratory cell type that is distinct from the *Doc2*- and *kon*-positive CVM cells (Figures 6E, 6G, 6I, and 6K) (see section “discussion”). *Grip*-positive cells ultimately are incorporated into both the longitudinal muscle of midgut and the hindgut (Figures 6N and 6O).

Last, cell state 1 (c10.1) is associated with genes expressed as early as st 5, containing cells marked by *sna*, *opa*, *Blimp-1*, *salm*, as well as *hhl* likely representing progenitor cell types of st 6/7 (Figures 6B and S6A–S6D). Among markers for c10.1 is a forkhead box (Fox) transcription factor, *Forkhead box L1* (*FoxL1*)

(Figure 6L). Fox proteins are involved in fate determination of different mesoderm cell populations during organogenesis.^{98–100} *FoxL1* has been shown to contribute to muscle development during late embryogenesis.¹⁰¹ We found that *FoxL1* transcripts can be first detected in embryos by *in situ* hybridization at st 5, in a caudal region anterior to the *HLH54F*-positive CVM precursors (left panel of Figure 6M). *FoxL1* expression remains distinct from *HLH54F* throughout gastrulation (Figures S6B and S6E), while it partially overlaps with *Blimp-1* in the caudal mesoderm until st 8 (middle panels of Figures 6M and S6D). Following gastrulation, *FoxL1* is expressed in a population that co-migrates with CVM precursors at st 11 (right panels of Figure 6M). These cells ultimately locate to the anterior and posterior midgut as well as hindgut in a circular pattern around the basal surface of the endoderm, in contrast to the midgut longitudinal muscle fate of CVM cells (Figures 6N and 6O). Therefore, our analysis demonstrates that these caudal mesoderm cells in c10.1, marked by *FoxL1*, *opa*, and *Blimp-1*, are a muscle progenitor cell type that is different from the progenitors that give rise to CVM cells. These *FoxL1*-expressing progenitor cells arise early at the cellular blastoderm stage, stay localized to a region in proximity to CVM precursors, then migrate along with them (assuming a position at the back of the migrating collective) until becoming incorporated into the developing gut.

With the trajectory analysis identifying cellular states associated with distinct transcription signatures (Figure 6B) and our subsequent investigation of marker genes by *in situ* hybridization (Figures 6D, 6G, 6I, 6K, 6M, and 6O), we show that single-cell profiling can help tease apart the temporal and spatial differences in gene expression within closely related cell populations, providing insights into cell fate decisions toward lineage differentiation.

Transcription programs specify, regionalize, then segment the mesoderm cells along the AP axis

This single-cell study highlighted transcription programs that control morphogenesis from blastula to gastrula stages. Our results favor the view that the blastoderm (nc14) embryo is first carved into three domains: trunk vs. anterior and posterior poles (Figure 7A). Additionally, centered on the patterning of mesoderm, it provides insights into both temporal and spatial gene regulation that contributes to the specification (Figure 7C) and subdivision of the mesoderm cell population (diagram,

Figure 6. The single-cell transcriptomic analysis provides both temporal and spatial information on the lineage commitment of the posterior trunk mesoderm

(A and B) Trajectory analysis reveals five cellular states along with marker genes (B) that represent both temporal and spatial changes in transcription in c10 (boxed area, A). States 1–5 refer to c10.1–c10.5.
(C, D, F, and G) Cropped UMAP of *HLH54F* (C), *hhl* (D), and *Doc2* (F) are shown with expression scales. *In situ* hybridization using hybridization chain reaction (HCR) probes shows the *HLH54F*-marked CVM primordium (red) is specified at the blastoderm stage, distinct from the trunk mesoderm that expresses *hhl* (green) (D), and migrating CVM cells express *Doc2* later in development (st10/11) (G).
(E and H–K) Diagram illustrating the position of *HLH54F*-expressing CVM cells located at the front (red) or back (yellow) of the migrating collective at st10–11, and depicting the position of *FoxL1*-labeled muscle precursors (green) that co-migrate with CVM (E). Expression of state 3 marker *Grip* and state 4 marker *kon* shown by UMAP (H and J) and *in situ* hybridization (I, I', and K) colocalizing with *HLH54F*. *Grip*+ cells occupy the back of CVM migrating cohort (arrows, I; magnified view of another embryo showing dorsal view in inset, I').
(L and M) State 1 cells are marked by *FoxL1* and *Blimp-1*, which label a distinct region of the c10 UMAP plot compared to *HLH54F* (compare L with C). Expression patterns of transcription repressors *Blimp-1* (blue) and *FoxL1* (green) partially overlap, but both are distinct from *HLH54F* (red) at st5–8 (M).
(N and O) Fate of the caudal mesoderm cells. Diagram illustrates that caudal mesoderm cells contribute to midgut (MG), hindgut (HG), and malpighian tubules (MTs) at st 13 (N). *FoxL1* (green), *HLH54F*, and *Grip* (red) expression detected by *in situ* hybridization with anti-Cut antibody labeling MTs (blue, asterisks in O). Genes and stages assayed are indicated. DAPI (blue, D, F, I, and K) for counterstaining. Scale bars, 50 μ m. See also Figure S6 and Table S7.

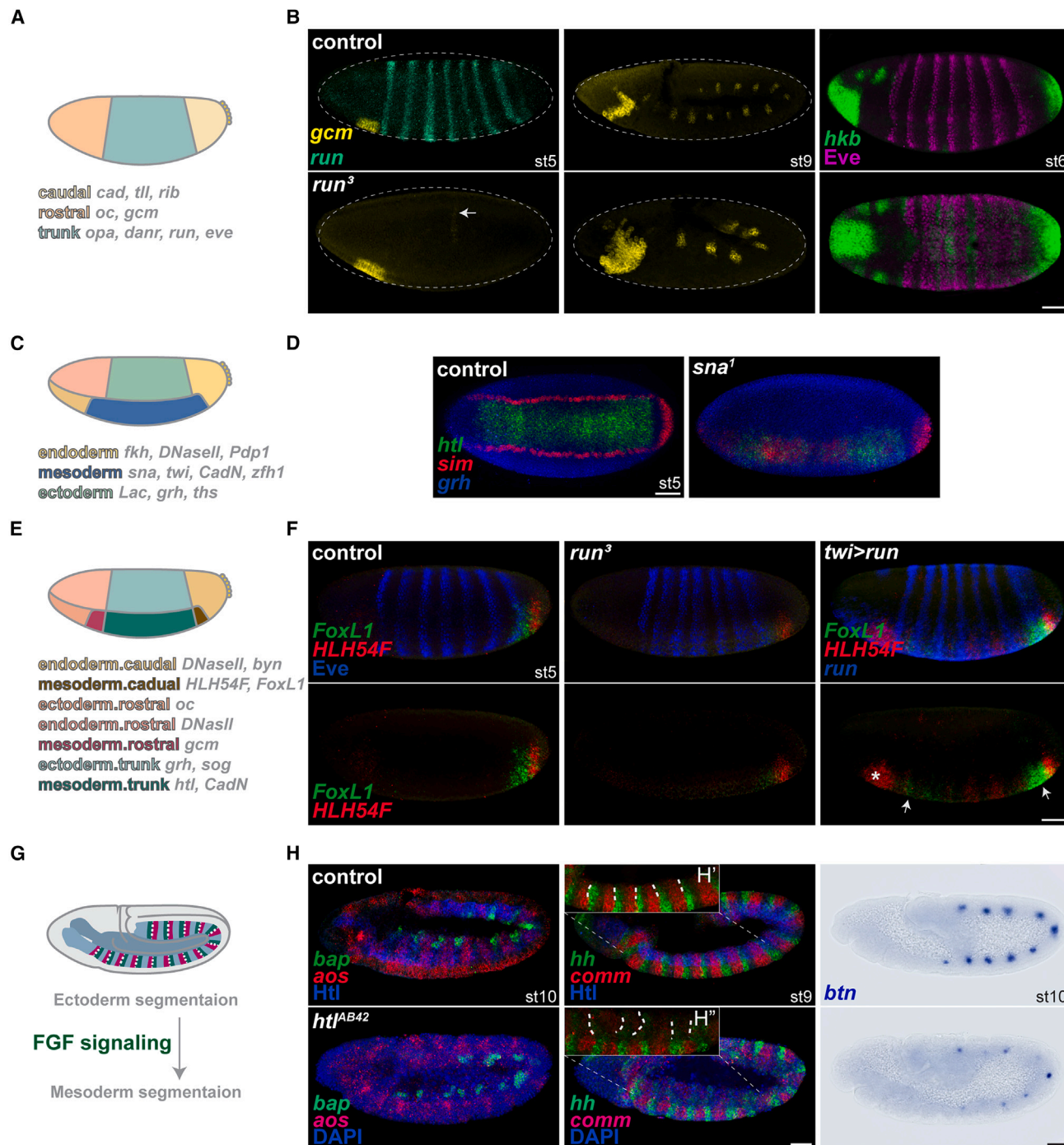


Figure 7. The temporally ordered transcription programs establish the spatial domains and tissue identities during early embryonic development

(A and B) The early blastoderm embryo is regionalized by transcription factors that specify the head, trunk, and tail, colored in beige, green, and yellow in the embryo diagram (A). *gcm*, *hkb*, and *Eve* are affected in *run*³ mutants (B).

(C and D) Three germ layers are specified during nc14 as mesodermal (blue) and endodermal (both anterior and posterior, yellow) transcription programs are clearly distinct from the ectoderm tissue (orange and green) (C). Defects in *htl*, *grh*, and *sim* expression are observed in *sna*¹ mutants (D).

(E and F) The embryo is further regionalized by the end of nc14 as the mesoderm is divided into head/rostral (fuchsia), trunk (teal), and posterior/caudal (brown) mesoderm domains (E). Loss- and gain-of-function perturbations of *run* have opposite effects on the posterior mesoderm gene expression (F).

(G and H) Mesoderm segmentation follows the ectoderm, which ensures that mesoderm cells migrate and differentiate normally (G). *Htl* antibody staining and DAPI counterstaining (both blue) were used to visualize mesoderm cells. The white dashed lines in magnified views indicate mesoderm cells located at para-segment boundaries; compare H' and H''. Genes detected and stages are labeled. Scale bars, 50 μm. See also Figure S7.

Figure 7E), which takes place prior to gastrulation. This process is followed by further specialization and lineage commitment of cells located in the posterior trunk region and segmentation of the trunk mesoderm with continuing expression of pair-rule and segment polarity genes from nc14 onwards (Figure 7G). To provide additional genetic evidence for this framework, we assayed mutants of patterning genes to examine their role in supporting such a stepwise process.

To test how disrupting the trunk gene expression program affects transcription in the poles and later on the posterior trunk cell fate, we utilized a loss-of-function allele of *run* (*run*), a pair-rule transcription factor known to act early at the blastula stage to regulate gene expression.¹⁰² *run* is initially expressed broadly in the trunk at early nc14 then quickly becomes restricted to five to seven transverse stripes from mid- to late-nc14 (Figure 2G).¹⁰² When *run* expression is broad, it is thought to inhibit anterior cell fates.^{102,103} In *run*³ mutants, we observed an expansion of *gcm* expression in the head mesoderm at nc14/st 5, becoming more apparent at st 9 (Figure 7B). The stripes associated with pair-rule gene *eve* appear compressed into a smaller area within the trunk in *run* mutants compared with WT embryos (Figure 7B, right). Furthermore, *hkb*, normally restricted to the poles prior to stage 8, is ectopically expressed in the trunk at st 6 in *run* mutants (Figure 7B, right). *opa*, on the other hand, appears to be dispensable for normal gene expression in the trunk (Figures S7B and S7C). While *run* is acting, loss of *opa* function does not lead to an obvious change in the *gcm* domain (Figure S7A).²⁰ However, *opa* is required later at gastrulation. For instance, not only does *Opa* promote gene expression within the trunk,^{20,22,24} but it also represses terminal gene expression from the trunk (Figure S7D). These results suggest that *run*, functioning in a similar way but earlier than *opa*, is a critical component of the trunk gene regulatory network that restricts the expression of terminal genes to the anterior and posterior poles.

We used *sna*¹ mutants to test how specification of one germ layer influences the other germ layers. *sna* is required for the specification of mesoderm as well as mesectoderm in two rows of cells abutting it. Loss of *sna* results in the loss of mesoderm cell fate as *htr* expression is not maintained (Figure 7D). In addition, the gene *single-minded* (*sim*) is expressed only at low levels within patches in ventral regions of the embryo, indicating a failure to define mesectoderm. Last, *grh* expression (normally restricted to the ectoderm) expands into the ventral domain overlapping with *htr* in the *sna*¹ mutant (Figure 7D). These observations indicate that disrupting the transcription program active in the mesoderm alters gene expression in other tissue types and that the emergence of mesoderm cells at mid-nc14, driven by *sna* and *twi*, is a crucial programmatic step toward gastrulation.

sna activity is followed by division of the mesoderm into head, trunk, and posterior domains. To assay how posterior mesoderm cells respond to dysregulation of the trunk program, we again turned to *run*, which is a marker of the c10.1 cell state that likely represents progenitor cells that give rise to visceral muscles (Figures S7E and S7G). *run* pair-rule expression pattern at st 5/6 is dynamic (Figure S7F). At st 5, the *run* seventh stripe at the posterior overlaps with both *FoxL1* and *HLH54F* domains. However, just a bit later at st 6 when the *run* pattern encom-

passes 14 stripes, one stripe arises at the posterior that is in line with the *HLH54F* positive domain (Figure S7F). This suggests that *run* may act to support both these distinct cell populations. In *run*³ mutants, *FoxL1* levels are greatly reduced (Figure 7F), whereas overexpressing *run* in the mesoderm using a *twi*-*GAL4* driver causes a concomitant increase in levels of *FoxL1* and *HLH54F* in the trunk as well as the anterior (arrows and asterisk, Figure 7F). *opa* is also enriched in c10.1 cells that express *FoxL1* (Figures 6B and S6C), but neither *FoxL1* or *HLH54F* levels are affected by loss of *opa*, consistent with a recent study in which no posterior patterning defects were detected by computer simulation.²¹ Together, these results suggest that mesoderm cell fates are tightly controlled within the trunk and that the size of a lineage precursor population is determined by cooperative regulations involving multiple transcription factors (see section “discussion”).

Our dataset also points to a segment polarity program in the mesoderm potentially contributing to their proper migration as a collective (Figures 4K–4M). The fibroblast growth factor (FGF) signaling pathway is essential to mesoderm development by coordinating cell division, cell-shape change, and movement during gastrulation as well as by promoting cardiac and somatic muscle differentiation later during organogenesis.^{49,104} Using a loss-of-function *htr*^{AB42} allele,¹⁰⁵ we tested whether FGF signaling is required for the maintenance of segment polarity within the mesoderm by examining the localized expression of markers identified in the c1 trajectory analysis (Figures 4A and 7H). Indeed, *bap* is a known FGF target and, as expected, expressed in irregular patches. However, *aos*, *hh*, and *comm* also all appear to be expressed at lower levels in *htr*^{AB42} mutants (Figure 7H). Intriguingly, *hh* and *comm* stripes in the mesoderm are uneven in the mutant within the mesoderm (Figure 7H; compare Figure 7H' with 7H'), suggesting that mesoderm cells fail to move in a coordinate manner with the ectoderm during GBE. Signaling pathways including FGF serve as bridges and messengers from the ectoderm to the mesoderm to ensure that the two germ layers are patterned concomitantly.^{66,106} We hypothesize that the continual expression of segment polarity genes in the mesoderm is crucial for supporting proper collective behavior during mesoderm cell migration.

DISCUSSION

By investigating gene expression trends using conventional genetic tools assisted by single-cell transcriptomic information, this study contributes to our understanding of how gene expression programs are orchestrated in a stepwise manner to lay down the basic body plan and its modular units to coordinate complex morphogenetic processes.

One of the advantages of our approach is that it illuminates several sequentially acting transcription programs that are associated with gene expression changes during early *Drosophila* embryogenesis, encompassing the transition from blastula to gastrula stages. In contrast to our initial assumption that we would see cells differentiated by early patterning programs relating to germ layer formation, we were surprised to find that AP position, specifically the trunk versus terminal programs, was more influential when grouping blastoderm cells according

to their transcriptional states. This result is consistent whether or not cells are fixed (i.e., f.c6 separates from f.c3; Figure S2D). Furthermore, our data that cells in the posterior pole, which express *cad* and *tl*, are preferentially associated with a “younger” cell state in the blastoderm (e.g., Figures 2B, 2E, 2F, S2A, S2C, and S2E) may relate to delay of maternal transcript degradation from the poles and/or to increased complexity of zygotic transcriptome associated with cells in the trunk versus termini. While it is also possible that products of germ plasm mRNAs enriched at the posterior pole act to repress transcription and cause a delay in maternal-to-zygotic-transition (MZT),^{107,108} further investigation is required to demonstrate their role in regulating gene expression outside germ cells and examine how they affect AP patterning in the blastoderm embryo. Recent studies have suggested that *Drosophila* embryos, despite being long germ-band embryos, have retained properties of an ancient short-germband program in which posterior cells act with multipotent potential to generate segments in a delayed manner relative to segment specification in the trunk.^{21,25} Our data support this view that patterning of termini is delayed and suggest that single-cell sequencing experiments can provide deep insights into how transcription timing is coordinated within different domains of the blastoderm, or, as manifested by transcription progression, how developmental speeds are differentially regulated in cells at different locations.

While this dataset serves as a resource for understanding the stepwise progression of all three germ layers, we specifically focused on the patterning and specialization of mesoderm lineages in our in-depth analyses, finding that allocation/divergence of the terminal lineages (rostral and caudal mesoderm) from early trunk mesoderm is accomplished prior to the morphogenetic movements associated with gastrulation, namely invagination, EMT, and GBE.

In the trunk, Opa continues (from the blastula stage) to act as a timing factor at the initial phase of EMT during gastrulation. Previously shown to drive segmental patterning in the ectoderm, we find that Opa is broadly expressed in the trunk mesoderm at EMT, preceding and likely required for the transcription of a number of segmentally expressed genes enriched within the same cluster. We show that several of these segmentation genes are implicated in patterning and ultimately also required for differentiation of mesoderm cell lineages through analysis of mutant phenotypes (e.g., *comm* and *mirr*; see Figures 4N and S4D). Other markers that were not investigated include *Toll* (*Tl*) and *derailed* (*drl*) in cell states 2 (c1 trajectory analysis) and *anterior open* (*aop*) and *argos* (*aos*) in state 4 (Figure 4A; Table S5). It is unlikely that Opa acts alone in regulating transcription timing, and potential roles for additional transcription factors were also highlighted by our results (e.g., Figure 3). Therefore, future experiments would focus on interrogating the mesoderm-specific requirement for *opa* during gastrulation and characterizing the contributions of these other factors to mesoderm patterning.

The unexpected segmentation gene expression we observed downstream of Opa is intriguing, as it signifies that part of the Opa-dependent transcription program also relates to promoting planar polarization of the migrating mesoderm cells. With the integrated dataset, an *opa*, *sna*-enriched cell cluster (i.c11) and two separate clusters corresponding to the anterior and posterior para-

segment identities (i.c4 and i.c5, respectively) emerge from those cells undergoing EMT (Figures S4F and S4G). We suggest that the Toll genes, as well as other segment polarity genes, play a role similar to their function in patterning the ectoderm during GBE,¹⁰⁹ regulating mesoderm morphogenesis relating either to establishment of cell polarity (planar, apicobasal, or front-back) and/or in the regulation of spatially distinct actomyosin activities to keep the relative position of mesoderm cells in check during migration. EMT, the process through which mesoderm cells acquire their migratory ability, is tightly coupled with GBE¹¹⁰ and these two processes may be regulated by the same segmentation gene network controlling AP patterning. Notably, mutations in segmentation genes affecting ectodermal planar cell polarity also lead to mesoderm spreading defects.^{65,66} Furthermore, mesoderm cells likely lose polarity information during division as EMT occurs (Figure 4M) and have to regain planar polarity (i.e., segmental polarity) to support coordinate posterior movement following GBE of the ectoderm. Such a process allows mesoderm cells to position themselves in association with ectodermal lineages in order to form the proper ratio of different muscle types.

Surprisingly, a gene expression program associated with ecdysone biosynthesis also plays a role as trunk mesoderm cells prepare for/undergo EMT (Figures 5D and 5E). This finding is interesting in that it suggests that a hormonal/metabolic cue through ecdysone influences EMT at gastrulation in *Drosophila* embryos. However, it remains unclear what tissues provide the source for embryonic ecdysone (E) and when ecdysone signaling is first required during embryogenesis. An even earlier role supported by maternal products of the pathway could indirectly affect the movements of mesoderm cells. Future experiments are needed to address those questions. For neural crest cell delamination, recent studies have identified that metabolic triggers are important for regulating gene expression.¹¹¹ It is possible that the *Drosophila* embryo similarly does not initiate the GBE/EMT program unless the proper metabolic environment is present. Since cell movement requires a significant expenditure of energy, this link between metabolism/hormones and cell movements is understandable.

This study also provides insight into how multipotent mesodermal progenitor cells are carved out early in embryogenesis through the combinatorial input of transcription factors, including timing factors. Our developmental trajectory analysis identified transcription factors, expressed in spatially distinct domains, that allocate the posterior ventral regions of the blastoderm to ultimately give rise to neighboring migrating cell types. We show that *FoxL1*, along with several other transcription regulators (e.g., *Blimp-1*, *salm*; see c10.1; Figure 6B), marks a progenitor cell type located anterior to CVM in the caudal mesoderm. Mesoderm cells in the caudal region are also alternatively named for the cell types they ultimately contribute to. For example, CVM corresponds to longitudinal visceral muscle (LVM), while the muscle cells in the hindgut are referred to as hindgut visceral muscles (HVMs).^{112,113} The *FoxL1*+ cells therefore represent a progenitor population established at stage 5 that likely contributes to the HVM as well as TVM. The transcription program responsible for the specification and later migration of these caudal mesoderm cell types are distinct from the trunk and relate to the terminal program (i.e., *torso*, *tl*, as well as timing factor *cad*). However, genes functioning in the trunk can still have an impact. For instance,

run, previously shown to repress *ocelliless* in the anterior,^{102,103} also functions in the posterior blastoderm as part of the transcription regulatory network to activate expression of *FoxL1* and possibly *HLH54F* in the progenitor populations.

While it is appreciated that CVM cells appear to also contribute to the MTs, the *Drosophila* kidney equivalent,¹¹⁴ direct evidence arguing for a secondary developmental outcome for *HLH54F*-positive mesoderm cells had not been demonstrated. Analyzing gene expression at single-cell resolution enabled us to discover subtle differences in transcriptomic signature even for patterns composed of a small number of cells. Here, we provide molecular support for the finding that CVM indeed contributes to MTs in addition to midgut longitudinal muscles. A possible hindgut fate is linked to the expression of *Grip*. Specifically, we found that there is a deterministic transcription program involving maintenance of *Grip* expression in a subset of CVM cells that accounts for how a subset of cells located at the back of the CVM migrating collective become MTs vs. other tissues and those located at the front become longitudinal muscles. Without the insight of Monocle trajectory analysis to further subdivide cells within clusters, we would not be aware that the CVM migrating collective has a different gene expression program for cells located at the front vs. back of the migrating collective.

In summary, by highlighting regulatory steps in transcriptional progression at single-cell resolution, this study contributes to our understanding of how animals control and coordinate morphogenetic processes and paves the way for future mechanistic analysis of gene regulatory networks and signaling pathways that account for the specification of multipotent progenitor cell types as well as collective cell migration. The single-cell transcriptomic map generated in this study also serves as a resource for future research focusing on gastrulation and, in particular, mesoderm development.

Limitation of the study

In this study, we chose to focus analysis on scRNA-seq data acquired from live cells isolated by embryo homogenization. As we wanted to decrease processing time and because the isolation procedure leads to a great deal of material loss, we could only infer the developmental stages of the embryos within the ~90-min interval based on the gene expression information (*in situ* hybridization data) and comparisons with temporal information provided by previous studies. In addition, the following assumptions were made for data analysis: first, that cells do not age, or rather transcription does not progress, once cells are placed on ice; and second, that the isolation procedure, including physical (i.e., Douncer) and chemical (i.e., Accumax) methods used, as well as incubation with the magnetic beads during dead cell removal, do not introduce transcriptional changes. Furthermore, for cell types such as the caudal mesoderm, cell numbers are low, and such analyses in the future could benefit from enriching rare cell populations.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.celrep.2023.113289>.

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AUTHOR CONTRIBUTIONS

A.S. and J.S. conceived the project. J.S. planned the experimental approach. A.S. directed the project. J.S. performed all experiments and F.G. performed the computational work. J.S., C.Z. and F.G. analyzed the data with input from A.S. The manuscript was written by J.S. and A.S. with input from F.G. and C.Z.

DECLARATION OF INTERESTS

The authors declare no competing interests.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit anti-Odd paired	This study	N/A
Guinea pig anti-Heartless	Stepanik et al. ¹¹⁵	N/A
Rabbit anti-Phantom (Phm)	Dr. Michael O'Connor (Minneapolis, MN, USA)	N/A
Mouse anti-Even-skipped	Developmental Studies Hybridoma Bank (DSHB)	2B8; RRID: AB_528230
Goat anti-GFP	Rockland Immunochemicals	Cat# 600-101-215; RRID: AB_218182
Rabbit anti-RFP	MBL International	Cat# PM005; RRID: AB_591279
Rabbit anti-Beta-Galactosidase	MP Biomedicals	Cat# 559761; RRID: AB_2687418
Monoclonal Anti-MAP Kinase, Activated	Sigma-Aldrich	Cat# M9692; RRID: AB_260729
Mouse anti-Biotin (Z021)	Thermo Fisher	Cat# 03-3700; RRID: AB_2532265
Rabbit anti-FITC polyclonal antibody	Thermo Fisher	Cat# A-889; RRID: AB_221561
Sheep anti-Digoxigenin polyclonal antibody	Thermo Fisher	Cat# PA1-85378; RRID: AB_930545
Anti-Digoxigenin-AP, Fab Fragments Antibody	Sigma-Aldrich	Cat# 11093274910; RRID: AB_2734716
Alexa Fluor 488 goat anti-mouse	Molecular Probes	Cat# A-21202; RRID: AB_141607
Alexa Fluor 488 goat anti-guinea pig	Molecular Probes	Cat# A-11073; RRID: AB_2534117
Alexa Fluor 488 donkey anti-guinea pig	Jackson Immuno Research Labs	Cat# 706-545-148; RRID: AB_2340472
Alexa Fluor 555 donkey anti-rabbit	Molecular Probes	Cat# A-31572; RRID: AB_162543
Alexa Fluor 555 donkey anti-sheep	Molecular Probes	Cat# A-21436; RRID: AB_2535857
Alexa Fluor 647 donkey anti-rabbit	Molecular Probes	Cat# A-31573; RRID: AB_2536183
Alexa Fluor 647 donkey anti-mouse	Molecular Probes	Cat# A-31571; RRID: AB_162542
Chemicals, peptides, and recombinant proteins		
Boitin RNA labeling mix	Roche	55612420
DIG RNA labeling mix	Roche	57127420
Fluorescein RNA labeling mix	Roche	59973820
T7 RNA polymerase	Roche	13644022
Critical commercial assays		
Chromium Next GEM Single Cell 3' v3.1 (Dual index)	10X Genomics	11684795910
Dead Cell Removal Kit	Miltenyi Biotec	130-090-101
LIVE/DEAD Cell Imaging kit (488/570)	Thermo Fisher	R37601
Accumax®	Innovative Cell Technologies	#AM-105
Hybridization Chain Reaction	Molecular Instruments	N/A
Deposited data		
Single-cell RNA-seq data	This study	GEO: GSE222660
Visualization of UMAP graph	This study	Mendeley data: https://doi.org/10.17632/s25d8y9pvz.1
Experimental models: Organisms/strains		
<i>D. melanogaster</i> : <i>ht^{AB42}/TM3,ftz-lacZ</i>	Bloomington Drosophila Stock Center (BDSC)	#5370
<i>D. melanogaster</i> : <i>yw;opa¹, slp1-DESE-lacZ, TM3, twi-GFP</i>	BDSC	#3312
<i>D. melanogaster</i> : <i>opa⁸/TM3, twi-GFP</i>	BDSC	#5335
<i>D. melanogaster</i> : <i>run³/FM7c, ftz-lacZ</i>	BDSC	#56499
<i>D. melanogaster</i> : <i>yw;UAS-run[[232]/CyO</i>	Prazak et al. ¹¹⁶	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
<i>D. melanogaster</i> : UAS- <i>opa</i> [D10]	Prazak et al. ¹¹⁶	N/A
<i>D. melanogaster</i> : <i>Adh</i> ¹⁷ <i>sna</i> ¹ <i>cn</i> ¹ <i>vg</i> ¹ /CyO	BDSC	#25127
<i>D. melanogaster</i> : <i>y</i> [1] <i>w</i> [*]; <i>P</i> { <i>w</i> [+ <i>mC</i>] =GAL4- <i>twi</i> .2xPE}3	BDSC	#58804
<i>D. melanogaster</i> : <i>y</i> [1] <i>w</i> [*]; <i>P</i> { <i>w</i> [+ <i>mC</i>] =GAL4- <i>twi</i> .2xPE}2	BDSC	#2517
<i>D. melanogaster</i> : <i>P</i> { <i>ry</i> [+ <i>t7.2</i>]= <i>hsFLP</i> }1, <i>y</i> [1] <i>w</i> [1118]; <i>P</i> { <i>w</i> [+ <i>mW.hs</i>]= <i>FRT</i> (<i>w</i> [<i>hs</i>]))G13 <i>P</i> { <i>w</i> [+ <i>mC</i>]=UAS- <i>mCD8::GFP.L</i> }LL5	BDSC	#5131
<i>D. melanogaster</i> : <i>y</i> [1] <i>w</i> [*]; <i>P</i> { <i>w</i> [+ <i>mC</i>]=UAS- <i>mCD8::GFP.L</i> }LL5, <i>P</i> {UAS- <i>mCD8::GFP.L</i> }2	BDSC	#5137
<i>D. melanogaster</i> : <i>P</i> { <i>w</i> [+ <i>mC</i>]= <i>otu</i> - <i>GAL4::VP16.R</i> }1, <i>w</i> [*]; <i>P</i> { <i>w</i> [+ <i>mC</i>]= <i>GAL4</i> - <i>nanos.NGT</i> }40; <i>P</i> { <i>w</i> [+ <i>mC</i>]= <i>GAL4::VP16</i> - <i>nanos.UTR</i> }CG6325[MVD1]	BDSC	#31777
<i>D. melanogaster</i> : <i>y w</i> ; <i>P</i> (<i>mat-tub-Gal4</i>) <i>mat67</i> ; <i>P</i> (<i>mat-tub-Gal4</i>) <i>mat15</i>	Staller et al. ⁸⁷	line 2318
<i>D. melanogaster</i> : <i>w</i> [*]; <i>P</i> { <i>w</i> [+ <i>mC</i>]=UAS- <i>EcR.A.W650A</i> }TP5	BDSC	#9451
<i>D. melanogaster</i> : <i>w</i> [*]; <i>P</i> { <i>w</i> [+ <i>mC</i>]=UAS- <i>EcR.B2.W650A</i> }TP5	BDSC	#9449
<i>D. melanogaster</i> : <i>y</i> [1] <i>sc</i> [*] <i>v</i> [1] <i>sev</i> [21]; <i>P</i> { <i>y</i> [+ <i>t7.7</i>] <i>v</i> [+ <i>t1.8</i>]= <i>TRiP.HMS01185</i> }attP2/ <i>TM3</i> , <i>Sb</i> [1]	BDSC	#34706
<i>D. melanogaster</i> : <i>y</i> [1] <i>w</i> [*]; <i>P</i> { <i>y</i> [+ <i>t7.7</i>] <i>w</i> [+ <i>mC</i>]= <i>E(spl)m5-HLH-GFP.FPTB</i> }attP40	BDSC	#66445
<i>D. melanogaster</i> : <i>y</i> [1] <i>w</i> [*]; <i>P</i> { <i>y</i> [+ <i>t7.7</i>] <i>w</i> [+ <i>mC</i>]= <i>E(spl)mdelta-HLH-GFP.FPTB</i> } attP40	BDSC	#68191
<i>D. melanogaster</i> : <i>y</i> [1] <i>w</i> [*]; <i>P</i> { <i>y</i> [+ <i>t7.7</i>] <i>w</i> [+ <i>mC</i>]= <i>mirr-GFP.FPTB</i> }attP40	BDSC	#68183
<i>D. melanogaster</i> : <i>y</i> [1] <i>w</i> [*]; <i>P</i> Bac{ <i>y</i> [+ <i>mDint2</i>] <i>w</i> [+ <i>mC</i>]= <i>btn-GFP.FPTB</i> }VK00037	BDSC	#56154
Oligonucleotides		
HCR probes ¹¹⁷ (Table S8)	Molecular Instruments	Table S8
PCR primers used to generate riboprobes (Table S8)	This paper	Table S8
Recombinant DNA		
<i>danr</i> cDNA clone RE72284	Drosophila Genomics Resource Center (DGRC)	RRID: DGRC_10042
<i>ich</i> cDNA clone RE65372	Drosophila Genomics Resource Center (DGRC)	RRID: DGRC_1135960
<i>mid</i> cDNA clone LP04777	Drosophila Genomics Resource Center (DGRC)	RRID: DGRC_1337803
Software and algorithms		
Zeiss Zen 3.0 (Blue edition)	Zeiss	RRID: SCR_013672
Adobe Photoshop v24.1	www.adobe.com	RRID: SCR_014199
Adobe Illustrator v27.1.1	www.adobe.com	RRID: SCR_010279
Cell Ranger v7.0	10x Genomics	RRID: SCR_017344
Seurat ¹¹⁸ v3.2.0, v4.1.0 ¹¹⁹	https://satijalab.org/seurat/	RRID: SCR_016341
DoubletFinder ¹²⁰	https://github.com/chris-mcginnis-ucsf/DoubletFinder	RRID: SCR_018771

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Monocle2 ^{121–123}	http://cole-trapnell-lab.github.io/monocle-release/docs/	RRID: SCR_016339
g:Profiler ¹¹⁷	https://biit.cs.ut.ee/gprofiler/gost	RRID: SCR_006809
Other		
Qubit™ dsDNA HS Assay	Thermo Fisher	Q32854
Bioanalyzer High Sensitivity DNA Analysis	Agilent	5067–4626
HiSeq X	Illumina	N/A
NovaSeq 6000	Illumina	N/A
R scripts and markdown files	This study	https://github.com/StathopoulosLab/Single_Cell_RNAseq_Analysis_supplementary ; https://zenodo.org/badge/latestdoi/668423939

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Angelike Stathopoulos (angelike@caltech.edu).

Materials availability

Drosophila strains and other reagents generated in this study will be available upon request from the [lead contact](#), or the commercial sources listed in the [key resources table](#).

Data and code availability

- The raw sc-RNA sequencing data generated in this study has been submitted to NCBI database (<https://www.ncbi.nlm.nih.gov/geo/>) and is available under accession number GEO: GSE222660
- A github repository was generated and publicly available with the analysis codes for all datasets (https://github.com/StathopoulosLab/Single_Cell_RNAseq_Analysis_supplementary or <https://zenodo.org/badge/latestdoi/668423939>)
- Additional Rds objects, R script and installation instructions for visualizing RNA sequencing UMAP graphs are deposited into Mendeley Data Respository (Mendeley data: <https://doi.org/10.17632/s25d8y9pvz.1>)
- Any additional information required to reanalyze the data shown in this paper is available from the [lead contact](#) upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

Fly stocks and husbandry

All stocks were kept at 22°C in the standard medium. Experimental crosses were kept in cages with apple juice agar plates supplemented with yeast paste. yw were used to show the wild-type expression pattern, and heterozygous embryos from the mutant collections were used as the controls. Embryos overexpressing run were generated by crossing *twi(2PE)-GAL4* females (BDSC#58804, #2517) with UAS-run (gift from Peter Gergen) males. *EcR* dominant negative zygotic mutants (F1) were generated by crossing MTD-GAL4 (BDSC#31777) virgin females with UAS-*EcR.B2.W650A* (BDSC#9449) males. Two step crosses were used to obtain the maternal mutants (F2). First, MTD-GAL4 males with UAS-*EcR.B2.W650A* virgin females, virgins collected from the progeny then crossed with UAS-*EcR.B2.W650A* males.

METHOD DETAILS

Embryos collection, cell isolation and library construction

Drosophila embryos produced by crossing *twi(2PE)-GAL4* females (BDSC#58804, #2517) and UAS-*mCD8GFP* (BDSC#5131, #5137) males were collected on apple agar plates at 22°C. Two sessions of 45 min pre-lay collections were performed in order to clear the older embryos held by the females. After collecting for 1 h and 30 min, embryos were aged for 3 h and 15 min. A similar timed embryo collection followed by immunostaining with a GFP antibody (1:2000, Rockland) was done in advance to estimate the percentage of embryos at a specific developmental stage obtained, indicating that our collection resulted in an embryonic sample in which the majority (~50%) were at st8, ~30% at st5-7, ~20% at st9-10 and very few older than st10 (~1%). This window allowed us to study both

the patterning of embryos that contributes to germ layer designation (st6-7), germband elongation (GBE) and mesoderm epithelial-to-mesenchymal transition (EMT) (stage 8), as well as the initial differentiation of tissues (stages 9–10).

Prior to cell isolation, Accumax was thawed, and the Douncer was cleaned. Schneider's medium, PBS, Douncer and low-binding tubes were kept on ice. Embryos were dechorionated with 50% bleach/water, rinsed well, then immediately transferred to the Douncer-containing cold media. Gross dissociation was done with the loose pestle by gently moving it up and down ~15 times. After brief centrifugation at 300 RCF to pellet the cells at 4°C, 5min room temperature Accumax treatment was performed to further dissociate the cells. Following washing in cold media twice, cells were filtered through a 30µm MACS cell strainer (Miltenyi Biotec). Lastly, the dead cell removal kit (Miltenyi Biotec) was used to clean up the dying cells and cell debris. The final cell stock was resuspended in cold PBS with 1% BSA, then promptly counted in order to be diluted to an ideal concentration (~900 cells/µl) for the GEM generation following the recommended protocol from 10x Genomics (CG000315 RevC). The LIVE/DEAD cell imaging kit (Invitrogen) was used in test runs to evaluate the viability of the cells (typically >95% after dead cell removal) in the final single-cell suspension. The Chromium Next GEM Single Cell 3' v3.1 kit (10x Genomics) with dual indexes was used for library construction and sequenced on an Illumina HiSeq X platform (Fulgent, Temple City, CA).

Care was taken to expedite processing such that homogenization to running the Chromium Controller occurred in about one and a half hours. Alternatively, protocols that homogenize and then fix cells can have lengthy processing steps. It is possible that particular gene expression bias could be associated with different protocols. This is also why we chose to validate our clusters using *in situ* hybridization or antibody staining of fluorescent protein-tagged genes; in every case (for over 40 genes examined), we detected expression in cells of the tissue type we had expected based on the clustering data. We did find that the mesoderm cell population was overrepresented (4865 out of the original 9845 cells for live wild-type#1, [Figure S1B](#)), which may relate to the fact that mesoderm cells are mesenchymal, are detached from each other during gastrulation, and therefore may be more likely to survive the cell isolation process. This was fortuitous, as our goal was to study patterning and signaling dynamics associated with the mesoderm at gastrulation.

Generation of additional datasets

Live WT#1 is the focus of this study, unless otherwise noted. Two additional wild-type datasets (live WT#2 and fixed WT#3) were generated as biological replicates. Embryos collection, cell isolation and dead cell removal were performed as described above. Methanol fixation and rehydration was done largely following the suggested protocol from 10x Genomics (CG000136 RevB). Briefly, 400µL of chilled methanol was added to 100µL of cell suspension drop by drop while gently mixing to prevent clumping. Cells were fixed on ice for 15 min before being transferred to –20°C for short term storage. For rehydration, cells were first placed on ice for 15 min to equilibrated to 4°C, centrifuged at 1000 RCF for 5 min at 4°C and then resuspended in rehydration buffer (1% BSA, 0.5u/µl RNase inhibitor in PBS). One repeat sample of live cell suspension (WT#2) and one sample from dehydrated methanol-fixed cells (WT#3) were processed together using the Chromium Next GEM Single Cell 3' v3.1 kit (10x Genomics). The constructed libraries were sequenced on an Illumina NovaSeq 6000 platform.

Cluster annotation and validation

Berkeley *Drosophila* Genome Project (BDGP) database was used for cluster annotation. With the expression patterns for ~8600 genes in staged embryos detected by *in situ* hybridization, it also provides standard vocabularies to annotate gene expression patterns during embryonic development. Following the list of markers ([Tables S1](#), [S4](#), and [S5](#)), we checked the top 20 genes enriched in each cluster, especially those with more restricted expression domains and time windows. The identity of the cluster was also verified with *in situ* hybridization or antibody staining.

Fixation, *in situ* hybridization, and immunostaining

Embryos were fixed following the standard protocols. Riboprobes for *wg*, *hh* and *run* were made from cDNA cloned into the pBlue-script vector and reverse transcribed with T7 RNA polymerase (Roche). For, *danr* (RE72284), *ich* (RE65372) and *mid* (LP04777), full-length cDNA clones from DGRC were used as templates. The rest of RNA probes used in this study were made by amplifying a 400–1100 bp region from a genomic DNA sample extracted from wild-type flies, with the reverse primers containing a T7 promoter sequence to facilitate reverse transcription. The antisense RNA probes labeled with Digoxigenin, Fluorescein and Biotin (Roche) were used in combination with primary antibodies (1:400) of different origin (Thermo Fisher) to detect the *in vivo* expression of target genes.¹²⁴ Alexa Fluor (488, 555, 647) secondary antibodies (1:500; Molecular Probes) were used for fluorescent signal amplification and detection. Anti-DIG-alkaline phosphatase (1:400; Sigma) with its NBT/BCIP substrates were used to detect *btn* expression in *htr^{AB42}* mutant background. Standard protocol for antibody staining was used. Dilutions for the primary antibodies were as such: guinea pig anti-Htl (1:800; in house), rabbit anti-Opa (1:200; in house, see below), goat anti-GFP (1:1000; Rockland immunochemicals), mouse anti-Eve (1:80; DSHB), rabbit anti-β-Gal (1:800; MP Biomedicals), rabbit anti-Phtn and guinea pig anti-Sro (1:500, both kind gifts from Dr. Michael O'Connor, Minneapolis, USA). Same secondary antibodies and dilution were used. Purified MBP-Opa amino acids Q61-T410, 6xHis protein was used to immunize a rabbit (Pocono Rabbit Farms and Lab, Canadensis, PA). Subsequently anti-Opa IgG was affinity purified from serum and concentrated by ammonium sulfate precipitation.

Hybridization chain reaction (HCR)

Hybridization chain reaction (HCR)¹²⁵ was performed as described previously¹²⁶ with modifications to initial steps to account for embryonic as opposed to ovarian tissue. These probes were used in the study: *eGFP*, *lacZ*, *grh*, *sna*, *htl*, *CadN*, *HLH54F*, *FoxL1*, *Blimp-1*, *Doc2*, *kon*, *Grip*, *run*, *hkb*, *fkh*, *cad*, *sog*, *DNasell*, *dhd*, *bnk* and *sim*.

QUANTIFICATION AND STATISTICAL ANALYSIS

Read processing, filtering, doublets removal and visualization

Illumina sequencing reads (294 million raw read pairs) were mapped to customized *Drosophila* dm6-transgene reference and quantified using Cell Ranger v7.0 (10x Genomics). A total of 9,845 single cells were recovered from the Cell Ranger pipeline with a mean value of 29,901 reads per cell and a median value of 2,910 genes and 15,251 unique molecular identifiers (UMIs) per cell. The scRNA-seq output matrix was imported to Seurat v3.2.0¹¹⁸ for analysis. Cells with more than 250 but less than 100,000 unique molecular identifiers (UMIs), less than 7000 unique genes, and less than 10% ribosomal reads were kept for downstream analysis (9834 cells). Gene expression counts were log normalized, and log transformed.

Standard data processing was performed as recommended including NormalizeData, FindVariableFeatures (with method set to 'vst'), ScaleData, RunPCA, RunUMAP (with dims set to 1:30), FindNeighbors (with reduction set to PCA and dims set to 1:30). DoubletFinder¹²⁰ was used for doublet detection and removed 707 cells.

After the above filtering step (718 cells were removed), the matrix of a total of 9,127 cleaned single cells was subject to normalization, variable gene identification (method as 'vst', nfeatures as 2000), scaling, and PCA analysis. As PCA elbow plot showed the plateau after the first 20 PCA components, dims were set to 1:20 for FindNeighbors and RunUMAP functions. With resolution set as 0.2 for FindClusters function, a total of 12 unique clusters were identified. Cell cluster marker genes were identified using FindAllMarkers function with the default settings. Top10 markers for each cluster were collected for generating expression heatmap (Table S1).

Data integration and clustering

The standard Seurat integration pipeline¹¹⁸ using Log Normalization method was used to integrate the presented dataset with two other wild-type datasets (repeat live WT, WT#2, and methanol-fixed WT, WT#3, shown in Figure S2, Tables S4, and S5). Before integration, all three WT dataset were filtered using Seurat package v4.1.0¹¹⁹ based on UMIs, unique gene features, and ribosomal reads. Cells with more than 250 but less than 100,000 unique molecular identifiers (UMIs), less than 7000 unique genes, and less than 10% ribosomal reads were kept for downstream analysis. DoubletFinder was used for doublet detection and removed 10% of the cells from the WT repeat and fixed WT datasets. The top 2000 variable gene features (FindVariableFeatures, method = 'vst') in each dataset were used for the SelectIntegrationFeatures function and FindIntegrationAnchors function. The three datasets are then integrated with the IntegrateData function using the LogNormalization method and a default PCA dimension of 1:30. A standard workflow of ScaleData, RunPCA, RunUMAP (with dims set to 1:30), FindNeighbors (with reduction set to PCA and dims set to 1:30) were run on the combined dataset. Finally, the UMAP was visualized with 30 PCs.

The same integration pipeline as described above but with Seurat v3.2.0 was used to integrate the presented dataset with a previously published dataset (GSE202987: GSM6145582)³³ obtained from stage 12 embryos shown in Figure S1 and Table S2. The stage 12 dataset was previously obtained by Seroka et al. through the 10x Chromium platform, sequences aligned through 10x Cell Ranger v3.1, and analyzed using Seurat package v3.1. Before integration, the loaded stage 12 dataset was filtered using Seurat package v3.2.0 based on UMIs, unique gene features, and ribosomal reads. Cells with more than 250 but less than 100,000 unique molecular identifiers (UMIs), less than 7000 unique genes, and less than 25% ribosomal reads were kept for downstream analysis (19465 cells). The filtered presented dataset (filter parameters described above) with 9,127 were used to combine with the filtered stage 12 dataset with 19465 cells. The standard integration workflow as described above was performed with a PCA dimension of 1:20. The UMAP was visualized with 20 PCs.

Gene Ontology enrichment analysis

GO enrichment analysis was performed using the cell signature genes predicted by Seurat. The gene list was analyzed using R version of g:Profiler (<https://biit.cs.ut.ee/gprofiler>)¹¹⁷ for drosophila GO: Biological Pathways and FDR threshold set at 0.05.

Subclustering, differential gene expression, and trajectory analysis

For cell subclustering analysis, the raw gene account matrix of a specific cluster identified in the global analysis was subject to Seurat clustering analysis using the same protocols (20 PCA, 0.5 resolution setting) that have been applied to global analysis. DEG/marker gene analysis (Table S3) was performed using FindAllMarkers function.

Pseudotime trajectory analysis for individual cell clusters was performed with Monocle2^{121–123} following the online tutorial. Top 200 (for analysis with cell clusters 5,6, and 9) and top 500 variable genes (for cluster 1 and 10) within the specific cluster detected by Seurat VariableFeatures function were used as ordering_gene for Monocle2 setOrderingFilter function (Tables S3, S6, and S7). Dimension reduction was performed using the DDRTree method. Pseudotime values of individual cells were projected onto a UMAP plot for visualization.

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Supplemental information

**Single-cell transcriptomics illuminates
regulatory steps driving anterior-posterior
patterning of *Drosophila* embryonic mesoderm**

Jingjing Sun, Chen Zhang, Fan Gao, and Angelike Stathopoulos

SUPPLEMENTAL INFORMATION

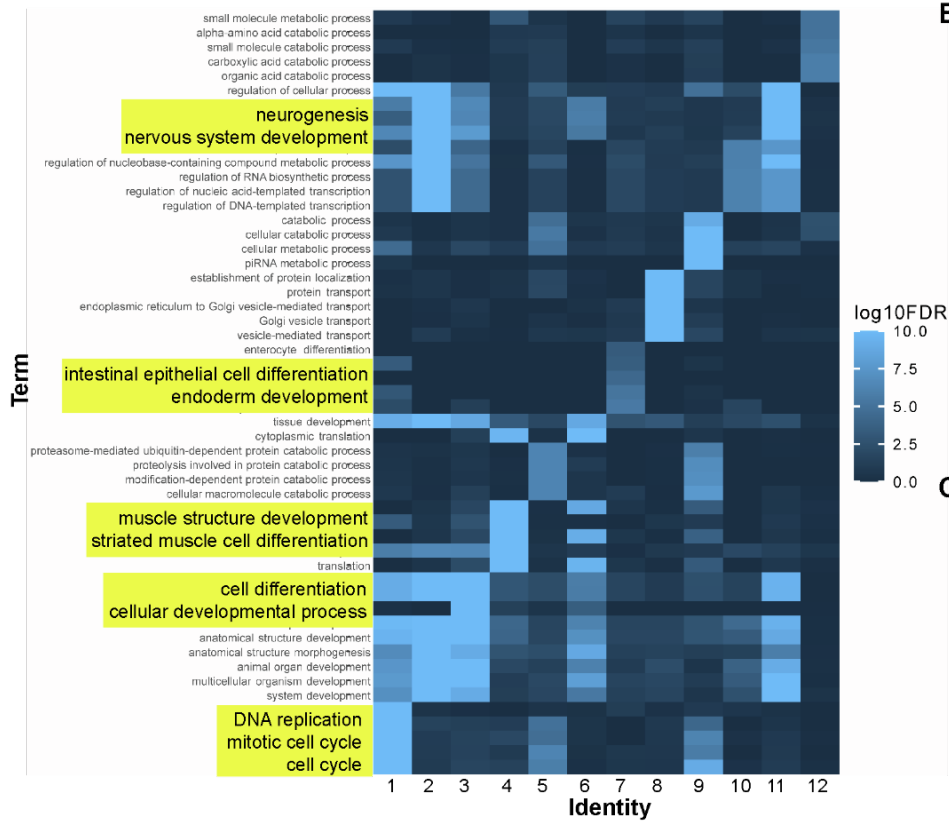
List of Supplemental Materials

Figures S1 – S7

Table S8

Supplemental Reference List

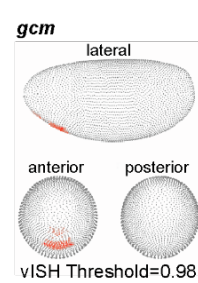
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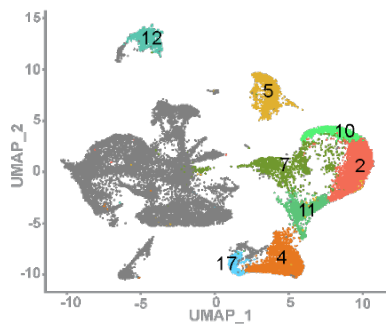
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Identity	Freq
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3	1175
4	859
5	792
6	718
7	656
8	541
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10	233
11	224
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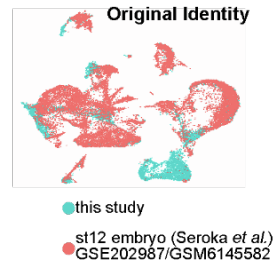
C



D



E



F

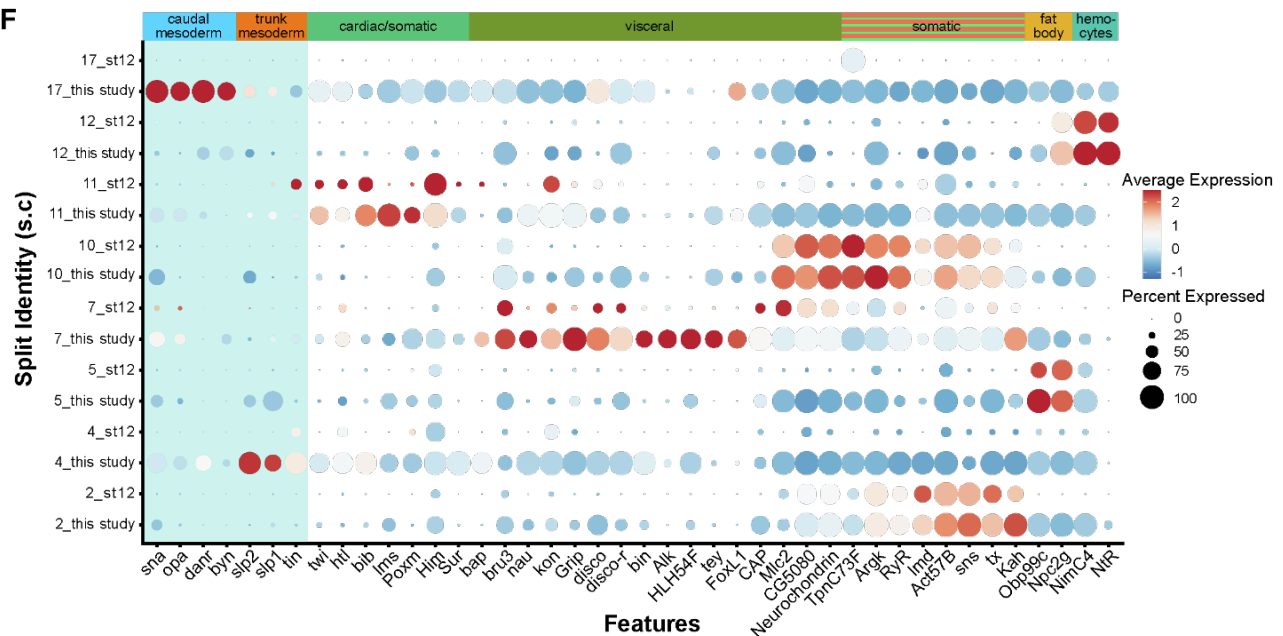


Figure S1. Single-cell transcriptomes of gastrulating *Drosophila* embryos. Related to Figure 1, Tables S1 and S2.

(A) Gene ontology (GO) enrichment analysis of marker genes associated with each cluster.

(B) Number of cells in each cluster after removing the doublets and low-quality cells.

(C) Predicted expression pattern of *gcm* at stage 6 according to DVEX^[S1] (<https://shiny.mdc-berlin.de/DVEX/>).

(D-F) Integration of the single cell transcriptomic data of gastrulation from this study with published data from stage 12 embryos^[S2]. UMAP plot showing the mesodermal cell lineages s.c2, 4, 5, 7, 10, 11, 12, 17 (D) of the integrated datasets: (i) this study live wild-type (WT) #1 and (ii) st12 embryo from Seuroka *et al.* (see Methods). Contribution of integrated datasets to individual cell clusters: cells from this study in blue vs. cells from st12 published dataset in red (E). Side by side visualization of marker gene expression in mesoderm derived cell clusters from the integrated datasets by dotplot (F). Expression levels (normalized among the mesoderm cells) and percentage of cells in the cluster (y-axis) that express each selected marker (x-axis) are shown. Cells that express early caudal and trunk mesoderm genes are associated with s.c4 and s.c17 and predominantly identified by this study (F; see also D,E).

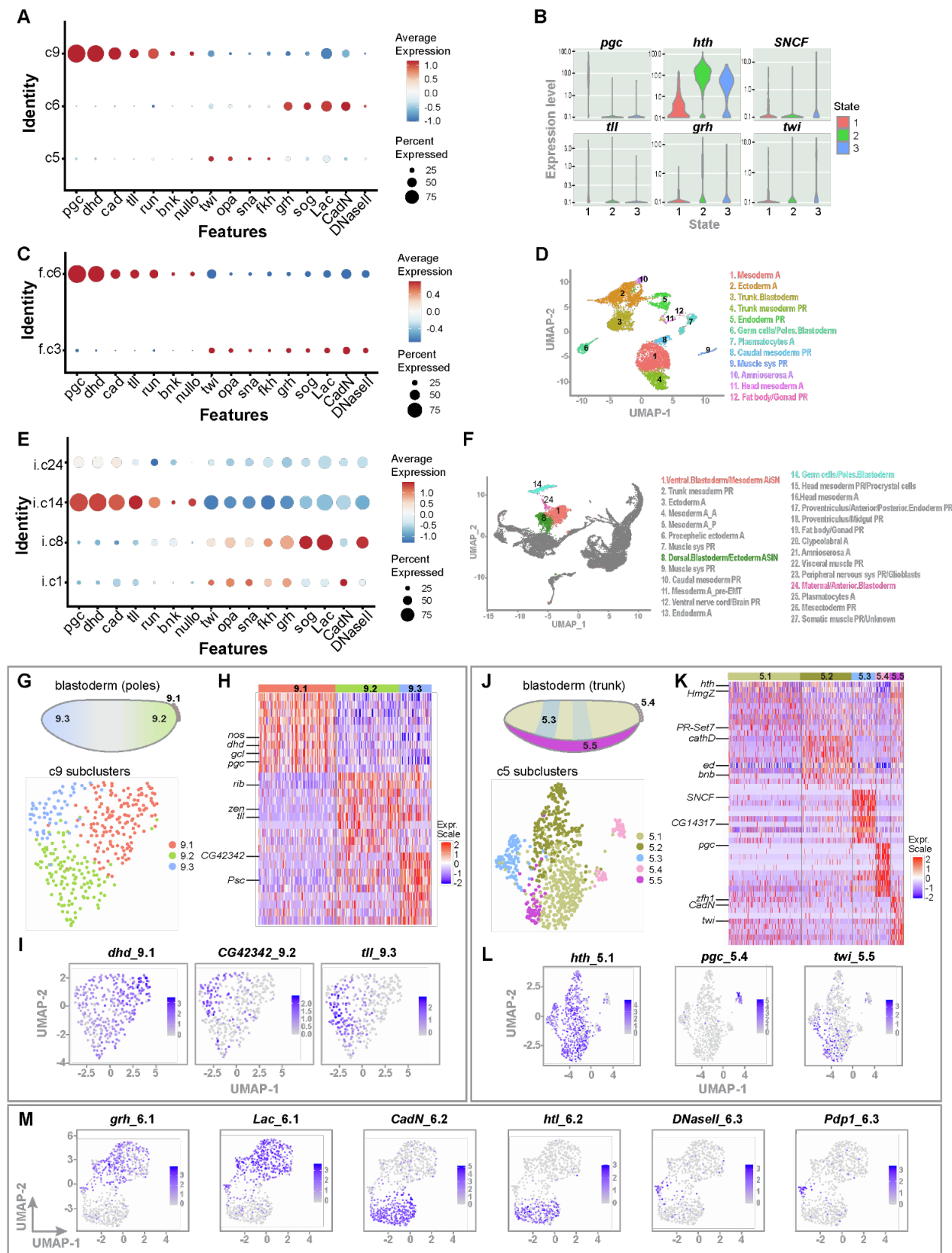


Figure S2. Featured gene expression during nc14. Related to Figure 2 and Tables S3-5.

(A) Dotplot showing representative gene expression in the cell clusters associated with three blastoderm-associated cell clusters of live WT#1 dataset (A).

(B) Violin plots showing expression of additional markers associated with the three cell states generated by trajectory analysis of c9, 5, 6 (Related to Figure 2E,F).

(C,D) Single cell transcriptomic profiling of methanol-fixed cells isolated from stage 5-10 embryos identified 12 cell clusters (D) that largely overlap with those from the live cells (A; live WT#1 shown in main text figures). (C) Dotplot showing expression of the same markers in the two blastoderm-associated cell clusters of fixed cell experiment (fixed WT#3 clusters: f.c3 and f.c6).

(E,F) Four blastoderm-associated cell clusters emerge (i.c1, i.c8, i.c14, i.c24) as a result of integrating three independent single-cell datasets: live WT#1, live WT#2, and fixed WT#3 (E; see Methods). Violin plot showing the trend of representative gene expression in the 27 cell clusters identified by PC analysis in which blastoderm cell clusters are highlighted (F).

(G-I) UMAP plot (G) and heatmap (H) of c9 cells from live WT#1 data (main text figures), which are divided into three subclusters (c9.1, c9.2 and c9.3). Diagram illustrates a blastoderm embryo with color-coded domains labeled with associated subclusters. (I) Subclustering UMAP plots of selected gene expression.

(J-L) UMAP plot (J) and heatmap (K) of c5 cells from live WT#1. Five subclusters (c5.1-5.5) are identified. Color-coded domains of the schematic blastoderm embryo are labeled with associated subcluster identity. (L) Subclustering UMAP plots of selected gene expression.

(M) Subclustering UMAP plots showing selected marker gene expression associated with the three subclusters identified in c6 from live WT#1 (see Figure 2K).

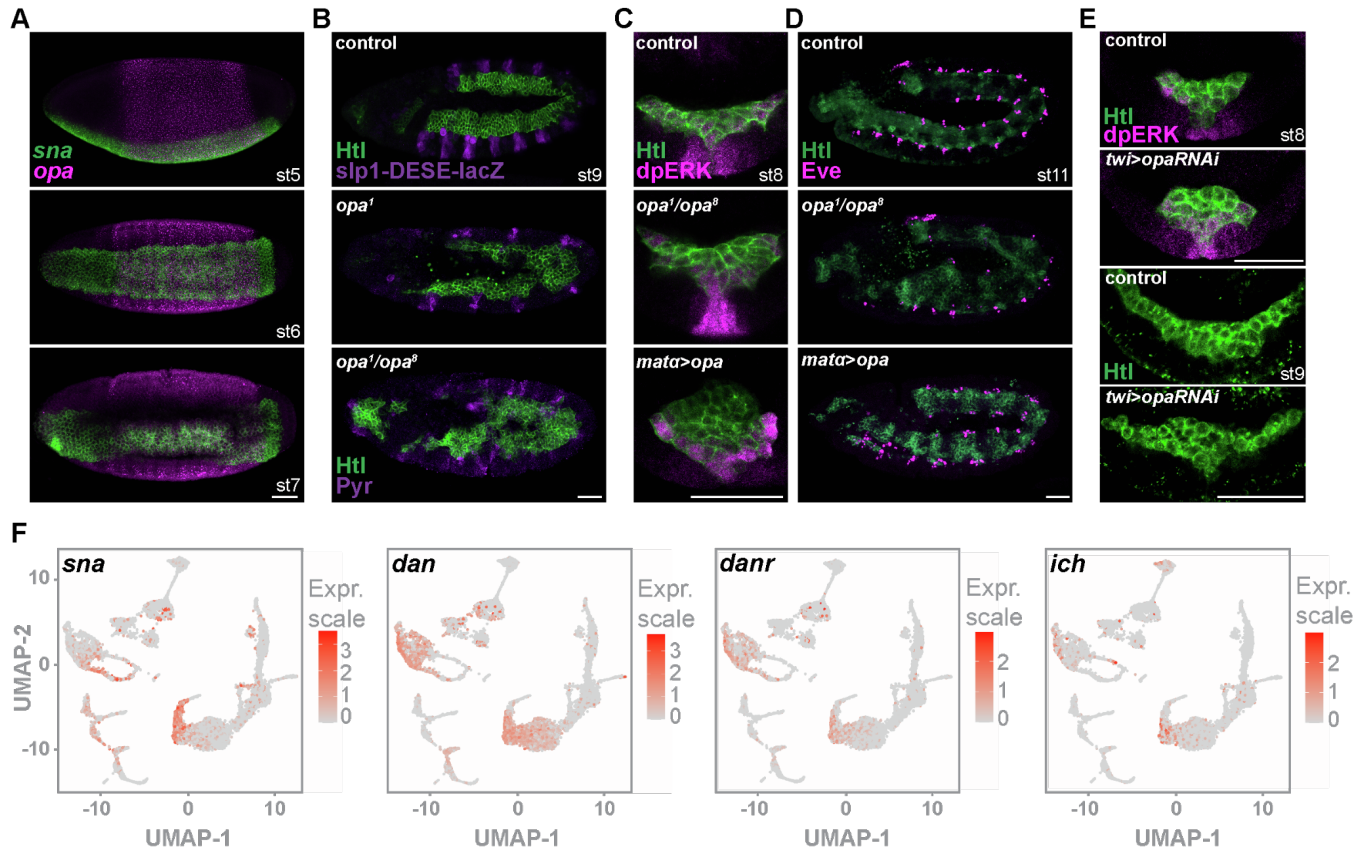


Figure S3. Pre-EMT expression of Opa contributes to mesoderm patterning and lineage differentiation and overexpression of *opa* in the mesoderm disrupts its normal development. Related to Figure 3.

(A) Detection of *opa* transcripts (purple) in the mesoderm (labeled by *sna* probe in green) during invagination.

(B-E) Mesoderm patterning associated with dysregulated *opa* expression. Whole mount side views and Cross sections showing spreading and differentiation defects in *opa* loss of function and overexpression mutants (B-D). Mesoderm cells are labeled with an anti-Htl antibody (green), dpERK antibody staining (purple, C) was used to assay active RTK signaling and Eve antibody (purple, D) to mark the pericardial cells. *opa* mutants exhibit uneven distribution of mesoderm cells and reduced numbers of *slp1* stripes as indicated by β -Galactosidase staining (dark purple, B). Spreading is asymmetric and a delay in spreading is observed in *mata>opa* embryos. The mesoderm spreading defect in these mutants leads to a reduction in the number of pericardial cells at stage 11. When *opa* is specifically knocked-down in the mesoderm by *twi(2xPE)GAI4* (see Methods), only minor mesoderm spreading defects were observed at st8-9 (E), and pericardial cell distribution appears normal.

(F) UMAP plots of *sna*, *dan*, *danr* and *ich* with red intensity indicating the relative expression levels. Genotypes and developmental stages are as indicated. Scale bars=50 μ m.

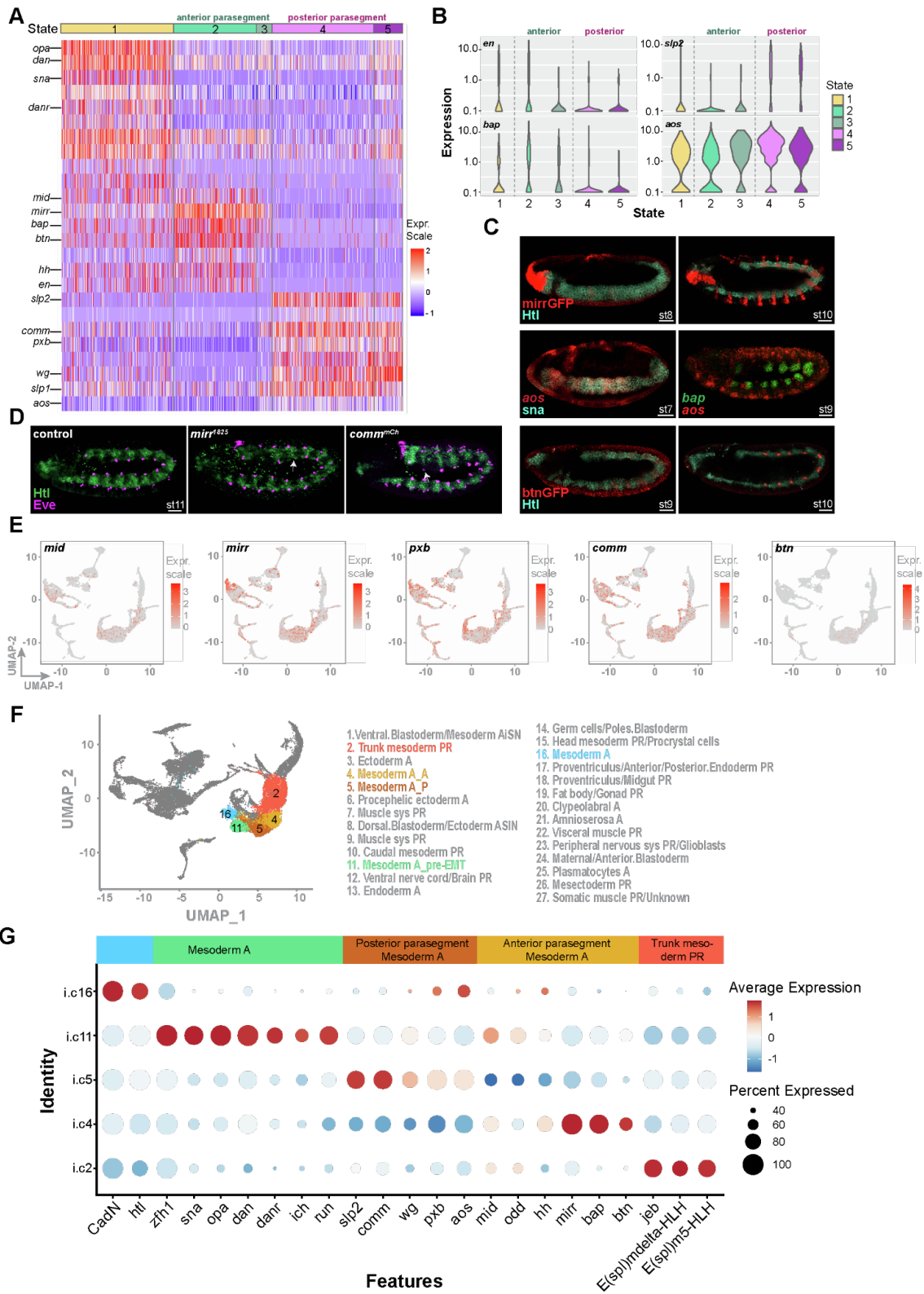


Figure S4. Trajectory analysis of c1 mesoderm cells illuminates patterning along the A-P axis. Related to Figure 4, Tables S5 and S6.

(A-B) Trajectory analysis generates a heatmap showing the top ten marker genes of each five cellular states (A). Violin plots of additional marker gene expression are shown in (B).

(C) Antibody staining of GFP-tagged *mirr* and *btn* or in situ hybridization for endogenous *aos* and *bap* to confirm their segmented expression patterns in st7-10 embryos.

(D) Expression of *mirr* and *comm* in the mesoderm is required for proper cardiac progenitor cells specification. Arrows indicate the missing and extra Eve-positive cell cluster in the *mirr* and *comm* mutant embryos.

(E) UMAP plots of selected markers with red intensity indicating relative expression levels.

(F,G) Five cell clusters (i.c2, 4, 5, 11, 16) relating to the trunk mesoderm are highlighted in the UMAP of integrated single cell datasets (F). Expression of mesoderm markers are shown in the dotplot (G). Three clusters from this integrated dataset (i.e. i.c4, 5, 11) are equivalent to c1 of live WT #1 (the dataset focused on in this study).

Scale bars=50µm.

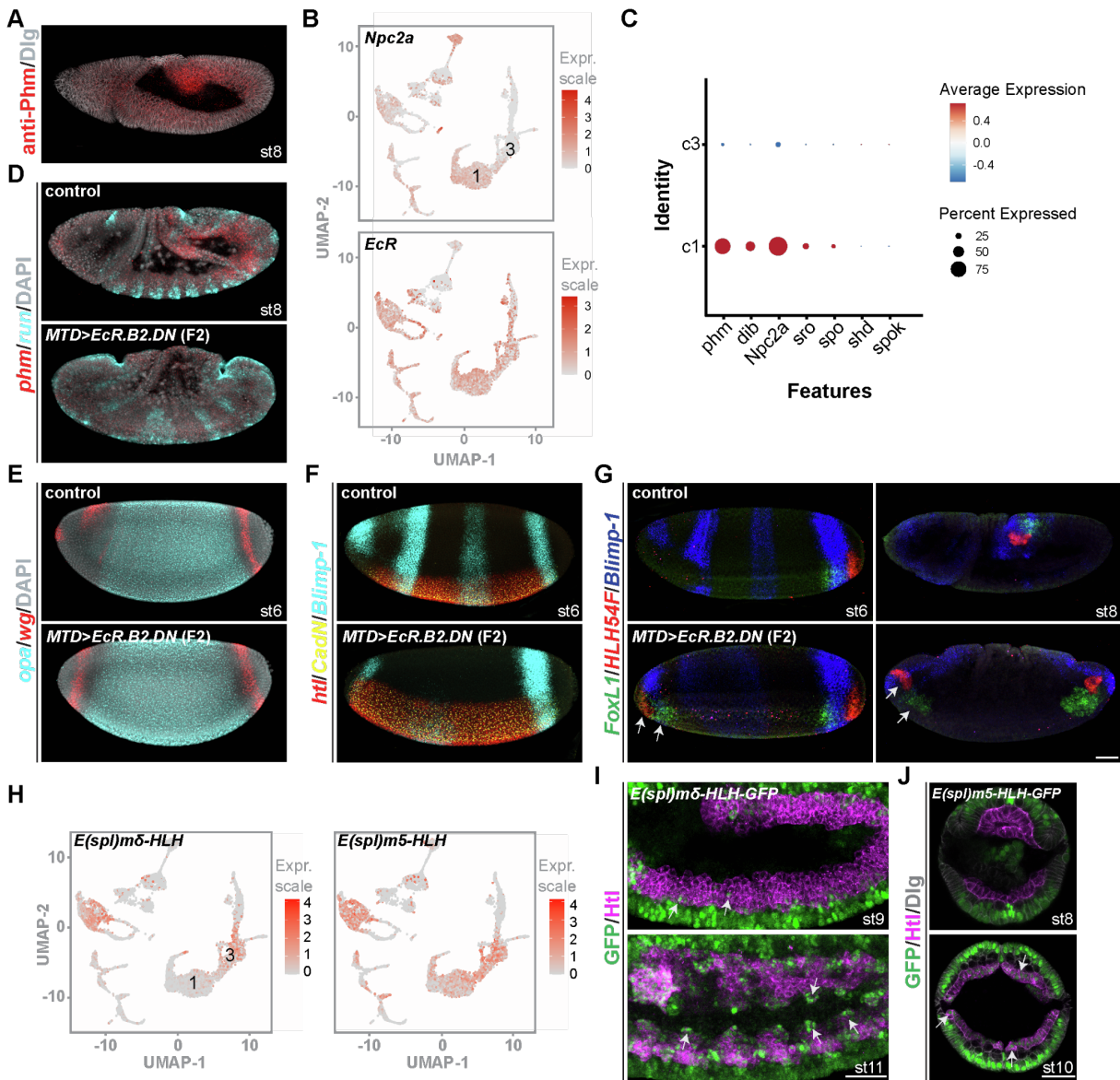


Figure S5. Patterning defects in maternal EcR dominant negative mutants and upregulation of Notch signaling targets post EMT. Related to Figure 5.

(A) Phm (red) and Dlg (white, labeling basolateral cell membranes) antibody staining of wild-type embryo showing Phm is expressed in the mesoderm at st8.

(B) Global UMAPs of *Npc2a* and *EcR*.

(C) Dotplot showing selected Halloween gene expression in mesoderm cell clusters c1 and c3.

(D-G) Defects in AP patterning are observed in F2 second generation offspring when a dominant negative form of EcR is maternally-expressed in the ovary using *MTD-GAL4*. *phm* is downregulated, and a posterized anterior domain is visible in the *MTD>EcR.DN* (F2) mutant (D). (E-G) Disrupting EcR function maternally leads to the anterior domain of embryos adopting gene expression patterns resembling the posterior domain (E), without affecting D-V patterning and mesoderm specification (F). Duplication of the posterior *wg* stripe (red) and slight anterior expansion of *opa* trunk expression (light blue) are seen in the mutant (E). Furthermore, in these mutants,

the anterior and central *Blimp-1* stripes are lost (cyan, F; see also dark blue, G). Duplicated expression of the posterior trunk mesoderm genes *FoxL1* (green) and *HLH54F* (red) is also associated with maternal expression of *EcR.DN* (arrows, G).

(H-J) *E(spl)* complex genes are upregulated in c3 post-EMT mesoderm (H), as confirmed by immunostaining of GFP-tagged *mδ-HLH* (arrows, I, side view of whole-mount embryos are shown) and *m5-HLH* (arrows, J cross-sections are shown).

Combinations of genes and developmental stages assayed by in situ hybridization are as indicated. Scale bars=50μm.

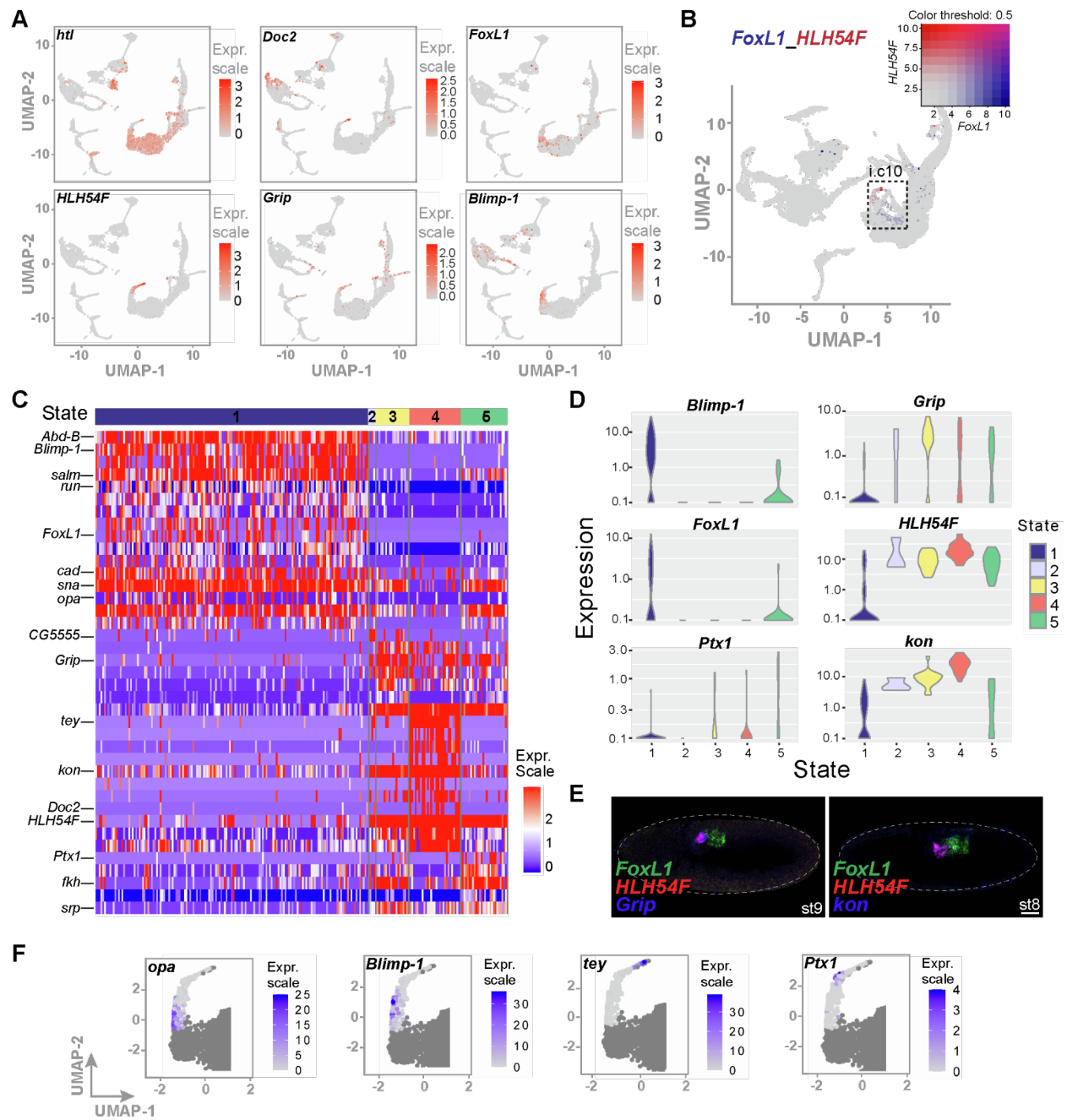


Figure S6. Transcription programs that specify the posterior trunk mesoderm. Related to Figure 6 and Table S7.

(A) UMAPs of selected marker genes in c10.

(B) UMAP plot showing expression of *FoxL1* (blue) and *HLH54F* (red) simultaneously in the integrated dataset (live WT#1, live WT#2 and fixed WT#3). Caudal mesoderm cluster i.c10 is marked by the dotted line box. Expression scale showing to the right.

(C-F) Trajectory analysis of c10 identifies five different cellular states. Heatmap of the top 10 marker genes for each state (C). Violin plots of selected markers with colors labeling different cellular states (D). Consistent with the trajectory analysis, *Grip* and *kon* (blue) are co-expressed with *HLH54F* (red) in the CVM precursors while distinct

from the *FoxL1*-expressing cells (green) at st8-9 (E). Cropped UMAP plots for selected cell state markers showing *opa* and *Blimp-1* (both state 1), *tey* (state 4) and *Ptx1* (state 5) occupy distinct space within c10 (F). Scale bar=50μm.

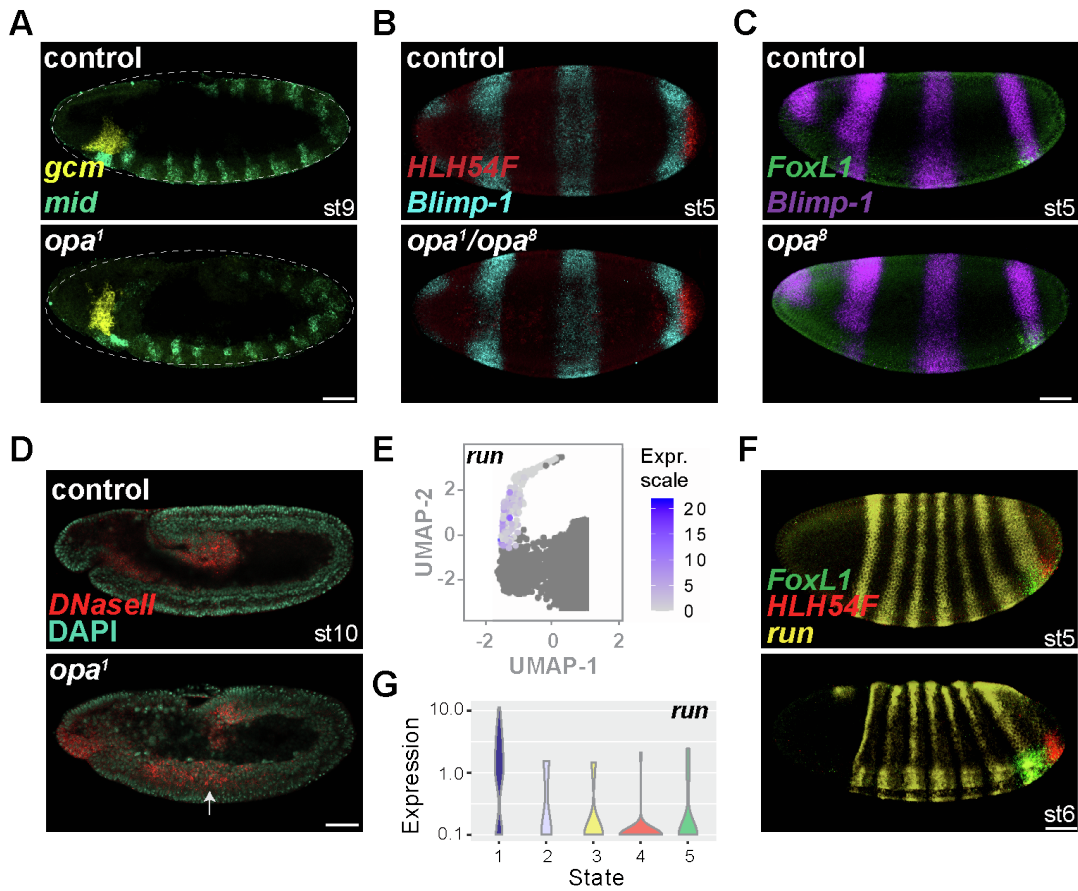


Figure S7. *opa* functions together with other factors in the trunk to spatially restrict gene expression to the head and caudal mesoderm. Related to Figure 7.

(A-C) Loss of *opa* does not lead to obvious changes in *gcm* (A, yellow), *HLH54F* (B, red) and *FoxL1* (C, green) expression.

(D) Ectopic expression of *DNaseII* (red) is observed in the trunk in the *opa*¹ mutant (arrow, D) suggesting Opa (directly or indirectly) normally supports repression of this gene from the trunk.

(E-G) Pair rule gene *run* is enriched in state 1 cells from trajectory analysis (G) within c10 (E). Cells comprising c10 state 1 also are associated with *FoxL1* transcripts (see Figures 6B, S6D). This result and the overlapping expression of *run* and *FoxL1* at stage 5 detected by *in situ* hybridization using HCR probes (F) suggested a role for Run in supporting *FoxL1* expression.

Genes detected and stages are as labeled. Scale bars=50µm.

Table S8, List of oligonucleotides used in this study, related to key resources table

Oligonucleotides (Table S8)		
D2eGFP RNA probe (HCR)	Molecular Instruments	d21260aa-84e9-4ae2-9adf-109eef0e85eb/PRH935
<i>HLH54F</i> RNA probes (HCR)	Molecular Instruments	4358/E576
<i>FoxL1</i> RNA probes (HCR)	Molecular Instruments	fd9e7d98-87b2-46e2-967a-a384167b8da6/PRR051
Blimp-1 RNA probes (HCR)	Molecular Instruments	fd9e7d98-87b2-46e2-967a-a384167b8da6/PRR053
<i>Doc2</i> RNA probes (HCR)	Molecular Instruments	5c204473-08ca-4546-bf24-1925b8003083/PRO669
<i>kon</i> RNA probes (HCR)	Molecular Instruments	4049/E192
<i>Grip</i> RNA probes (HCR)	Molecular Instruments	9d28cc4d-0e82-4330-9564-96b40a58248c/RTB563
<i>hkb</i> RNA probes (HCR)	Molecular Instruments	9d28cc4d-0e82-4330-9564-96b40a58248c/RTB565
<i>run</i> RNA probes (HCR)	Molecular Instruments	9d28cc4d-0e82-4330-9564-96b40a58248c/RTB564
<i>cad</i> RNA probes (HCR)	Molecular Instruments	d5cdac09-6920-42eb-b229-b5d2421b995c/RTB831
<i>tll</i> RNA probes (HCR)	Molecular Instruments	d5cdac09-6920-42eb-b229-b5d2421b995c/RTB299
<i>grh</i> RNA probes (HCR)	Molecular Instruments	6bb0a64d-d4b8-4b1a-a83b-7d76fe5d9a77/PRR869
<i>sna</i> RNA probes (HCR)	Molecular Instruments	4614/E928

<i>CadN</i> RNA probes (HCR)	Molecular Instruments	6bb0a64d-d4b8-4b1a-a83b-7d76fe5d9a77/PRR868
<i>htl</i> RNA probes (HCR)	Molecular Instruments	6bb0a64d-d4b8-4b1a-a83b-7d76fe5d9a77/PRB870
<i>sim</i> RNA probes (HCR)	Molecular Instruments	9d28cc4d-0e82-4330-9564-96b40a58248c/RTB560
<i>lacZ</i> RNA probes (HCR)	Molecular Instruments	d5cdac09-6920-42eb-b229-b5d2421b995c/RTB830
<i>bnk</i> RNA probes (HCR)	Molecular Instruments	5fa8f306-142e-4909-8555-3a57ae81b8a5/RTF902
<i>dhd</i> RNA probes (HCR)	Molecular Instruments	5fa8f306-142e-4909-8555-3a57ae81b8a5/RTF901
<i>DNaseII</i> RNA probes (HCR)	Molecular Instruments	94ed6100-de8d-4db1-82ca-3d3a69c9fb59/RTF683
<i>sog</i> RNA probes (HCR)	Molecular Instruments	3285/C529
<i>bap</i> RNA probe	Trisnadi and Stathopoulos ^[S3]	N/A
<i>aos</i> RNA probe	Trisnadi and Stathopoulos ^[S3]	N/A
ATGATGATGAACGCCTTCAT	This study	opa-F
AAGTAATACGACTCACTATAGGGA GATGTATGCGTCCGCTTGTGGA	This study	opa-R
AATCACTGGCGGTCGAACAA	This study	run:FP1
TAATACGACTCACTATAGATACGG ATGGAAGGCGTGTG	This study	run:RP1T7
ATTAACCCTCACTAAAGGGAGCAG ATCCCTATCTTCACGC	This study	byn-F
TAATACGACTCACTATAGGTGGGA TAGGAGAACACGGAC	This study	byn-R
AGGAGCGCGAGACAAGTCT	This study	odd-F

AAGTAATACGACTCACTATAGGGA GATGTATGCGTCCGCTTGTGGA	This study	odd-R
TAGCGCTCACAGTCTCCACTTTCA	This study	slp2-F
AAGTAATACGACTCACTATAGGGA GACCACCGAGAGTTGCCTTTGAT	This study	slp2-R
CGATTAAGACGCACCACAGTTC	This study	oc-F
TAATACGACTCACTATAGCATATTC GCGTACTTATCCTGC	This study	oc-R
CCTTTACAACTGCCCAAGC	This study	DNaseII-F
TAATACGACTCACTATAGTTGTCTG TAGTTGGTGACCAG	This study	DNaseII-R
GCAATGGTCGCTTGGAATC	This study	gcm-F
TAATACGACTCACTATAGCTGTTG ACTGGGTGATATG	This study	gcm-R
GCTATCCCATGACGCGTTCG	This study	pxb-F
TAATACGACTCACTATAGCTTGGT TTCGGTTGGGTAGC	This study	pxb-R
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GACTGTTACGCAGATTGTTCG	This study	mirr-F
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CTGTGCGTCGGAAGTGAATC	This study	phm-F
TAATACGACTCACTATAGACAACC TAATCCTGCACCAC	This study	phm-R

SUPPLEMENTAL REFERENCES LIST

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