Temporal Coordination of Collective Migration and Lumen Formation by Antagonism between Two Nuclear Receptors

HIGHLIGHTS
- E75 antagonizes DHR3’s function in inducing lumen formation of border cells (BCs)
- E75 and DHR3 temporally coordinate collective migration and lumen formation of BCs
- DHR3 is required and sufficient for chitin secretion into the lumen
- DHR3 and βFtz-f1 downregulate JNK signaling and cell adhesion in the BCs

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Temporal Coordination of Collective Migration and Lumen Formation by Antagonism between Two Nuclear Receptors

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SUMMARY
During development, cells undergo multiple, distinct morphogenetic processes to form a tissue or organ, but how their temporal order and time interval are determined remain poorly understood. Here we show that the nuclear receptors E75 and DHR3 regulate the temporal order and time interval between the collective migration and lumen formation of a coherent group of cells named border cells during Drosophila oogenesis. We show that E75, in response to ecdysone signaling, antagonizes the activity of DHR3 during border cell migration, and DHR3 is necessary and sufficient for the subsequent lumen formation that is critical for micropyle morphogenesis. DHR3’s lumen-inducing function is mainly mediated through βFtz-f1, another nuclear receptor and transcription factor. Furthermore, both DHR3 and βFtz-f1 are required for chitin secretion into the lumen, whereas DHR3 is sufficient for chitin secretion. Lastly, DHR3 and βFtz-f1 suppress JNK signaling in the border cells to downregulate cell adhesion during lumen formation.

INTRODUCTION
During development, a group or population of cells often has to undergo multiple, distinct morphogenetic processes in a certain temporal order (e.g. A, then B...) to form a tissue or organ (Webb and Oates, 2016). If the correct temporal order is not followed (e.g. process B occurring before process A), that tissue or organ would not form correctly (Rougvie, 2001; Thummel, 2001). Besides the correct order, the time interval between two processes is another important aspect of the temporal control for the morphogenetic processes. Making the interval too long or too short would also be detrimental to the formation of the organ or tissue. Despite their importance in development, our current understandings on how the temporal order and time intervals are regulated and determined are very limited.

The somatic follicle cells of the Drosophila egg chamber have served as an excellent model system to study multiple morphogenetic processes (Horne-Badovinac and Bilder, 2005). Specifically, during stage 9 of oogenesis, a group of about eight cells detaches from the anterior follicle epithelium and undergoes collective migration between the germ-line nurse cells in a posterior direction (Montell, 2003). By early stage 10A, this coherent cluster of cells would have migrated a distance of about 150 μm in 6 h, reaching the border between oocyte and nurse cells, hence the name border cells. About 6 h later, by stage 10B, the cluster of eight border cells would have migrated dorsally a short distance along the border, eventually stopping at the dorsal-most position of the border. Four hours later, by stage 12, this border cell cluster undergoes a second morphogenetic process to eventually form the tip of micropyle, a tubular structure required for sperm entry into the mature oocyte (Montell et al., 1992). Therefore, the formation of micropyle tip by border cells requires two distinct morphogenetic processes in a certain temporal order: first, the well-studied, stereotyped, collective migration process and then a largely uncharacterized morphogenetic process that transforms these border cells into the tip of the tubular structure. Furthermore, an interval of about 16 h exists between the beginning of collective migration and the start of the micropyle formation. However, whether and how the temporal order and the time interval between the two morphogenetic processes are regulated remain largely unknown.

Previous studies have shed light on the temporal regulation of border cell migration. The steroid hormone ecdysone, its receptor heterodimer ecdysone receptor (EcR) and ultraspiracle (USP), and their co-activator

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Taiman (Tai) had all been shown to be required for the initiation of border cell migration (Bai et al., 2000; Cherbas et al., 2003; Jang et al., 2009). Ecdysone and the EcR signaling had long been known to play important roles in coordination of growth and developmental timing during embryogenesis, larval molting, and metamorphosis in Drosophila (Jia et al., 2017; Kozlova and Thummel, 2003; Yamanaka et al., 2013). Active form of ecdysone is also made in the adult Drosophila ovaries to regulate progression of oogenesis (Ables et al., 2016; Buszczak et al., 1999; Carney and Bender, 2000). 20-Hydroxyecdysone, the active form of ecdysone, is locally synthesized by the follicle epithelium in individual egg chambers and reaches its highest levels around stages 9 and 10 (Domanitskaya et al., 2014; Margaret et al., 1989). Even small patches of wild-type follicle cells in mosaic stage 9 egg chambers were shown to produce a sufficient level of active ecdysone that allows the border cells to begin migration (Domanitskaya et al., 2014). The sufficiency of ecdysone/EcR signaling on initiation of border cell migration was further demonstrated by Jang and co-workers, in which early expression of the activated form of the co-activator Tai can precociously initiate border cell migration (Jang et al., 2009). However, what cellular processes in the border cells are directly regulated by EcR signaling and whether EcR also temporally regulates micropyle formation are currently unknown.

In this study, we show that E75 and DHR3, two nuclear receptors/transcription factors downstream of EcR signaling, regulate both the temporal order and time interval between border cell migration and micropyle formation. We find that EcR's migration-initiating ability is partially through E75's function. During border cell migration, EcR signaling activates the expression of both E75 and DHR3, with E75 repressing DHR3's function. Furthermore, de-repression of DHR3 function after completion of border cell migration switches on lumen formation, turning the cluster of border cells into the tip of micropyle. Such antagonist relationship between E75 and DHR3 (while both under the control of EcR signaling) provides the regulatory mechanism of temporal order and time interval between two distinct morphogenetic processes essential for the formation of a functional micropyle.

RESULTS

RNAi Screen Identifies E75 Acting downstream of EcR Signaling

Ecdysone signaling was known to be critical for the temporal control of initiation of border cell migration (Bai et al., 2000; Jang et al., 2009), but the cellular processes directly regulated by ecdysone/EcR signaling are largely unknown. To identify them, we carried out a small-scale RNAi screen of candidate genes that were previously reported to be responsive to ecdysone in Drosophila larvae and pupae and in cell lines (Beckstead et al., 2005; Gauhar et al., 2009; Sap et al., 2015). We first screened through the well-established response genes of ecdysone signaling (Ashburner and Richards, 1976; Huet et al., 1995; Richards, 1995), including E74, E75, E93, Br-c, and DHR3. Two to three different RNAi lines were used to confirm that phenotypes were not due to off-target effects, and two RNAi lines for the EcR gene were used as positive controls. A border-cell-specific Gal4 driver, slbo-Gal4, was used to drive expression of various RNAi transgenes in border cells beginning at late stage 8 of oogenesis, before border cells initiate their migration at early stage 9. As expected, both EcR RNAi lines (9327 and v35078 lines) resulted in phenotypes of strong migration delay or block, consistent with the previous reported roles of EcR in initiating and promoting border cell migration (Figures 1A, 1B, and S1A–S1C) (Bai et al., 2000; Jang et al., 2009). In comparison, border cell clusters from the control (wild type) stage 10 egg chambers almost always reached the 100% migration position, with only 6% of clusters displaying moderate delay (stopping at the 75% position) (Figures 1B and S1A–S1C). Interestingly, of all the five ecdysone response genes tested, only E75 displayed strong migration defects. In fact, all three RNAi lines (v44851, 26717, Thu1738) consistently resulted in severe migration block and delay phenotypes as well as dramatic redistribution of F-Actin, when compared with the control (Figures 1A, 1B, and S1C). We then screened an additional collection of 20 genes that were considered ecdysone response genes or putative target genes of EcR/USP in previous reports (Beckstead et al., 2005; Gauhar et al., 2009; Li and White, 2003). However, none of the genes, when knocked down, displayed strong migration defects (Figures S1A–S1C). Only mild-to-moderate migration phenotypes were observed in a few of the RNAi experiments.

We then proceeded to determine whether E75 acts downstream of EcR to initiate and promote border cell migration. Three distinct isoforms of E75 (A, B, C) were shown to be involved in different developmental and cellular processes (Huet et al., 1995; Li et al., 2016; Terashima and Bownes, 2004). Both E75A and E75C contain ligand-binding domain and DNA-binding domain, whereas E75B contains only the ligand-binding domain and no DNA-binding domain (Bialecki et al., 2002; Segraves and Hogness, 1990). E75B
Figure 1. E75 Antagonizes DHR3 during Border Cell Migration

(A) Confocal images of egg chambers stained with phalloidin (red, for F-actin) and DAPI (blue, for nuclei) with indicated genotypes. The boxed regions are enlarged and shown to the right. Border cells expressing EcR RNAi displayed strong migration defects but exhibited a similar morphology and F-actin distribution pattern to that of wild-type (WT) border cells, whereas E75 RNAi border cells with migration defects displayed a very different morphology and F-actin distribution pattern to that of the EcR RNAi and WT border cells. Co-expression of DHR3 RNAi rescued E75 RNAi’s morphology and F-actin defects. Ri is the abbreviation for RNAi for this and all subsequent figures. Posterior is to the right and anterior is to the left for this and all subsequent figures.

(B) Quantification of border cell migration with indicated genotypes. EcR-Ri denotes EcR RNAi, “+” indicates that these genotypes include both EcR RNAi and one of the denoted genotypes (UAS-GFP, UAS-E75A, UAS-E75B, and UAS-E75C). The “+” below E75 RNAi indicates that these genotypes include both E75 RNAi and one of the denoted genotypes (UAS-GFP, DHR3 RNAi). The x axis denotes the percentage of stage 10 egg chambers examined for each genotype that exhibited various degrees of migration, as represented by the five color-coded bars (see Figure S1 for details).

(C and E) Confocal images displaying β-galactosidase (β-gal) staining (C) and DHR3 staining (E) of stages 8, 9, and 10 egg chambers. Boxed region is enlarged to the right, showing a high-magnification view of the border cells.
can heterodimer with DHR3 and inhibit its activity during metamorphosis (White et al., 1997). The sequences used in the three RNAi lines for E75 are all within the common region and would have knocked down all three isoforms. Therefore, we overexpressed each isoform to test its individual rescue ability on border cell migration defects as caused by EcR RNAi. We found that E75B overexpression markedly rescued EcR RNAi’s migration defects, whereas E75C displayed a much weaker rescue effect and E75A showing no significant rescue (Figure 1B). Moreover, we found that E75’s transcription levels (as represented by a previously used reporter E75-lacZ (Manning et al., 2017)) within border cells at stages 9 and 10 were much higher than those at stage 8 (Figure 1C), consistent with ec dysone signaling being significantly increased beginning at stage 9. And mosaic border cell clusters containing a clone of EcR RNAi-expressing cells demonstrate that E75B protein levels are drastically decreased when EcR function is reduced (Figure 1D), indicating that EcR activity is required for E75B expression during stage 9. Taken together, these results indicate that E75B is likely the major isoform and downstream factor to mediate EcR’s temporal control on border cell migration. Other downstream factors besides E75B may also mediate ec dysone/EcR signaling because E75B can only partially rescue EcR’s loss-of-function phenotype in border cells. Consistently, a recent study using microarray analysis also identified E75 as one of the target genes that are responsive to ec dysone signaling in the migratory border cells (Manning et al., 2017).

E75 Antagonizes DHR3’s Function during Collective Migration of Border Cells

During metamorphosis, ec dysone-activated EcR turns on the expression of E75B, which then binds to DHR3 and antagonizes its activity (White et al., 1997). E75B and DHR3 are both nuclear receptors/transcription factors and are both induced by ec dysone, and E75B’s inhibition of DHR3 function leads to suppression of DHR3’s transcriptional activation of its target genes essential for metamorphosis (Caceres et al., 2011; Reinking et al., 2005; White et al., 1997). To determine whether antagonistic interaction also exists between E75 and DHR3 during border cell migration, we co-expressed DHR3 RNAi and E75 RNAi in border cells. We found that DHR3 reduction strongly rescued E75 RNAi’s migration defects (Figures 1A and 1B), as well as the morphological defects of border cells (Figure 1A, also described in the section below). On the other hand, overexpression of DHR3 resulted in similar phenotypes of migration and morphology to those of E75 RNAi (Figures 1G and 1H), only with DHR3 overexpression’s defects more severe than those of E75 RNAi (Figures S3A and S3B). Furthermore, E75 overexpression can in turn suppress DHR3 overexpression’s severe defects (Figures 1G and 1H), with all three of its isoforms (E75A, E75B, E75C) displaying similar suppressing abilities. This is consistent with previous reports that both E75A and E75B isoforms can heterodimerize with DHR3 to inhibit DHR3’s transcription activation ability (Sullivan and Thummel, 2003; White et al., 1997). Furthermore, all three isoforms contain the ligand-binding domain, which was previously shown to be involved in heterodimerization with DHR3 and inhibition of its activity (Reinking et al., 2005). Lastly, we showed that DHR3’s levels were also increased in border cells beginning at stage 9 (Figure 1E), similar to E75’s temporal expression pattern (Figure 1C), and its levels also depended on EcR’s activity (Figure 1F). Together, these data demonstrate an antagonistic relationship between E75 and DHR3 during border cell migration, with both their expressions activated by EcR during the migratory process.

Conversely, we found that expressing E75 RNAi (B26717) or DHR3 in border cells significantly reduced the level of ec dysone response activity or EcR signaling (Figures S2A–S2C and S2F), which is represented by expression levels of EcRRE-lacZ, a common reporter of EcR activity used in previous studies (Jang et al., 2009; Koelle et al., 1991). Moreover, expression of DHR3 RNAi could rescue E75 RNAi’s EcRRE-lacZ levels (Figures S2D–S2F). These data indicate that E75 can exert a positive feedback on EcR signaling by antagonizing DHR3’s inhibitory effect on EcR signaling. This conclusion is consistent with previous studies that showed DHR3 physically interacted with EcR and suppressed its activity (Lam et al., 1997; White et al., 1997). It should be noted that DHR3 could also exert its inhibitory effect on EcR indirectly.
**E75 Antagonizes DHR3’s Function in Lumen Formation during Border Cell Migration**

We noted that border cell clusters with E75 knockdown or DHR3 overexpression displayed different morphology and F-actin staining pattern from border cells with reduced EcR function (Figure 1A), as indicated by our EcR RNAi result and previous reports (Hackney et al., 2007). The delayed border cell clusters with EcR RNAi often displayed a coherent and front-polarized morphology with F-actin enriched in the periphery of the cluster, similar to that of the wild-type clusters (Figure 1A). On the contrary, E75 RNAi- or DHR3-overexpressing border cells (driven by slbo-Gal4) lost the front-polarized cluster morphology that is characteristic of front-back polarity, and F-actin is instead enriched in the center of the cluster in a ring-like structure (Figures 1A and 1G), which is unique and never observed in any of the previously reported mutant phenotypes of border cells (to our knowledge). Closer examination revealed that this unique structure is not within individual border cell’s cytoplasm but is instead composed of portions of outer border cells’ inside membranes, which join together to form a continuous supra-cellular ring (Figures 2A–2E). And a similar but smaller ring can also form between two border cells in a flip-out clone that expresses E75 RNAi by Ay-Gal4 (Figure S3C). Moreover, such supra-cellular structure is also enriched with molecules that are typically associated with apical membranes (aPKC, Crb, Baz/Par3, PIP2-GFP reporter) (Figures 2A, 2C, 2E, and S3C) but not with lateral membranes (Dlg) (Figure 2A). In addition, E-cadherin re-localization was also observed, with E-cadherin sometimes ectopically enriched in a region adjacent to that of aPKC staining (Figure S8D). A typical supra-cellular ring encloses a space that resembles a lumen with significant depth (about 5 μm, Video S1) in the center of cluster, effectively displacing the two central polar cells to the side and underneath. To address whether polar cells are involved in the formation of the lumen, we closely followed the position of polar cells and found that lumens were mostly enclosed by border cells and that the ring-like lumen structures were not derived from the circumference of polar cells (Figures 1A, 1G, 2A, 2C, 2E, and 2B; polar cells marked by *). slbo-Gal4 expressed E75 RNAi or DHR3 only in border cells and not in polar cells, raising the possibility that differential expression at the interface of the border and polar cells could result in the unique lumen phenotype. We then used c306-Gal4 to drive E75 RNAi expression in both border cells and polar cells (Figure S3D), and we found similar lumen phenotype to that resulted from slbo-Gal4, suggesting differential expression of E75 RNAi or DHR3 in border cells and polar cells is not the cause of lumen formation.

The strong and specific enrichment of apical markers such as aPKC in the membranes enclosing the luminal space suggests that the border cell cluster has undergone a lumen formation process to become a tubular structure with the apical membrane facing the central lumen. Interestingly, the E75 RNAi border cells displayed a range of lumen-like phenotypes. Half of them (50.0%) showed a clear lumen phenotype that is similar to that of the DHR3 overexpressing border cells, whereas majority of the rest (39.0%) exhibited little luminal space and discontinuous apical membrane patches as labeled by aPKC (Figures S3A and S3B), which resemble the previously reported structure of pre-apical patches (PAP) that are present during the intermediate stages of de novo lumen formation in several model systems (Bryant et al., 2010; Ferrari et al., 2008; Yang et al., 2013). These moderate phenotypes may reflect incomplete lumen formation or intermediate stages of lumen formation in the border cells, whereas the large lumen structure present in vast majority of the DHR3-overexpressing clusters and half of the E75 RNAi clusters may indicate complete lumen formation.

Formation of a tube and its enclosing lumen from non-epithelial cells is referred to as de novo lumen formation (Sigurbjörnsdóttir et al., 2014), which is a fundamental morphogenetic process central to animal development. Extensive studies in various in vitro and in vivo model systems have revealed that the initial stage of de novo lumen formation involves establishment of a new apical-basal polarity, which requires re-routing of multiple cellular processes and components including polarized intracellular trafficking, polarized actin and microtubule cytoskeleton, polarized distribution of apical markers, and newly synthesized membrane (Akhtar and Streuli, 2013; Datta et al., 2011; Sigurbjörnsdóttir et al., 2014). We found that in addition to the re-distribution of apical markers to the lumen-facing membrane, the intracellular traffic as well as cytoskeleton was also dramatically re-organized in the E75 RNAi- or DHR3-overexpressing border cells. Staining with Rab11 and Sec15 antibodies revealed that recycling endosome and exocyst were enriched in the cytoplasmic regions near the lumen-facing apical membrane, indicating a polarized transport toward the lumen (Figures 2G and 2H). Furthermore, F-actin and sometimes aPKC were observed localizing to large vacuole-like compartment adjacent to the lumen-facing membrane (Figures 2B and 2F), suggesting that these large vesicles could be in the process of fusing with the adjacent apical membrane. This phenomenon was similar to previous reports of VACs (vacular apical compartments) forming in the MDCK.
cells that are undergoing de novo lumen formation (Brignoni et al., 1993; Vega-Salas et al., 1988). In addition, the actin and microtubule cytoskeletons were re-organized in such a way that they are now mostly localized in and adjacent to the lumen-facing membrane. Interestingly, β-tubulin was re-organized into a distribution pattern that seems to radiate away from the central lumen (Figure 2D). Lastly, marked increase
of intracellular membrane levels as indicated by Myr-RFP and PIP2-GFP was observed in the cytoplasm of DHR3 expressing border cells, suggesting that high levels of newly synthesized membrane are needed for formation and expansion of lumen-facing membrane (Figure 2E). Taken together, these results indicate that during border cell migration E75 acts to suppress DHR3’s lumen formation function, which includes re-routing of endocytic recycling, re-distribution of apical markers, re-polarization of actin and microtubule cytoskeletons, and increased levels of membrane components.

DHR3 Is Later Required for the Formation of Micropyle Tip

We next sought to understand why E75 needs to suppress DHR3’s lumen formation function during border cell migration. After border cells finish their anterior migration to the border at stage 10A, they would further migrate a short distance dorsally and finally stop at the dorsal border between nurse cells and oocyte at stage 10B. About four hours later, around stages 12, this cluster of border cells will undergo a morphogenetic transformation to form part of the micropyle, which is a tubular structure essential for sperm entry (Montell et al., 1992). Such a morphogenetic process is not well characterized and understood. Therefore, we wonder whether the lumen-forming phenotype from E75 knockdown or DHR3 over-activation represents the precocious occurrence of the morphogenetic event involved in micropyle formation. If that is the case, E75 may actually be preventing a late morphogenetic process from occurring earlier (i.e. before or during border cell migration). Therefore, E75 and DHR3 may function together to keep the correct temporal order between the two morphogenetic processes. To address this possibility, we first sought to characterize and understand the process that enables wild-type border cell cluster to be transformed into the tip of micropyle.

Collective migration of border cells has been extensively studied, but the morphogenetic process that turns the border cells into micropyle tip is little studied. Previous work by Montell and coworkers first demonstrated that border cells develop into the tip of micropyle and contribute to the cellular process thought to maintain a functional opening, whereas the centripetal follicle cells form the bulk of the micropyle structure. Furthermore, in the absence of border cells, a slightly smaller micropyle structure could still form, but it lacks the functional opening required for sperm entry (Montell et al., 1992). We sought to describe and characterize such morphogenetic process in details, using markers of actin cytoskeleton, membrane, and apical polarity (Figures 3A–3D, Videos S2, S3, and S4). Similar to migratory border cells at stage 9, border cells at stage 10 (a period of about 10 h, Figure 3A) mostly retain the coherent cluster morphology as well as the distribution pattern of F-actin and apical polarity proteins. During stages 9 and 10, Par6-GFP was shown to localize between adjacent border cells in a thin section of junctional region (Figure 3B, Video S2), which was subsequently retracted and significantly shortened during stage 11 (a period of about 0.5 h). During stage 12, Par6-GFP localization is further remodeled, with its pattern shifted from junctional region between adjacent border cells to the membrane facing the lumen-like cavity (Figure 3B, Video S2). Consistently, Lifeact-GFP and a PIP2 membrane reporter (PIP2-GFP) both demonstrate a similar remodeling in their distribution patterns from stage 9 to stage 12, with Lifeact-GFP and PIP2-GFP highly enriched in the same membrane region surrounding the luminal space in wild-type border cells at stage 12 (Figures 3C and 3D, Videos S3 and S4).

A very small percentage of wild-type stage 11 or 12 egg chambers would contain border cells that failed to migrate properly and reach the oocyte border (Figures S4A–S4E). Interestingly, we found that those stages 11 and 12 border cells with migration defects also displayed lumen formation that was accompanied by the remodeling of apical markers, F-actin, and PIP2-enriched membrane and was similar to the DHR3-induced lumen formation process at stages 9 and 10 (Figures S4A–S4E). This result indicates that the remodeling process is autonomously initiated in border cells and is under strict temporal control. Finally, DHR3 knockdown or E75 overexpression each led to disruption of the remodeling process (Figures 3E and 3F). As shown by the PIP2-GFP marker (Figure 3E), most of DHR3 RNAi or E75 overexpressing border cell clusters at stages 12 and 13 displayed a cluster morphology that is characteristic of border cells at stages 9 and 10 (Figure 3C, Video S3), where the PIP2-GFP is broadly localized in membranes between adjacent border cells. Consequently, these border cells failed to develop into the anterior tip of micropyle that surrounds a lumen-like cavity.

Together, these results demonstrate that DHR3 activity is required for the morphogenetic process of lumen formation that is essential to micropyle formation. Interestingly, the morphogenetic remodeling process involved in micropyle formation is similar to the DHR3-induced lumen formation process occurred...
precociously in border cell cluster during stages 9 and 10, suggesting that DHR3 is not only required but also sufficient for all the remodeling events necessary for lumen formation.

**Reduction of EcR Signaling and E75 Levels Causes De-repression of DHR3 Activity**

We next sought to understand how DHR3 function is temporally regulated to limit its lumen-forming activity only to the period of micropyle formation and not to the period of collective migration. We reasoned that DHR3's activity in border cells had to be inhibited by E75 during stages 9 and 10, as shown by our results above (Figure 2). Afterward, DHR3's activity would need to be de-repressed beginning at stage 11 to start the morphogenetic process of lumen formation. We already showed that DHR3 function is...
antagonized by E75 and that both E75 and DHR3 are expressed by EcR during border cell migration at stage 9. We then examined the temporal expression patterns of E75 and DHR3 as well as the levels of EcR signaling. We found that EcR signaling, as reflected by its well-established reporter EcRE-lacZ, reaches its highest levels during stage 10B, and then dramatically declines from stage 11 to stage 13 (Figures 4A and 4E). Accordingly, both expression levels of the E75-lacZ reporter, which reflects the transcription levels of E75 (Figures 4B and 4F) (because E75 antibody cannot penetrate and stain stages 11–13 egg chambers very well), and the protein levels of DHR3 as detected by DHR3 antibody also decrease from stage 11 to stage 13 (Figures 4C and 4G). These results suggest that as ecdysone signaling decrease dramatically (beginning at stage 11) E75 levels should also decrease to a low level (at stages 11 and 12), which could be below the threshold level for inhibition of DHR3’s activity (note: the real E75 protein level could be lower than the level detected by the antibody).

Figure 4. Temporal Expression Patterns of EcRE-lacZ, E75-lacZ, DHR3, and βFtz-f1-lacZ from Stage 10B to Stage 13

(A–D) Confocal images showing antibody staining of β-gal that is expressed by the EcRE-lacZ reporter (A), E75-lacZ enhancer trap (B), and βFtz-f1-lacZ enhancer trap (D), as well as antibody staining of DHR3 (C), from stage 10B to stage 13. Boxed regions are enlarged and shown at the right of all panels. Areas encircled by yellow dotted lines (based on labeling of GFP as expressed by slbo-Gal4) highlight the border cell clusters (A–D) at stages 12 and 13. Scale bars, 10 μm.

(E–H) Quantification of antibody staining of border cell clusters in (A–D) from stage 10B to stage 13. (E, F, G, and H) corresponds to (A, B, C, and D) respectively. The number of border cell clusters examined (n) for each stage is given at the x-axis. Statistical analysis was performed using two-tailed Student’s t test. Error bars indicate S.E.M.

See also Figure S5.
even lower than the E75-lacZ staining level due to β-galactosidase’s long perdurance). To test the above idea, we need a good activity reporter for DHR3’s function.

Previous literatures indicate that DHR3’s immediate downstream target gene during metamorphosis is βFtz-f1 (Liu et al., 2017; Kagayama et al., 1997; Lam et al., 1997), whose expression levels serve as a readout for DHR3 activity. We obtained an enhancer trap line for βFtz-f1, βFtz-f1-lacZ, which supposedly could reflect the transcription level of βFtz-f1. We found that its expression could serve as a bona fide reporter for DHR3 activity, based on the following results. First, βFtz-f1-lacZ expression is initially at non-detectable levels at stages 9 and 10A (Figure S5A), and at very low levels at stage 10B (Figures 4D and 4H), then it abruptly reaches much higher levels at stages 11, 12, and 13 (Figures 4D and 4H). Therefore, βFtz-f1-lacZ’s temporal expression pattern is highly consistent with our above prediction about the temporal regulation of DHR3 activity. Second, DHR3 overexpression led to precocious expression of βFtz-f1-lacZ within border cells during stages 9 and 10, whereas coexpression of E75B and DHR3 suppressed such precocious expression (Figures 5A and 5C). Conversely, DHR3 RNAi, E75B overexpression, or E75B and DHR3 coexpression, each suppressed βFtz-f1-lacZ’s normal expression in border cells during stage 11 (Figures 5B and 5D). Furthermore, DHR3 overexpression in the follicle cells at the stage 9, when βFtz-f1-lacZ is not normally expressed, ectopically turned on βFtz-f1-lacZ’s expression in the follicle cells (Figure S5B). Third, expression of βFtz-f1-RNAi in the background of E75 RNAi partially rescues border cell’s migration defects (Figures 5E and 5A), suggesting that βFtz-f1 functions downstream of DHR3. Lastly, expressing βFtz-f1-RNAi in the border cells resulted in the disruption in the formation of micropyle tip, similar to the loss-of-function defects of DHR3 RNAi (Figures S6B and S6C). On the other hand, overexpression of βFtz-f1 in stage 10 border cells resulted in actin-enriched patches that are similar to PAP from moderate E75 RNAi defects (Figures S3A, S3B, and S6A), suggesting an incomplete lumen forming phenotype. Taken together, these results support the conclusion that reduction in Ecd signaling and E75 levels leads to de-repression of DHR3 activity (as represented by the βFtz-f1-lacZ reporter) beginning at stage 11, which serves to switch on lumen formation for micropyle formation (during stages 11–13). Moreover, βFtz-f1 acts downstream of DHR3 to mediate micropyle formation.

DHR3 Is Required and Sufficient for Chitin Secretion into the Lumen

An essential feature of de novo lumen formation in the vertebrates is the secretion of glycoprotein such as the negatively charged podocalyxin into the lumen to keep the lumen membranes apart and promote the expansion of luminal space (Bryant et al., 2014; Strilic et al., 2010). Although Drosophila does not possess a podocalyxin homolog, the tube formation during Drosophila tracheal development requires the secretion of chitin into the lumen (Devin et al., 2005). Chitin is a long-chain polymer of N-acetylglucosamine, which is also a primary component of the Drosophila exoskeleton (Moussian et al., 2005; Zhu et al., 2016).

We then proceeded to determine whether chitin is present in the lumen enclosed by the border cells and whether DHR3 acts to promote secretion of chitin into the lumen. Interestingly, we found that chitin (labeled by FB-28) is only present in the extracellular space adjacent to wild-type border cells during and after stage 11 (Figures 6B, 6D, 6E, and S7E), whereas it is not present around the border cells before stage 11 (Figures 6A, 6C, and S7E). A very small percentage of wild-type stage 12 egg chambers would contain border cells that failed to migrate properly and reach the oocyte border, and we found that chitin is present within the lumen surrounded by those border cells (Figure 6F). Together, these results indicate that chitin is present in the lumen within the border cell cluster. In addition, the temporal and localization patterns of chitin suggest that it is secreted by border cells beginning at stage 11 during lumen formation for the micropyle tip. Furthermore, we found that expressing E75 RNAi or DHR3 specifically in the border cells (by slbo-Gal4) each resulted in chitin being precociously localized within the lumen of border cell clusters that failed to migrate to the oocyte border at stage 10 (Figures 6A and 6C), indicating that DHR3 activation is sufficient to induce chitin secretion. On the contrary, DHR3 knockdown or E75 overexpression in border cells led to loss of extracellular chitin adjacent to border cells (Figures 6B and 6D), indicating that DHR3 is required for chitin secretion by the border cells. Together, these results indicate that DHR3 activity is necessary and sufficient for chitin secretion by the border cells during lumen formation. To further test whether βFtz-f1 is also sufficient for chitin secretion, we examined and found no chitin secretion in βFtz-f1 overexpressing border cells at stage 10 (Figures S7A and S7C). This result could be due to the aforementioned fact that βFtz-f1 overexpression only resulted in PAP (incomplete lumen formation, Figure S6A). Hence, it is conceivable that chitin secretion may only occur after lumen formation progresses to a certain degree. On the other hand, we found that βFtz-f1 is required for chitin secretion by the border cells during
micropyle formation (Figures S7B and S7D), similar to DHR3’s role (Figures 6B and 6D). These results suggest that βFtz-f1 may act downstream of DHR3 to partially mediate DHR3’s chitin secretion role.

DHR3 and βFtz-f1 Suppress JNK Signaling in the Border Cells
Lastly, we sought to explore what signaling pathways DHR3 regulates in border cells. We tested reporters for a number of signaling pathways previously known to play essential roles in the border cells, including JAK/STAT (Beccari et al., 2002; Silver et al., 2005), Notch (Wang et al., 2007), JNK (c-Jun N-terminal kinase) (Llense and Martin-Blanco, 2008; Melani et al., 2008), and Dpp (Luo et al., 2015). Among them, JNK was the only signaling found to be severely affected by E75 knockdown or DHR3 overexpression (Figure S8A). Staining for Puc-lacZ, a widely used reporter for JNK signaling, revealed that both E75 RNAi and DHR3

Figure 5. βFtz-f1 Acts Downstream of DHR3 and βFtz-f1-lacZ Serves as a Reporter of DHR3 Activity
(A) βFtz-f1-lacZ levels in border cells at stage 10 as represented by β-gal antibody staining. Compared with wild-type (WT) control, E75 RNAi and DHR3 overexpression both result in significant increase of βFtz-f1-lacZ levels (quantified in C), whereas double knock down of E75 and DHR3 (E75 Ri + DHR3 Ri) and overexpression of both DHR3 and E75 (UAS-DHR3 + UAS-E75) abolish the increase (quantified in C).
(B) βFtz-f1-lacZ levels in border cells at stage 11 as represented by β-gal staining. Compared with wild-type (WT) control, DHR3 RNAi and E75 overexpression both result in significant reduction of βFtz-f1-lacZ levels (quantified in D). Yellowed dotted lines (A and B) outline individual border cell clusters, as labeled with GFP expressed by slbo-Gal4. Boxed regions (A, B) are enlarged and shown at the right of all panels. Scale bars, 50 μm for egg chambers; 10 μm for border cells.
(C and D) Quantification of βFtz-f1-lacZ levels. The number of border cell clusters examined for each genotype is indicated within its corresponding column. Statistical analysis was performed using two-tailed Student’s t test. Error bars indicate S.E.M. **, p < 0.01; ***, p < 0.001; ****, p < 0.0001; ns, not significant.
(E) Quantification of rescue of border cell migration defects of E75 RNAi by co-expression of βFtz-f1 RNAi. Represented images for the indicated genotypes are shown in Figure S6A.

DHR3 and βFtz-f1 Suppress JNK Signaling in the Border Cells
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overexpression caused strong reduction of Puc-lacZ reporter activity in stage 10 (Figures 7A–7C), indicating that increased DHR3 activity suppresses JNK signaling. Indeed, knockdown of DHR3 in the background of E75 RNAi rescued the level of Puc-lacZ expression back to the wild-type level in stage 10 (Figures 7A and 7B). Furthermore, overexpressing bsk (encoding Drosophila JNK) in the background of E75 RNAi or DHR3 overexpression partially rescued the severe migration defects and precocious lumen formation of border cells that were resulted from E75 loss of function (Figures 7D and S8B). Together, these results demonstrate that reduction of E75 or increase of DHR3 activity leads to downregulation of JNK signaling in the migratory border cells at stages 9 and 10.

JNK signaling pathway was previously reported to be required for cell-cell adhesion between adjacent border cells during their collective migration (Llense and Martin-Blanco, 2008; Melani et al., 2008). Down-regulation of JNK signaling in border cells resulted in strong disruption of the coherent cluster morphology, with the most severe cases showing individual border cells dissociated from the cluster. We found that reduction of E75 or increase of DHR3 activity leads to similar but milder defects in cluster morphology. Twenty percent of E75 RNAi expressing border cell clusters (n = 64) at 29°C displayed an incoherent and outstretched cluster morphology (Figure S8C, second row). Furthermore, 27% of DHR3 expressing border cell clusters at 25°C (n = 46) exhibited a more severe incoherent and outstretched cluster morphology.
It should be noted here that our experiments involving Gal4 were normal, i.e., to maximize the efficiency of UAS/Gal4 system and phenotypic severity. Almost all of the DHR3 expressing clusters at 29°C/C14C displayed severe migration block phenotype (no migration, Figure S3B), whereas a majority of DHR3 expressing clusters at 25°C/C14C could detach and migrate.

Figure 7. DHR3 and bFTZ-f1 Downregulate JNK Signaling in the Border Cells
(A) Puc-lacZ expression levels in migratory border cells at stage 9 or 10 as indicated by representative confocal images of β-gal antibody staining. (B) Quantification indicates that E75 RNAi, DHR3 and bFTZ-f1 overexpression as shown in (A) each results in strong and significant decrease of Puc-lacZ levels as compared to wild type (WT) control, while coexpression of DHR3 RNAi and E75 RNAi returns the Puc-lacZ levels to that of WT.

Figure S8C, third row). It should be noted here that our experiments involving Gal4 were normally carried out at 29°C to maximize the efficiency of UAS/Gal4 system and phenotypic severity. Almost all of the DHR3 expressing clusters at 29°C displayed severe migration block phenotype (no migration, Figure S3B), whereas a majority of DHR3 expressing clusters at 25°C could detach and migrate.
Importantly, we only observed incoherent cluster morphology in those DHR3 clusters (25°C) and E75 RNAi clusters (29°C) that have detached and undergone migration, suggesting that strong cell-cell adhesion is especially needed for migratory border cells to maintain their coherent cluster morphology. Lastly, E75 RNAi expression in a small Flip-out clone of two border cells resulted in these two border cells losing cohesion and almost dissociating themselves from the rest of cluster (Figure S3C). Taken together, these results demonstrate that E75 RNAi or DHR3 expression could result in significant cell-cell adhesion defects, indicating that increased DHR3 activity could downregulate cell adhesion.

We next tested whether JNK signaling was negatively regulated by DHR3 and βFtz-f1 during micropyle formation. We showed that in the wild type the level of JNK signaling was reduced from stage 10 to stage 11 and then further reduced from stage 11 to stage 12 (Figures 7E and 7F). Knockdown of βFtz-f1 in the border cells at stage 12 increased the originally low JNK signaling to a much higher level, which is comparable with the level at stage 10 (Figures 7E and 7F). These results suggest that JNK signaling needs to be suppressed in order for lumen formation to proceed properly during stages 11 and 12. Indeed, overexpression of bsk and hence increase of JNK signaling resulted in disruption of lumen formation during formation of the micropyle tip at stage 12 (Figures 7G and 7H). Taken together, these results suggest that JNK-mediated cell adhesion between border cells is temporally and differentially regulated during two different morphogenetic processes—collective migration and micropyle formation—and that its downregulation by DHR3 is essential for lumen formation in the latter process.

DISCUSSION

We demonstrate that two nuclear receptors, E75 and DHR3, are critical for temporal regulation and coordination between two very different morphogenetic processes of the border cell cluster, namely its collective migration and its lumen formation. First, our results revealed that the levels of E75 and DHR3 (in response to ecdysone) are the underlying control of the temporal order (Figure 8). Strong loss of function of E75 or DHR3 overexpression disrupts the temporal order and causes lumen formation to occur first. Consequently, collective migration could not take place afterward, because of the unique nature of the lumen structure, which precludes migration from occurring. Second, levels of E75 and DHR3 together with the antagonism between the two nuclear receptors underlie the mechanistic control of the time interval between the two morphogenetic processes (Figure 8). E75 acts as a molecular timer. Its expression level determines the length of interval between migration and lumen formation (Figure 8). Very little E75 (strong loss-of-function) causes lumen formation to occur before migration could take place, effectively resulting in no interval between the two morphogenetic processes. Moderate E75 loss-of-function phenotype demonstrates that collective migration could take place at early stage 9 (Figures S3A and S3B) but accompanied with a precocious occurrence of lumen formation at late stage 9 or stage 10, indicating a shortened interval. On the other hand, too much E75 (E75 overexpression) results in reduced occurrence of lumen formation at stage 12 or 13 (Figure 3F), suggesting an expanded interval. Finally, it is important to note that border cells that supposedly contain the wild-type levels of E75 and DHR3 but display migration defects exhibit a normal time interval (Figures S4A–S4F, and 6F). During tissue or organ formation, it is not uncommon for a certain cell population to undergo two vastly different morphogenetic processes. This study provides a novel mechanistic insight into the molecular machinery that coordinates both the order and time interval between morphological processes.

It is well known in the metamorphosis literatures that E75 and DHR3 genes are sequentially activated by ecdysone (Huet et al., 1995; White et al., 1997). As a result, E75 is considered as an “early response gene,” whereas DHR3 is considered as an “early-late response gene” (Huet et al., 1995; White et al., 1997). A previous study reported that E75b and DHR3 reached their respective peak protein levels at different time points (about 2 h apart) immediately after pupariation (White et al., 1997). Because E75b level began to decrease a couple of hours before DHR3 did, DHR3 was eventually de-repressed, resulting in the gene activation of βFtz-f1 and appearance of βFtz-f1 protein about 2 h after the E75b decrease began, a time point where E75 was at a low level and DHR3 at a medium level (White et al., 1997). It had been suggested that the ecdysone response genes that control pupal transition were also used reiteratively during oogenesis (Ables et al., 2016). Therefore, it is conceivable that E75 and DHR3 genes could also be sequentially activated by ecdysone in the border cells during early stage 9, and their protein levels reach their respective peaks at different time points during stage 10 (1–2 h apart, Figure 8). It should be noted that such a small time difference (1–2 h) would not be readily detectable with our reporters and antibodies. Consequently, as the E75 level drops below a threshold level immediately before stage 11, the DHR3
protein level, although declining, is still at an intermediate level (Figure 8). As a result, a sufficient amount of DHR3 is de-repressed (activated) to activate \(\beta\)Ftz-f1 expression and start the lumen formation process.

Our study also uncovers a surprising mechanism of how a nuclear receptor controls the process of de novo lumen formation. DHR3 seems to act as a master switch or potent inducer for lumen formation because it is necessary and sufficient for lumen formation of border cells both during stage 9 and during stages 11–13. Activation of DHR3 function in border cells seems to simultaneously induce multiple cellular processes that were previously demonstrated to be essential for de novo lumen formation in other systems (Sigurbjörnsdottir et al., 2014). In addition, DHR3 is necessary and sufficient for the secretion of chitin into the lumen of border cells both at stage 9 and at stage 12. Chitin had been previously shown to be required for tube expansion and maturation during Drosophila tracheal morphogenesis (Devine et al., 2005). Its function seems to provide an extracellular matrix support (Moussian et al., 2006; Wang et al., 2006). The mechanism by which chitin affects tube morphogenesis remains poorly understood. How DHR3 induces chitin synthesis and secretion and whether chitin is required for lumen formation and tube maturation in micropyle remain to be further determined.

Furthermore, we demonstrate that DHR3’s lumen-inducing function is mainly mediated through \(\beta\)Ftz-f1, a nuclear receptor and transcription factor that has been well established to be DHR3’s immediate target gene during metamorphosis. However, \(\beta\)Ftz-f1 does not seem to mediate all of DHR3’s functions because overexpression of \(\beta\)Ftz-f1 could not induce complete lumen structure and chitin secretion, suggesting that other factors downstream of DHR3 may contribute to lumen formation. Lastly, we show that JNK signaling is downregulated by DHR3 and \(\beta\)Ftz-f1, suggesting that cell adhesion between adjacent border cells needs to be reduced during lumen formation. This is consistent with the idea that remodeling of apical polarity,
cytoskeleton, and membrane during lumen formation may require downregulation of cell-cell adhesion. Given the multiple functions as demonstrated for DHR3, it will be interesting to test whether these lumen inducing functions will be conserved in other developmental contexts in Drosophila and vertebrate. It is reported that E75 and DHR3 are expressed in tissues that form luminal structures including gut, trachea, and salivary gland (Wilk et al., 2013). Moreover, the ligand sensor of DHR3 displays its activity in gut, trachea, and Malpighian tubules (Palanker et al., 2006). Further study is needed to determine whether E75 and DHR3 are involved in lumen formation in those tissues. Interestingly, previous studies reported that the mammalian homolog of DHR3, RORα, was enriched in human mammary duct, and its inactivation impaired polarized acinar morphogenesis (Xiong et al., 2012; Xiong and Xu, 2014), raising the possibility of a similar lumen-forming role in vertebrate.

Although treated as an excellent model system for collective migration, border cells’ physiological function during oogenesis is to make a functional opening within the micropyle for sperm entry. How the border cell cluster develops into the anterior tip of the tubular structure of micropyle is poorly understood. Our study reveals a dynamic remodeling of apical polarity molecules, F-actin, and PIP2-enriched membrane, which is consistent with the process of de novo lumen formation. The functional roles of DHR3, βFtz-f1, EcR, E75, and JNK during micropyle formation, as demonstrated by our study, provide the first detailed analysis of this morphogenetic process. We suggest that in addition to collective migration, border cells could also serve as a model system to study de novo lumen formation in Drosophila.

Limitations of the Study
In this study, we demonstrate that DHR3’s lumen-inducing function is mainly mediated through βFtz-f1, a nuclear receptor and transcription factor that has been well established to be DHR3’s immediate target gene during metamorphosis. However, what target genes are directly activated or repressed by βFtz-f1 are still unknown. Further studies will be needed to reveal those target genes, for their discovery will be critical to understanding the molecular mechanisms of lumen formation.

Resource Availability
Lead Contact
Further information and requests for reagents should be directed to and will be fulfilled by the Lead Contact, Jiong Chen (chenjiong@nju.edu.cn).

Materials Availability
This study did not generate new unique reagents.

Data and Code Availability
The published article includes all datasets generated or analyzed during this study.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101335.

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AUTHOR CONTRIBUTIONS
Conceptualization, J.C., X.P.W., S.L., and G.E.; Methodology, X.P.W., J.C., and H.W.; Investigation, X.P.W., L.L.; Resources, S.L., G.E.; Visualization, X.P.W. and H.W.; Writing, J.C., X.P.W., H.W., and G.E.; Supervision, J.C. and H.W.

DECLARATION OF INTERESTS
The authors declare no competing financial interests.

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Supplemental Information

Temporal Coordination of Collective Migration and Lumen Formation by Antagonism between Two Nuclear Receptors

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Supplemental Information

Supplemental figures

Figure S1 A small-scale RNAi screen, rescue of DHR3 RNAi and E75 antibody staining (Related to Figure 1).

(A) Migratory border cells in wild type, EcR RNAi and E75 RNAi egg chambers respectively. Ri is the abbreviation of RNAi. White boxes mark their typical positions at stage 10. RNAi expression was driven by the slbo-Gal4 line. (B) Diagram of an egg chamber at early stage 10. The color-coded bars indicate positions along the migratory route of border cells. 100% indicate the position that wild type border cells would have normally reached at stage 10, while the 75%, 50%, 25% and 0% positions
represent various degrees of migration defects (with increasing severity). (C) Quantification of border cell migration. The x-axis denotes the percentage of stage 10 egg chambers examined for each line that exhibited each degree of migration, as represented by the five color-coded bars in (B). For example, 92.9% of wild type control (WT; line 6458) egg chambers at stage 10 exhibited 100% migration, 5.3% of WT egg chambers were in the 75% migration delay category, and the remaining 1.8% of WT egg chambers were in the 50% migration delay category. The number of stage 10 egg chambers examined for each line is given (n). The stock number is listed within the parenthesis next to the RNAi of the indicated gene (see Methods for details). (D) Border cells expressing DHR3 RNAi displayed cluster morphology and F-actin distribution pattern that were similar to the wild type. DHR3 RNAi expression could rescue defects of cluster morphology and F-actin distribution in DHR3 overexpressing border cells. (E, F) E75B antibody staining for migratory border cells. (E) E75B staining of border cells that have already reached the oocyte border. (F) E75 staining of border cells during migration. A small Flip-out clone of border cells expressing E75-RNAi within a mosaic cluster shows absence of E75B staining, demonstrating the specificity of the E75B antibody. Scale bars: 50 µm for egg chambers, 10 µm for border cells.
Figure S2 E75 exerts a positive feedback on EcR signaling (Related to Figure 1). 

(A-F) EcRE-lacZ levels in border cells at stage 10 as represented by β-gal staining. Compared to wild type control (A), E75 RNAi (B) and DHR3 overexpression (C) both result in significant and strong reduction of EcRE-lacZ levels (quantified in F), while double knock down of E75 and DHR3 (D) largely abolish the reduction (quantified in F). DHR3 RNAi alone does not cause any significant change in EcRE-lacZ levels (E, F). Statistical analysis was performed using two-tailed Student’s t-test. Error bars indicate S.E.M. **, P<0.01; ***, P<0.001; ****, P<0.0001; ns, not significant. Scale bars: 50 µm for egg chambers, 10 µm for border cells.
Figure S3 A range of lumen forming phenotypes by *E75 RNAi* and *DHR3* overexpression (Related to Figures 2 and 3).

(A) *DHR3* overexpressing border cells exhibits more severe migration and lumen phenotypes than those of *E75 RNAi* border cells, which displays a range of phenotypes with various severity. Compared to the WT migratory border cells during mid stage 9, *DHR3* expressing border cells at stage 10 exhibits the most severe phenotype of strong lumen formation coupled with migration being blocked at 0% migratory distance (lumen at 0%, last row). *E75 RNAi* border cells display a range of similar but less severe phenotypes, including formation of pre-apical patches with migration blocked at 0% distance (PAP at 0%, second row), PAP formation with migration stopped at 25% to 75% of migratory distance (PAP at 25%-75%, third row), and complete lumen structure with migration stopped at 25% to 75% distance (lumen at 25%-75%, fourth row). aPKC labels the apical membrane. DAPI labels nuclei. GFP as expressed from *slbo-Gal4* labels the border cells, cells with lack of GFP but DAPI labeling are the two polar cells and region with no GFP and DAPI labeling is the lumen space. but not the polar cells. (B) Quantification of the range of the phenotypes described in (A). Number of egg chambers examined for each genotype is denoted within each column. (C) *E75 RNAi* expression in a small Flip-out clone of two border
cells (clone marked by GFP) that were driven by Ay-Gal4 resulted in a small lumen-like structure (label by aPKC) at the interface between the two border cells. (D) aPKC staining of E75 RNAi border cells expressed by c306-Gal4. A central lumen outlined by aPKC is shown. Scale bars: 50 μm for egg chambers, 10 μm for border cells.

**Figure S4 Lumen formation in delayed WT border cells (Related to Figure 3).**

(A-D) Delayed wild type (WT) border cells that failed to reach oocyte at stages 11-12 display the lumen structure that is outlined by a ring of aPKC (A), Par6-GFP (B), Lifeact-GFP (C), and PIP2-GFP (D). The stage 10 border cells (first rows, A-D) are used as references showing their cluster morphology, which are distinct from the lumen morphology shown by the stages 11 and 12 border cells in the second rows.
(A-D). (E) A stage 13 egg chamber showing its border cell cluster with a lumen structure outlined by phalloidin stained F-actin. (F) A stage 11-12 egg chamber showing its delayed WT border cell cluster with a lumen outlined by F-actin (as labeled by Lifeact-RFP). A polar cell (marked by *) is shown to be displaced at the side. (G, H) WT stage 11 and stage 12 border cell undergo normal micropyle formation, with position of polar cells clearly marked (*). Lifeact-GFP labels F-actin, which is enriched in the region where lumen formation initiates in stage 11 border cells (G). F-actin also outlines the large lumen cavity at stage 13 (H). Judging from their positions, polar cells appear not to be closely involved in lumen formation. Scale bars: 50 µm for egg chambers, 10 µm for border cells.

Figure S5 βFtz-f1-lacZ expression in stage 9 border cells and follicle cells (Related to Figures 4 and 5).

(A) Lack of βFtz-f1-lacZ expression in border cells at early stage 9, mid stage 9 and stage 10a, as shown by β-gal antibody staining. (B) Ectopic increase of βFtz-f1-lacZ levels induced by DHR3 expression in flip-out clone of follicle cells (labeled by GFP) of an early stage 9 egg chamber. Scale bars: 50 µm for egg chambers, 10 µm for border cells.
**Figure S6** βFtz-f1 RNAi's rescue effect on E75 RNAi and βFtz-f1 RNAi's micropyle defects (Related to Figure 5).

(A) Representative images for the migratory border cells of indicated genotypes quantified in Figure 5E. (B) βFtz-f1 RNAi caused disruption of lumen formation during micropyle formation, as compared to morphology of wild type border cells (outlined by PIP2-GFP) at stage 12. Their cluster or lumen morphology is quantified in (C). (C) 79.9% of stage 12 wild type border cells display obvious lumen morphology, whereas 81.8% of βFtz-f1 RNAi border cells display cluster morphology, which is characteristic of the wild type border cells at stage 10. Scale bars: 50 µm for egg chambers, 10 µm for border cells.
Figure S7 Chitin staining in WT and βFtz-f1 RNAi border cells (Related to Figure 6).

(A, B) Confocal and DIC images showing chitin staining in stage 10 (A) and stage 11 (B) egg chambers. Chitin is labeled with the Fluorescent Brightener 28 (FB28) dye. (A) Chitin is not detected within or close to wild type (WT) and βFtz-f1 overexpressing border cells (quantified in C). (B) Starting at stage 11, chitin is detectable adjacent to WT border cell cluster, but βFtz-f1 RNAi abolishes chitin staining (quantified in D). Yellow dotted lines outline individual border cell clusters as labeled with GFP expressed by slbo-Gal4. (C, D) Quantification of chitin levels of border cells for the indicated genotypes. Statistical analysis was performed using two-tailed Student’s t-test. Error bars indicate S.E.M. ****, P<0.0001; ns, not significant. (E) Confocal and DIC images showing chitin staining in stages 10-13 egg chambers. Lifeact-GFP labels F-actin. Scale bars: 50 μm for egg chambers, 10 μm for border cells.
Figure S8 JNK signaling identified from screen and its partial rescue of *E75 RNAi* and *DHR3* overexpression (Related to Figures 7 and 2).

(A) Activity reporters for JAK-STAT, Dpp, JNK and Notch signaling pathways were tested for in the wild type (WT) and *E75 RNAi* border cells, they include Stat-GFP, Dad-lacZ, Puc-lacZ, and M7-lacZ respectively. (B) Confocal images showing *bsk* overexpression partially rescued the severe migration defects and ectopic lumen formation of border cells as resulted from *E75 RNAi* or *DHR3* overexpression (migration rescue quantified in Figure 7D). (C) 20% of *E75 RNAi* expressing border cell clusters (n=64) at 29°C displayed an incoherent and outstretched cluster morphology. 27% of *DHR3* expressing border cell clusters at 25°C (n=46) exhibited a
more severe incoherent and outstretched cluster phenotype, with individual cells almost dissociated from the main cluster. E-cadherin (E-cad) labels the cell-cell junction between border cells. (D) Significant remodeling and relocalization of E-cadherin was observed in E75 RNAi expressing border cells. A range of E-cadherin relocalization phenotypes were observed, some with E-cadherin strongly enriched in a ring adjacent to that of aPKC staining (3rd row), others displaying some relocalization but not a strong enrichment of E-cadherin near the lumen (2nd row). Scale bars: 50 µm for egg chambers, 10 µm for border cells.

**Transparent methods**

*Drosophila genetics*

Flies were cultured and maintained on standard cornmeal media with sugar and yeast at 25°C. Progenies of crosses between UAS-RNAi or UAS-transgenes and slbo-Gal4 were cultured at 29°C for two days for specific gene’s knockdown and overexpression. For moderate knockdown or overexpression, flies were cultured at 29°C for only one day or cultured at 25°C, as indicated in the figure legends. To generate flip-out clones of border cells and follicle cells, female flies were heated-shocked for 3 minutes at 37°C and then kept at 29°C for 1 day before dissection.

Fly stocks listed below were obtained from different labs and stock centers, including Bloomington Stock Center (BDSC), National Institute of Genetics Stock Center, Japan (NIG), Vienna Drosophila RNAi Center (VDRC) and Tsinghua Fly Center (THFC).

| STOCK NAME   | SOURCE   | IDENTIFIER              |
|--------------|----------|-------------------------|
| UAS-E74 RNAi | VDRC     | v45900                  |
| UAS-E74 RNAi | NIG      | 6273R-3                 |
| UAS-E74 RNAi | NIG      | 6285-1R-1               |
| UAS-E75 RNAi | VDRC     | v44851 (used for Figure 1B) |
| UAS-E75 RNAi | BDSC     | 26717 (used for         |
| Gene          | Source        | Version/Stock Center | Notes                                      |
|--------------|---------------|----------------------|--------------------------------------------|
| UAS-E75 RNAi | THFC          | THU1738 (used for most of Figures) |
| UAS-E93 RNAi | VDRC          | v45855               |
| UAS-E93 RNAi | VDRC          | v45856               |
| UAS-Br-C RNAi| BDSC          | 38526                |
| UAS-Br-C RNAi| NIG           | 11514R-3             |
| UAS-EcR RNAi | VDRC          | v35078               |
| UAS-EcR RNAi | BDSC          | 9327                 |
| UAS-DHR3 RNAi| VDRC          | v12204               |
| UAS-DHR3 RNAi| VDRC          | v106837              |
| UAS-crol RNAi| VDRC          | v104313              |
| UAS-Hr39 RNAi| VDRC          | v37694               |
| UAS-Hr39 RNAi| VDRC          | v37695               |
| UAS-Su RNAi  | VDRC          | v105675              |
| UAS-Hr4 RNAi | VDRC          | v101856              |
| UAS-cact RNAi| BDSC          | 34775                |
| UAS-E2F RNAi | VDRC          | v15886               |
| UAS-Sox14 RNAi| VDRC       | v107146              |
| UAS-Sox14 RNAi| VDRC       | v10856               |
| UAS-Sox14 RNAi| BDSC       | 26221                |
| UAS-brat RNAi| VDRC          | v105054              |
| UAS-Kr-h1 RNAi| VDRC       | v51282               |
| UAS-Kr-h1 RNAi| VDRC       | v31333               |
| UAS-Ef4A RNAi| VDRC          | v45686               |
| UAS-Ef4A RNAi| VDRC          | v107846              |
| UAS-Kis RNAi  | VDRC          | v109414              |
| UAS-E63F-1 RNAi| VDRC       | v26899               |
| UAS-Cyp4e2 RNAi| VDRC      | v108025              |
| UAS-bip1 RNAi| VDRC          | v26104               |
| UAS-Impl2 RNAi| VDRC       | v106543              |
| UAS-Past1 RNAi| VDRC        | v22253               |
| UAS-srp RNAi  | VDRC          | v109521              |
| UAS-Eip78C RNAi| BDSC      | 28851                |
| UAS-Eip78C RNAi| BDSC      | 26718                |
| UAS-vrille RNAi| VDRC       | v5650                |
| UAS-βFtz-f1 RNAi| BDSC  | 27659                |
| UAS-βFtz-f1 RNAi| VDRC    | v104463              |
| Slbo-gal4,UAS-GFP/Cyo | BDSC | 6458                |
| C306-gal4 | Kyoto Stock Center | 107215              |
| Slbo-PH(PLCδ)-4xGFP,Ub-His-tone-RFP,Slbo-gal4,Upd-Gal4 | Gift from Pernille Roth (Cliffe et al., 2017) | N/A (used for Figures 3C, 3E, 7G and S6B) |
| **UMAT-Lyn-tdTomato** | **BDSC** | **4411** |
|------------------------|----------|---------|
| **Ay-Gal4,UAS-GFP**    | **BDSC** | **N/A** |
| **Slbo-lacZ,Slbo-Gal4/Cyo** | **Gift from Pernille Roth** | **N/A** |
| **UAS-E75A**           | **Gift from Henry M. Krause (Caceres et al., 2011)** | **N/A** |
| **UAS-E75B.Flag**      | **Gift from Oren Schuldiner (Rabinovich et al., 2016)** | **N/A** |
| **UAS-E75C.Flag**      | **Gift from Oren Schuldiner (Rabinovich et al., 2016)** | **N/A** |
| **UAS-DHR3**           | **Gift from Henry M. Krause (Caceres et al., 2011)** | **N/A** |
| **UAS-Bsk.B**          | **BDSC** | **9310** |
| **UAS-βFtz-f1**        | **BDSC** | **64290** |
| **UAS-LifeAct.GFP**    | **BDSC** | **35544** |
| **UAS-LifeAct.RFP**    | **BDSC** | **N/A** |
| **UAS-PLCδ-PH-EGFP**   | **BDSC** | **39693 (used for Figure 2E)** |
| **UAS-Par6.GFP**       | **BDSC** | **65847** |
| **UAS-Myr.RFP**        | **BDSC** | **7119** |
| **UAS-GFP**            | **BDSC** | **4776** |
| **UAS-Puc**            | **Gift from Lei Xue (Ma et al., 2011)** | **N/A** |
| **EcRE-lacZ**          | **BDSC** | **4517** |
| **E75-lacZ**           | **BDSC** | **11712** |
| **βFtz-f1-lacZ**       | **BDSC** | **11598** |
| **PucE69-lacZ**        | **Gift from Xue Lei (Ma et al., 2013)** | **N/A** |
| **E(sp)m7-lacZ**       | **From Zizhang Zhou and Qing Zhang (Tseng et al., 2014)** | **N/A** |
| **Dad-lacZ**           | **From Zizhang Zhou and Qing Zhang (Ninov, 2010)** | **N/A** |
| **10xStat-GFP**        | **BDSC** | **26197** |
| **Crb-HA**             | **Gift from Juan Huang (Huang et al., 2009)** | **N/A** |

**Immunostaining**

Female flies were raised on fresh food with yeast at 29°C for 2 days. Ovaries were dissected in PBS, and then fixed in 100µl devitellinizing buffer (7% formaldehyde) and 600µl heptane, with strong shaking for 10 min, then washed 3 x10min with PBS,
and 3x10 min with PBST. For egg chamber staining, egg chambers were blocked with 10% goat serum in PBST for 30 min after fixed and washed, and then incubated with primary antibody at 4°C overnight. Ovary samples were then washed 3x20 min with PBST, blocked with 10% goat serum in PBST for 30 min, incubated with secondary antibody at 1:200 in PBST for 2 hours. DAPI (Santa Cruz, sc-3598) was added and stained for 30 min during secondary staining. Lastly, ovaries were washed again with 20 min PBST for three times, mounted on microscope slide with 40% glycerol. Primary antibodies used include mouse anti-lacZ (1:100, DSHB,401-a), mouse anti-E75B (1:20, gift from Henry M. Krause)(Caceres et al., 2011), rabbit anti-DHR3 (1:100, gift from Carl S. Thummel ) (Ruaud et al., 2010), Guinea Pig anti-Sec15 (1:200, gift from Hugo J.Bellen)(Mehta et al., 2005), Mouse anti-Rab11 (1:200, BD, 610656), Rabbit anti-p-JNK (1:50, Promega, V7932), Rat anti-E-cad (1:50, DSHB, 5D3), Rabbit anti-PKCζ (1:100, Santa Cruz, C-20), Mouse anti β-tubulin (1:100, DSHB,E7), mouse anti-Dlg (1:100, DSHB, 4F3) , mouse anti-HA(1:100, Santa Cruz, F-7), rabbit anti-Baz (1:400, gift from A. Wodarz). Secondary antibodies were used including Cy5-AffiniPure Goat Anti-Rabbit IgG (1:200, Jackson ImmunoResearch), Cy3-AffiniPure Goat Anti-Rat IgG (1:200, Jackson ImmunoResearch), Cy3-AffiniPure Goat Anti-Mouse IgG (1:200, Jackson ImmunoResearch), Cy5-AffiniPure Donkey Anti-Guinea Pig IgG (1:200, Jackson ImmunoResearch). F-Actin was labeled by TRITC-conjugated Phalloidin (1:200, Sigma-Aldrich, P1951) without blocking. Confocal images were obtained with Leica SP5 confocal microscope and analyzed by software from Leica, Image J and Imaris.

Quantification of fluorescence and statistical analysis

A region of interest was first chosen and outlined. Then fluorescence intensity of the outlined region was measured and determined by Image J. To quantify nuclear staining for E75-lacZ, DHR3, βFtz-f1-lacZ, and Puc-lacZ (Figures 4F-4H, 5C, 5D, 7B and 7F), a representative border cell nucleus was selected for each border cell cluster. The selected nucleus is usually situated well in the focal plane and could be clearly viewed within the confocal section (usually displaying a full cross-sectional view, e.g.
full circle), other nuclei within the same cluster may not show a full view (partial cross-sectional view, e.g. crescent, half circle, or other irregular shapes) in the confocal image, due to their various z-axis positions. DAPI staining was sometimes used to help outline the nuclei. After the fluorescence intensity of the outlined nuclear region was determined, fluorescence intensity of a nearby cytoplasmic region (including nurse cell cytoplasmic region) of similar size was determined. Intensity of individual nucleus was then normalized to the intensity of nearby cytoplasmic region to obtain the relative intensity.

To quantify non-nuclear staining for EcRE-lacZ (Figures 4E), a region including all the border cells in view was outlined and its fluorescent intensity determined by Image J. For example, in Figure 4A (third and fourth rows), the outlined regions are areas encircled by yellow dotted lines, which are based on labeling of GFP as expressed by \textit{slbo-Gal4}. Intensity of the outlined region was then normalized to the intensity of nearby nurse cell cytoplasmic region of comparable size to obtain the relative intensity. To quantify FB-28 staining (Figures 6C and 6D), a region including the FB-28 staining was outlined and its fluorescent intensity determined by Image J. \textit{slbo-Gal4} expressed GFP and DIC views also help outline the region. Fluorescence intensity of a nearby nurse cell cytoplasmic region of similar size was also determined as the background intensity. Intensity of the outlined region was then normalized to the intensity of nearby nurse cell cytoplasmic region to obtain the relative intensity. Statistical analysis was performed with GraphPad Prism 6 using unpaired two-tailed Student's t-test, significance of p<0.05 was used as the criterion for statistical significance and indicated with *, p<0.01 was indicated with **, p<0.001 with three stars (***), and p<0.0001 with ****, not significant was indicated with “ns”.

**3-D imaging of border cell cluster**

We used 2 coverslips (0.13-0.17mm thick) as bridges to mount egg chambers so that there is ample space in the z-axis to avoid compression of border cell clusters. Individual confocal section was captured every 0.4 µm for each z-series of border cell
cluster. The z-series was then processed by Imaris software to view 3-D distributions of aPKC, Lifeact-GFP, Par6-GFP and PIP2-GFP (Figure 3, Videos 1-4).

Stage identification of egg chambers

Stages 9-13 egg chambers were staged according to previous literatures (Jia et al., 2016; Spradling, 1993). At stage 9, oocyte occupies about 1/3 of egg chamber volume, and border cells undergo anterior migration toward the border. For stage 10A egg chambers, oocyte occupies half of egg chamber volume, border cells have completed migration and reached the anterior border of oocyte. Moreover, anterior follicle cells have also completed their migration and arrived at the border between oocyte and nurse cells. At stage 10B, an anterior-most group of anterior follicle cells, named centripetal cells, migrate inwardly toward the anterior of oocyte, completely enclosing the oocyte. While oocyte still takes about 1/2 of egg chamber volume, the entire egg chamber is larger in volume than the stage 10A egg chamber. At stage 11, nurse cells begin the dumping process, allowing dramatic transfer of cytoplasm from nurse cells to oocyte. As a result, oocyte now occupies significantly more than 1/2 of egg chamber volume. At stage 12, nurse cells complete their dumping process, resulting in oocyte taking the vast majority (80-90%) of egg chamber volume and little space between adjacent nurse cell nuclei. At stage 13, some nurse cell nuclei become degraded and some still remain, dorsal filaments are visible.

Chitin staining

Ovaries were fixed as described for immunostaining but without blocking. Fluorescent Brightener 28 (FB28) (Sigma, F3543) was used as a chitin dye as previously reported (Zhao et al., 2018). We used FB28 (50mg/ml) with dilution of 1:400, stained ovaries in PBST for 30 min, washing 3x20 min with PBST.

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