A switch in the mode of Wnt signaling orchestrates the formation of germline stem cell differentiation niche in *Drosophila*

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Abstract

Germline stem cell (GSC) self-renewal and differentiation into gametes is regulated by both intrinsic factors in the germ line as well as extrinsic factors from the surrounding somatic niche. dWnt4, in the escort cells of the adult somatic niche promotes GSC differentiation using the canonical β-catenin-dependent transcriptional pathway to regulate escort cell survival, adhesion to the germ line and downregulation of self-renewal signaling. Here, we show that in addition to the β-catenin-dependent canonical pathway, dWnt4 also uses downstream components of the Wnt non-canonical pathway to promote escort cell function earlier in development. We find that the downstream non-canonical components, RhoA, Rac1 and cdc42, are expressed at high levels and are active in escort cell precursors of the female larval gonad compared to the adult somatic niche. Consistent with this expression pattern, we find that the non-canonical pathway components function in the larval stages but not in adults to regulate GSC differentiation. In the larval gonad, dWnt4, RhoA, Rac1 and cdc42 are required to promote intermingling of escort cell precursors, a function that then promotes proper escort cell function in the adults. We find that dWnt4 acts by modulating the activity of RhoA, Rac1 and cdc42, but not their protein levels. Together, our results indicate that at different points of development, dWnt4 switches from using the non-canonical pathway components to using a β-catenin-dependent canonical pathway in the escort cells to facilitate the proper differentiation of GSCs.

Author summary

Germ line association with the somatic cells is critical for various aspects of germ cell biology, including migration, self-renewal and differentiation. In *Drosophila* females, soma-germ line association begins during embryogenesis and continues until the mature egg is formed. In the adult, the somatic escort cells promote differentiation of the germline stem cell daughter using Wnt signaling. dWnt4, a Wnt ligand, acts in an autocrine manner in these escort cells, using the canonical pathway to regulate survival, division and
encapsulation of the stem cell daughter, a function critical for differentiation. Here, we show at an earlier stage, in the larvae, the same ligand uses components of Wnt non-canonical pathway, RhoA, Rac1 and cdc42, to regulate proper mingling of escort cell precursors between the germ cells. Thus, dWnt4 uses different modules of signaling at different points in development to promote cell movement and control cytoplasmic protrusions. As Wnts have been associated with cancers, understanding how Wnts modulate cell movement by switching on and off different modules may lead to insights into the etiology and progression of cancers.

Introduction

Stem cell self-renewal and differentiation are critical for maintaining the organ systems of multicellular organisms. Loss of stem cell self-renewal leads to aging, due to an inability to replenish these organs; while loss of differentiation leads to tumors, which can progress toward diseases such as cancer [1–3]. Thus, identifying triggers of stem cell differentiation is pivotal for understanding the etiology of degenerative diseases and cancer. Germline stem cells (GSCs) self-renew and differentiate to produce gametes [4–7]. The balance between self-renewal and differentiation is critical for a steady supply of gametes for increased reproductive success. Dysregulation of GSC self-renewal and differentiation manifests itself as changes in fecundity without markedly altering the organism’s growth or survival. Additionally, processes regulating GSC differentiation are conserved in other stem cell systems [8–10]. Therefore, GSCs make an excellent system to identify triggers of stem cell differentiation.

Drosophila GSCs are well characterized and genetically tractable [5]. GSC development starts during the larval stages. The female larval gonad is made up of the somatic niche and primordial germ cells (PGCs) [11,12]. The late larval somatic niche is comprised of intermingled cells (ICs), the terminal filament and the cap cells [11–14]. The PGCs that give rise to the GSCs in the adults are interspersed with ICs, precursors of the adult escort cell [11,15,16] (Fig 1A). Most of the PGCs remain undifferentiated during the larval stage [15,17,18]. As development progresses, the larval gonad transforms first into a pupal gonad and then into an adult ovary. During this transition, the PGCs closest to the cap cells acquire a stem cell fate and become GSCs while the rest directly differentiate [15,18]. Each adult ovary consists of 16–18 individual units called ovarioles. GSCs are located in the germarium, present at the anterior end of each ovariole [19]. GSCs divide to give rise to a self-renewed GSC and a stem cell daughter, the cystoblast (CB) (Fig 1B). The CB expresses a differentiating factor called Bag of marbles (Bam), undergoes differentiation and four incomplete divisions to form a sixteen-cell cyst. Of the sixteen cells, fifteen become nurse cells and one is specified as an oocyte [20–24] (Fig 1B). Loss of GSC differentiation results in loss of or delayed progression towards becoming an oocyte.

Soma-germ line interaction is critical for proper GSC self-renewal and differentiation [8,25,26]. Both PGCs in the larval gonad and GSCs in the adults are surrounded by somatic cells that constitute the somatic niche. Close contact and coordinated signaling between the surrounding somatic niche and the germ line is pivotal for self-renewal and differentiation [26,27]. In the larval stages, ICs regulate PGC proliferation [11]. Additionally, proper intermingling of ICs in the larval gonad promotes proper escort cell function in the adults [28,29]. In the adults, the somatic niche can be broadly divided into two regions; the self-renewal niche and the differentiation niche [26,30]. The terminal filament and cap cells comprise the self-renewal niche that is required for regulating GSC self-renewal (Fig 1B) [24,26,31–36]. Loss of
adherens junction proteins, such as DE-Cadherin and Armadillo/β-catenin, in either germline or somatic cells within the self-renewal niche, leads to loss of contact and thus loss of GSC self-renewal [37,38]. The differentiation niche, formed by the escort cells, makes extensive contact with CBs by means of escort cell protrusions that encapsulate the CB and promote GSC differentiation. The downstream Wnt non-canonical components are required in the escort cells for proper germline stem cell differentiation. (A) Schematic of a female late larval gonad. The primordial germ cells (PGCs) (blue) are interspersed with the intermingled cells (ICs) (red), precursors of escort cells. (B) A schematic of the Drosophila female gerarium present at the anterior end of the ovarioles. The germ line consists of germline stem cells (GSCs) (blue) that are attached to the self-renewal somatic niche made by the terminal filament and cap cells (orange). The GSCs divide to give rise to the cystoblast (blue) that differentiates into the escort cells (red) by means of escort cell protrusions that encapsulate the CB and promote GSC differentiation. (C) An illustration of the dWnt non-canonical pathway. dWnt binds to its receptor, Frizzled/Frizzled2 (Fz/Fz2) and activates DAAM1, RhoA, Rac1, and cdc42, downstream of Dsh. RhoA activates ROCK. These pathways are required for actin remodeling, cell movement and cell polarity. (D-I) Germaria of c587-GAL4 (control) and dsh, DAAM1, RhoA, Rac1 and cdc42 depleted escort cells stained with 1B1 (red) and Vasa (blue) showing an accumulation of >3 undifferentiated cells (yellow line). (J) Percentage of the germaria with >3 spectroso mes in dsh, DAAM1, RhoA, Rac1 and cdc42 depleted escort cells showing a significant difference in mutants (n = 50). (K-P) dWnt4 heterozygote, dWnt4/dsh1, dWnt4/DAAM, dWnt4/RhoA, dWnt4/Rac1 and dWnt4/cdc42 trans-heterozygote stained with 1B1 (red) and Vasa (blue) showing an accumulation of >3 undifferentiated cells (yellow line) in the trans-heterozygotes. (Q) Percentage of the differentiation defects in dWnt4 heterozygote, dWnt4/dsh1, dWnt4/DAAM, dWnt4/RhoA, dWnt4/Rac1 and dWnt4/cdc42 trans-heterozygote showing a significant difference in trans-heterozygotes (n = 50). Scale bar for all images is 20 μm. 

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differentiation (Fig 1B) [39–41]. Thus, soma-germline contact throughout development is a critical extrinsic cue that controls GSC differentiation.

Signaling from the somatic niche coordinates the balance of self-renewal and differentiation. In the larval gonad, EGF/Spitz signaling from the PGCs is required for IC survival and intermingling [11]. In the adult germaria, decapentaplegic (dpp) signaling in the self-renewal niche regulates GSC self-renewal by phosphorylating the transcriptional regulator, mothers against dpp (pMAD), to repress expression of the differentiation factor, Bam [20,24,31]. Ecdysone and dWnt4, among others, have been shown to function in the differentiation niche to promote the differentiation of the CB. Ecdysone signaling in the escort cells regulates the formation of escort cell protrusions through an unknown mechanism [42,43]. dWnt4 autocrine signaling in the escort cells is required for escort cell survival, repression of dpp expression and for formation of escort cell protrusions to encapsulate the CB [44–46].

Canonical Wnt signaling plays a critical role in the adult differentiation niche. Wnts can signal through either a canonical or a non-canonical pathway to affect downstream changes [47]. In the escort cells, dWnt4 is known to regulate differentiation through the canonical pathway [44–46]. dWnt4 binds to the receptor, Frizzled2 and co-receptor, Arrow (Arr), and relays a signal to the multidomain protein, Dishevelled (Dsh), leading to stabilization of β-catenin [47–50]. β-catenin, a cytoskeletal protein that is also a transcription factor, translocates into the nucleus and initiates transcription of downstream targets including Frizzled3 (Fz3) [51–57]. Depletion of any of the canonical pathway factors, such as β-catenin or co-receptor arr, specifically in the escort cells leads to loss of CB differentiation by regulating escort cell number [44–46]. Additionally, loss of dWnt4 leads to downregulation of a Wnt canonical reporter, Fz3 promoter fused to RFP (Fz3RFP), and adhesion molecules, β-catenin, Innexin2 and DE-Cadherin [46]. These adhesion molecules promote escort cell encapsulation of the CB that is required for its differentiation [46]. Thus, dWnt4 uses components of the canonical pathway to control both the number and function of escort cells.

Wnt ligands, such as dWnt4, can modulate the Planar Cell Polarity (PCP) of cells [58]. Polarity within a plane of individual cells or of tissues is known as PCP. PCP can specify the proximal and distal end of a cell, as in the Drosophila wing, or the dorsal-ventral and anterior-posterior side of a cell, as in the Drosophila eye [58–64]. Two systems are known to independently regulate PCP—the core components and the Daschous (Ds)/Fat (Ft) system [65,66]. The core components consist of six proteins—Frizzled (Fz), Dishevelled (Dsh), Diego (Dgo), Van Gogh (Vang)/Strabismus (Stbm), Prickle (Pk), and Flamingo (Fmi). These proteins form two complexes that are asymmetrically arranged on the opposite ends of a cell—the Fz, Dsh and Dgo, and the Vang/Stbm, Pk complex. Fmi is enriched on both sides of the cell and stabilizes the complexes [59–61,67]. Small Rho GTPases such as Ras homolog gene family, member A (Rhoa), Ras-related C3 botulinum toxin substrate 1 (Rac1) and Cell division control protein 42 (cdc42) can function as the downstream effectors of the core complex [60,68–70]. The Ds/Ft system consists of two atypical transmembrane cadherin proteins—Ds and Ft. Ds and Ft are present on opposite ends of each cell and bind to each other at cell-cell junctions. A golgi associated kinase, Four Jointed (Fj) phosphorylates the extracellular domains of Ds and Ft and thereby promotes their binding [66,71]. The role of the core components and the Ds/Ft system has been well established in specifying PCP in the wing and the eye.

Binding of Wnt to Frizzled2, can also activate pathways downstream of Dsh: Dishevelled Associated Activator of Morphogenesis 1 (DAAM1), Ras homolog gene family member A (Rhoa), Rac1 and cdc42 pathway. This signaling cascade is also referred to as Wnt/Fz non-canonical pathway. Activated Rhoads activates Rh associated kinase (ROCK/dRok) [47,61,72–75]. Together, these pathways regulate the cell division, cytoskeleton, cell movement and cell polarity [76–81] (Fig 1C). It has been previously shown that, like dWnt4 mutants, flies carrying
mutations in \(dsh^1\), a PCP specific allele, show disrupted oogenesis [82]. In contrast, loss of other genes that regulate PCP such as \(fz,\) \(vang/stbm,\) \(fmi,\) \(dgo,\) and \(pk\) do not show such a defect, suggesting that PCP might not be operating during oogenesis [82]. However, it is not known if and how downstream components of the Wnt/Fz non-canonical pathway promote oogenesis.

Here, we find that \(RhoA,\) \(Rac\) genes and \(cdc42,\) the downstream Wnt non-canonical components, but not members of the core components that define PCP, such as \(vang,\) \(ds,\) \(fz\) and \(ft,\) regulate CB differentiation. We find that \(dWnt4\) genetically interacts with the downstream non-canonical components: \(dsh^1,\) \(DAAM1,\) \(RhoA,\) \(Rac1\) and \(cdc42.\) \(RhoA,\) \(Rac1\) and \(cdc42\) are expressed at higher levels and are more active in the ICs of the late larval gonad than in the adult escort cells. Conversely, the Wnt canonical reporter is not expressed in the late larval stages but is expressed in the adults. Consistent with this observation, we find that these downstream non-canonical pathway components are required in the late larval stages for regulating intermingling of ICs with the germ cells but their function is not essential in the adults to regulate differentiation. Additionally, we find that \(dWnt4\) regulates the activity, but not the protein levels, of \(RhoA,\) \(Rac1\) and \(cdc42\) in the larval gonad. Thus, \(dWnt4\) regulates assembly of the somatic differentiation niche of the GSCs by switching the mode of signaling during development.

**Results**

**dWnt4 acts through the downstream non-canonical pathway components to regulate differentiation**

To determine if the Wnt non-canonical pathway components regulate differentiation, we depleted these components in the escort cells. We made use of RNA interference (RNAi) in conjunction with \(c587-GAL4,\) which is expressed in the escort cells, to deplete \(dsh,\) \(DAAM1,\) \(RhoA,\) \(Rac1\) and \(cdc42\) [43,83,84]. Both control and mutant germaria were stained with 1B1 and Vasa. 1B1 marks the endoplasmic reticulum rich organelle, the spectrosome, in undifferentiated GSCs and CBs; and it marks the branched structures, the fusomes, in differentiated cysts [85]. 1B1 also marks somatic cell membranes [86]. Vasa, an RNA helicase, marks the germ line [87]. We assayed for differentiation defects, defined as an accumulation of greater than 3 undifferentiated cells marked by spectrosomes. Depletion of \(dsh,\) \(DAAM1,\) \(RhoA,\) \(Rac1\) and \(cdc42\) resulted in germline differentiation defects compared to control (Fig 1D–1J) (S1A Fig) (Table 1). We also depleted the mediators that define the PCP of a cell utilizing previously validated RNAi lines in conjunction with \(c587-GAL4.\) We found that for \(vang,\) \(ds,\) \(ft\) and \(fz,\) we did not observe any significant differentiation defect [42,88–90] (S1B–S1I Fig) (Table 1). These results are consistent with Cohen et al’s observation that mutations in genes that regulate PCP of a cell such as \(fz,\) \(stbm,\) \(fmi,\) \(dgo\) and \(pk\) do not lead to defects in oogenesis [82]. Thus, we can conclude that downstream components of the Wnt non-canonical pathway such as \(dsh,\) \(DAAM1,\) \(RhoA,\) \(Rac1\) and \(cdc42\) are required in the escort cells for promoting GSC differentiation. Additionally, these results also suggest that the core components that mediate PCP such as \(vang/stbm,\) \(ds,\) \(fz\) and \(ft\) do not play a significant role in the escort cells to regulate GSC differentiation.

\(DAAM1,\) \(RhoA,\) \(Rac1\) and \(cdc42\) are critical proteins required for various basic cellular processes, such as progression of the cell cycle and cell movement [76–80,91]. It is not surprising that they are required in the escort cells for proper function. We asked if these proteins act downstream of \(dWnt4\) to promote differentiation. To answer this, we generated trans-heterozygous flies that contained one genetically reduced copy of \(dWnt4\) and one reduced copy of \(dsh,\) \(DAAM,\) \(RhoA,\) \(Rac1\) or \(cdc42\) [92–96]. \(Dsh\) is a multi-domain protein that uses distinct domains to interact with either the Wnt canonical or the Wnt non-canonical proteins [97].
We used dsh1, a PCP specific allele, to test if dWnt4 uses the downstream components of the PCP system [68,98]. If the downstream PCP components act through the same pathway, then these trans-heterozygotes could exhibit GSC differentiation defects, but if they act parallel or independent of dWnt4, then these trans-heterozygotes should not exhibit differentiation defects. We stained these trans-heterozygote ovaries for 1B1 and Vasa to assay for differentiation defects. Compared to 17.5% (n = 50) of heterozygous dWnt4 germaria, 98% (n = 50) of dsh1/dWnt4 trans-heterozygote, 85% (n = 50) of DAAM/dWnt4 trans-heterozygote, 80% (n = 50) of RhoA/dWnt4 trans-heterozygote, 78% (n = 50) of Rac1/dWnt4 trans-heterozygote, and 68% (n = 50) of cdc42/dWnt4 trans-heterozygote germaria showed accumulation of spectrosomes (Fig 1K–1Q) (S2A–S2E Fig) (Table 1). As Dsh acts downstream of Wnt signaling, we also stained dsh1, RhoA and Rac1 heterozygous, and trans-heterozygous flies with one mutant copy of dsh1 and one mutant copy of RhoA or Rac1 for 1B1 and Vasa. Compared to heterozygous dsh1, RhoA and Rac1 germaria, we found significantly higher number of spectrosomes in dsh1 mutants, dsh1;RhoA trans-heterozygotes, and dsh1;Rac1 trans-heterozygotes

Table 1. Quantitation of undifferentiated cells in mutants compared to control germaria.

| Genotype                  | Spectrosomes | n  | P-value / %Differentiation defect |
|---------------------------|--------------|----|-----------------------------------|
| c587-GAL4                 | 2.9 ± 0.7    | 50 |                                   |
| c587-GAL4>dsh1 RNAi       | 7.0 ± 1.4    | 50 | 2.6966E-29                        |
| c587-GAL4>dDAAM1 RNAi     | 7.8 ± 1.5    | 50 | 4.7793E-38                        |
| c587-GAL4>dRhoA RNAi      | 7.2 ± 2.5    | 50 | 4.6713E-24                        |
| c587-GAL4>dRac1 RNAi      | 4.0 ± 1.5    | 50 | 4.38037E-11                       |
| c587-GAL4>cdc42 RNAi      | 7.2 ± 4.5    | 50 | 6.05032E-12                       |
| c587-GAL4>vangogh RNAi    | 2.6 ± 1      | 50 | 0.194156                           |
| c587-GAL4>daichous RNAi   | 3.0 ± 1.3    | 50 | 0.333138                           |
| c587-GAL4>frizzled RNAi   | 2.5 ± 1      | 50 | 0.0907                             |
| c587-GAL4>fat RNAi        | 2.9 ± 0.8    | 50 | 0.353701                           |
| dWnt4/CyO                 | 2.7 ± 1.1    | 50 | 17.5% Differentiation Defect       |
| dsh1/+                    | 2.8±1.3      | 50 |                                   |
| DAAM/Fm7                  | 2.4 ± 1      | 50 | 16% Differentiation Defect         |
| RhoA/CyO                  | 2.9 ± 1.2    | 50 | 24% Differentiation Defect         |
| Rac1/TM6                  | 2.7 ± 1.1    | 50 | 22% Differentiation Defect         |
| Cdc42/Fm6                 | 3.9 ± 1.7    | 50 | 28% Differentiation Defect         |
| DAAM+/dWnt4/+             | 50 Z-score = 7.1376, P-value = 0 (compared to dWnt4/CyO) | 8.0064, P-value = 0 (compared to DAAM/Fm7) |
| dsh1+/dWnt4/+             | 50 Z-score = 7.1376, P-value = 0 (compared to dWnt4/CyO) | 1.41784E-13 (compared to dsh1/+)
| RhoA+/dWnt4/+             | 50 Z-score = 4.0718, P-value = 0 (compared to dWnt4/CyO) | 3.4338, P-value = 0.0006 (compared to RhoA/CyO)
| Rac1+/dWnt4/+             | 50 Z-score = 4.4236, P-value = 0 (compared to dWnt4/CyO) | 4.2418, P-value = 0 (compared to Rac1/TM6)
| dsh1                      | 4.1 ± 1.5    | 50 | 2.30222E-05 (compared to dsh1/+)
| dsh1/+; RhoA/+            | 4.1 ± 1.3    | 50 | 5.9375E-06 (compared to dsh1/+)
| dsh1/+; Rac1/+            | 4.1 ± 1.5    | 50 | 7.4036E-05 (compared to dsh1/+)
| Cdc42+/dWnt4/+            | 50 Z-score = 4.7801, P-value = 0 (compared to dWnt4/CyO) | 3.0298, P-value = 0.00032 (compared to Cdc42+/Fm6)
(S2D–S2H Fig) (Table 1). Taken together, these results suggest that \( dWnt4 \) can function via \( dsh \) and the downstream Wnt non-canonical components to regulate GSC differentiation.

**Downstream Wnt non-canonical components regulate CB encapsulation to promote differentiation**

If \( dWnt4 \) acts through the downstream Wnt non-canonical components to regulate GSC differentiation, then \( dWnt4 \) mutants and \( RhoA, Rac1 \) and \( cdc42 \) depleted escort cell mutants should phenocopy each other. Loss of \( dWnt4 \) leads to an accumulation of pre-CBs that do not express Bam due to loss of escort cell encapsulation and a reduction of escort cell number [44–46]. To test if depletion of the downstream non-canonical components in the escort cells also results in an accumulation of pre-CBs, we stained control and mutants for pMAD, which marks the GSCs, and used GFP under the control of \( bam \) promoter to analyze bam transcription and hence mark the differentiating progeny [99]. In addition, we also stained control and mutants for pMAD and BamC that marks proper translation of \( bam \) in the differentiating progeny. Similar to \( dWnt4 \) mutants, \( RhoA, Rac1 \) and \( cdc42 \) depleted escort cell mutants showed an accumulation of pMAD negative, and BamGFP and BamC negative cells, suggesting that these mutants accumulate pre-CBs (S3A–S3E Fig) (S4A–S4D2 Fig). Interestingly, in addition to early pre-CBs, \( cdc42 \) mutants also showed an accumulation of BamGFP and BamC positive CBs, suggesting that \( cdc42 \) may have an additional role in promoting CB differentiation.

Pre-CBs that accumulate in \( dWnt4 \) mutants are capable of differentiation upon ectopic expression of \( bam \). To test if the undifferentiated germ cells that accumulate due to depletion of downstream components of PCP in the escort cells are also capable of differentiating, we ectopically expressed \( bam \) by using a transgene that expresses \( bam \) under the control of heat-shock promoter (hs-\( bam \)) [21]. We found, post heat-shock, these mutants showed loss of undifferentiated cells and an accumulation of cysts, as marked by the presence of fusomes comparable to the control (90% for \( c587\text{-GAL4} \); 86% for \( c587\text{-GAL4}>\text{RhoA RNAi} \), P-value = 0.5353; 76% for \( c587\text{-GAL4}>\text{Rac1 RNAi} \), P-value = 0.0629 and 92% for \( c587\text{-GAL4}>\text{cdc42 RNAi} \), P-value = 0.7263 [for all \( n = 50 \)] (S5A–S5E Fig). We also analyzed the expression of Bruno in \( RhoA, Rac1 \) and \( cdc42 \) depleted escort cell mutants that carry the hs-\( bam \) transgene, without heat-shock and post heat-shock. Bruno, a translational repressor, is expressed at very low levels in the undifferentiated cells but is expressed at high levels post-differentiation from 16-cell cysts onwards [100,101]. We found, post heat-shock, cysts in \( Rac1 \) and \( cdc42 \) mutants that carry the hs-\( bam \) transgene expressed Bruno, while cysts in \( RhoA \) mutants carrying the hs-\( bam \) transgene only weakly expressed Bruno (S5F–S5M1 Fig). Altogether, these results suggest that the downstream Wnt non-canonical pathway components are required extrinsically in escort cells, to promote differentiation in the germ line.

Loss of escort cell number and their encapsulation results in loss of pre-CB differentiation [39–41,102]. To determine if loss of the downstream Wnt non-canonical pathway components leads to loss of encapsulation, we visualized the cytoplasmic protrusions using FaxGFP, a somatic cell membrane marker [39]. It has been previously demonstrated that overexpression in the escort cells of dominant-negative Rho (Rho\( ^{DN} \)), that disrupts Rho function, leads to loss of escort cell encapsulation [40]. We found that, similar to \( dWnt4 \) mutants and germaria where Rho\( ^{DN} \) was overexpressed in the escort cells, depletion of \( RhoA, Rac1 \) and \( cdc42 \) in the escort cells also resulted in loss of encapsulation (Fig 2A–2D1). To determine if this was due to decreased number of escort cells, we counted the number of Tj positive escort cells in these mutants. We found that these mutants exhibited significantly lower number of escort cells (16.5 ± 2.1 for \( c587\text{-GAL4} \); 9.9 ± 3.7 for \( c587\text{-GAL4}>\text{RhoA RNAi} \), P-value = 0.00013; 11 ± 1.5 for \( c587\text{-GAL4}>\text{Rac1 RNAi} \), P-value = 4.1403E-06 and 9.3 ± 2.2 for \( c587\text{-GAL4}>\text{cdc42 RNAi} \),
P-value = 9.5478E-07 (for all n = 10)). However, the escort cells that were present in the ger- maria failed to extend cytoplasmic protrusions. These results demonstrate that like dWnt4, downstream Wnt non-canonical pathway components in the escort cells regulate both encapsu- lation and their numbers, and thus CB differentiation [44–46]. These results taken together suggest that dWnt4 acts through Dsh and the downstream Wnt non-canonical pathway com- ponents to regulate differentiation.

Components of Wnt non-canonical pathway are expressed at high levels and are active in the larval gonad

Wnt proteins are known to regulate signaling through either the canonical or the non-canoni- cal pathway [47]. Surprisingly, our data suggests that dWnt4 also acts through the non-canoni- cal pathway components in the escort cells. There is little precedent for canonical and non- canonical Wnt pathways acting in the same cell at the same time. We therefore hypothesized that either dWnt4 signals via the canonical or non-canonical arm in distinct subsets of escort cells or at different stages of escort cell development. If there are subsets of cells that respond

Fig 2. The downstream Wnt non-canonical pathway components are required in the escort cells for cystoblast encapsulation. (A–D1) Germaria of c587-GAL4, RhoA, Rac1 and cdc42 depleted escort cells stained with Tj (red), GFP (green) and Vasa (blue) showing loss of encapsulation in RhoA, Rac1 and cdc42 depleted escort cells. Fax-GFP marks somatic cell membranes. GFP channel is shown in A1, B1, C1 and D1. Scale bar for all images is 20μm.

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to canonical signaling but not to non-canonical signaling, then we could observe the canonical reporter to be present only in some escort cells. If the regulation is temporal, we hypothesized that we could see differences in either levels or activity of canonical and non-canonical reporters as a function of development.

To test these hypotheses, we analyzed the expression patterns of downstream Wnt non-canonical pathway components and a Wnt canonical reporter during late larval stages, late pupal gonads and in the adult ovaries. Transgenic lines with GFP tagged RhoA, Rac1 or cdc42 that report endogenous expression were used for the Wnt non-canonical components [103]. These reporters were stained for their respective fluorescent proteins and Tj, which marks ICs in the larval gonad and all somatic cells, except for the terminal filament, in the adult gonad [29]. Vasa was used to mark the germ line. We found that RhoA, Rac1 and cdc42 showed high expression in the larval gonad (Fig 3A–3B1 and 3D) (S6A–S6D1 Fig). In contrast, we found that RhoA, Rac1 and cdc42 were expressed at low levels in the escort cells of the pupal and adult germaria (Fig 3C and 3D) (S6E–S6I1 Fig). This different expression pattern suggests that the Wnt non-canonical pathway members are regulated in a temporal manner in the escort cells and their precursors during oogenesis.

RhoA, Rac1 and cdc42 are members of the Rho family of small GTPases. These proteins are inactive when bound to GDP and active when bound to GTP [104,105]. The active forms of these proteins regulate various downstream signaling cascades [76,77,106,107]. We observed that downstream components of the Wnt non-canonical pathway, RhoA, Rac1 and cdc42 are expressed at high levels in the larval gonad. We asked if these proteins are present in their activated form in the larval gonad. To test this, we performed live-imaging of larval gonads of transgenic lines that report the active form of RhoA/Rac and cdc42 [108]. In these biosensors, GFP is fused to the Rho family GTPase Binding Domain (RBD) of downstream effector proteins. Expression of GFP suggests the presence of active forms of these proteins. For RhoA/Rac activity, the downstream effector protein, Protein Kinase N (Pkn) was fused to GFP and placed under squash (sqh) promoter [108]. For cdc42 activity, the cdc42 binding domain of Wiskott-Aldrich Syndrome protein (WASp) was fused with GFP and this was placed under the control of sqh promoter [108]. In order to observe the germ cells in these biosensors, we used kusabira-orange (KO) fused germline marker, Vasa-KO [109,110]. Similar to the expression pattern of the GFP tagged RhoA, Rac1 and cdc42 we observed the presence of active form of RhoA, Rac1 and cdc42 in the ICs of the larval gonad (Fig 3E–3F2). We also asked if these proteins are present in their active forms in the escort cells of the adult germaria. We found that RhoA/Rac activity was attenuated and cdc42 was present in its activated state in the escort cells, albeit at lower levels compared to the larval gonad (S7A–S7C Fig). Together, these results demonstrate that the Wnt non-canonical components are not only present, but also active in the ICs. To determine the activity of the canonical pathway in the larval gonad, we used Fz3RFP [57]. We found that while Fz3RFP showed no expression or was expressed at background levels in the ICs of the larval gonad, it was expressed at higher levels in the escort cells of pupal and adult stages [46,111,112] (Fig 3G–3I) (S7D–S7D1 Fig). Additionally, all the escort cells in the adult germarium expressed the canonical reporter (n = 10, 99.5% Tj positive cells expressed Fz3RFP). Thus, we concluded that Wnt canonical and Wnt non-canonical pathway components, and their activity, are temporally regulated in escort cells and their precursors during oogenesis.

**Components of the Wnt non-canonical pathway play a critical role in the larval gonad**

To determine if the different expression patterns of the downstream Wnt non-canonical pathway components mirrored their role in the two developmental stages, we used the UAS-GAL4
system to deplete genes using RNAi in the escort cells at specific developmental time points. The UAS-GAL4 system is temperature dependent and therefore, different temperatures can be used to attenuate genes at various developmental time points. Maximal RNAi activity is attained at 29°C whereas minimal RNAi activity is attained at 18°C [113]. c587 is expressed in the larval gonad, including the ICs, but is only expressed in the escort cells of the adult inner germarium (S8A–S8C1 Fig). To elucidate if the downstream Wnt non-canonical pathway components are required in the adults to regulate CB differentiation, the flies were kept at 18°C until they eclosed. Once eclosed, the flies were shifted to 29°C and kept at this temperature for 7 days [114] (Fig 4A). The germaria were then stained for 1B1 and Vasa. This strategy has been previously used to show that the dWnt4 canonical pathway is required in the adults to

Fig 3. The downstream Wnt non-canonical pathway components are temporally regulated. (A–B1) Larval gonad of transgenic fly with GFP tagged to RhoA stained for Tj (red), Vasa (blue) and GFP (green) showing high expression of RhoA in the ICs. 63x of larval gonad is shown in B and B1. GFP channel is shown in A1 and B1. (C–C1) Adult germarium of transgenic fly with GFP tagged to RhoA stained for Tj (red), Vasa (blue) and GFP (green) showing low expression of RhoA in the adult escort cells. GFP channel is shown in C1. (D) Quantification (n = 7) of GFP in the ICs, pupal escort cells and adult escort cells showing that the downstream Wnt non-canonical components are highly expressed in the ICs in comparison to the adult escort cells. (E–F2) Larval gonads of transgenic flies that report active form of RhoA/Rac (E–E2) and cdc42 (F–F2). While RhoA/Rac is active in the ICs, cdc42 is highly active in the ICs and terminal filament and comparatively less active in the Primordial Germ Cells (PGCs). Germline is marked by Vasa-KO. GFP channels and KO channels are shown in E1, E2, F1 and F2. (G–G1) Larval gonad stained for 1B1 (green), Vasa (blue) and RFP (red) showing low expression of the Wnt canonical reporter, Frizzled3 in the ICs. RFP channel is shown in G1. (H–H1) Adult germarium stained for Tj (green), Vasa (blue) and RFP (red) showing high expression of Frizzled3 in the adult escort cells. RFP channel is shown in H1. (I) Quantification (n = 7) of RFP in the ICs, pupal escort cells and adult escort cells showing that the canonical reporter is expressed at higher levels in the adult escort cells in comparison to the ICs. Scale bar for B and B1 is 10μm. Scale bar for all other images is 20μm.

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regulate CB differentiation [45]. We depleted dWnt4, RhoA, Rac1, cdc42 and also expressed cdc42\textsubscript{DN} in the escort cells to assay for differentiation defects. We found that compared to control, 18 °C-29 °C temperature shift dWnt4 mutants exhibited a differentiation defect (Fig 4B and 4C and Fig 4G). In contrast, compared to control, 18 °C-29 °C temperature shift RhoA, Rac1, cdc42 and cdc42\textsubscript{DN} mutants did not exhibit a significant accumulation of undifferentiated cells, suggesting that the downstream Wnt non-canonical pathway components do not play a critical role in the adult to regulate CB differentiation. We depleted dWnt4, RhoA, Rac1, cdc42 and also expressed cdc42\textsubscript{DN} in the escort cells to assay for differentiation defects. We found that compared to control, 18 °C-29 °C temperature shift dWnt4 mutants exhibited a differentiation defect (Fig 4B and 4C and Fig 4G). In contrast, compared to control, 18 °C-29 °C temperature shift RhoA, Rac1, cdc42 and cdc42\textsubscript{DN} mutants did not exhibit a significant accumulation of undifferentiated cells, suggesting that the downstream Wnt non-canonical pathway components do not play a critical role in the adult to regulate CB differentiation. We depleted dWnt4, RhoA, Rac1, cdc42 and also expressed cdc42\textsubscript{DN} in the escort cells to assay for differentiation defects. We found that compared to control, 18 °C-29 °C temperature shift dWnt4 mutants exhibited a differentiation defect (Fig 4B and 4C and Fig 4G). In contrast, compared to control, 18 °C-29 °C temperature shift RhoA, Rac1, cdc42 and cdc42\textsubscript{DN} mutants did not exhibit a significant accumulation of undifferentiated cells, suggesting that the downstream Wnt non-canonical pathway components do not play a critical role in the adult to regulate CB differentiation. We depleted dWnt4, RhoA, Rac1, cdc42 and also expressed cdc42\textsubscript{DN} in the escort cells to assay for differentiation defects. We found that compared to control, 18 °C-29 °C temperature shift dWnt4 mutants exhibited a differentiation defect (Fig 4B and 4C and Fig 4G). In contrast, compared to control, 18 °C-29 °C temperature shift RhoA, Rac1, cdc42 and cdc42\textsubscript{DN} mutants did not exhibit a significant accumulation of undifferentiated cells, suggesting that the downstream Wnt non-canonical pathway components do not play a critical role in the adult to regulate CB differentiation.
CB differentiation, the flies were kept at 29 °C until they eclosed. Once eclosed, the flies were shifted to 18 °C and kept at this temperature for 7 days [114] (Fig 4H). Compared to control, 29 °C-18 °C temperature shift dWnt4, RhoA, Rac1, cdc42 and cdc42^DN^ mutants resulted in an accumulation of undifferentiated cells (2.5 ± 1 for c587-GAL4; 5.6 ± 2 for c587-GAL4>dWnt4 RNAi, P-value = 1.0481E-17; 5 ± 1.5 for c587-GAL4>Rhoa RNAi, P-value = 1.43965E-15; 5 ± 1.5 for c587-GAL4>Rac1 RNAi, P-value = 1.55386E-18; 5 ± 2.5 for c587-GAL4>cdc42 RNAi, P-value = 2.34120E-18 and 5 ± 1 for c587-GAL4>cdc42^DN^, P-value = 1.49752E-17) (for all n = 50) (Fig 4I–4N) (S9C–S9F and S9I Fig). Taken together, these results demonstrate that the Wnt non-canonical pathway components act primarily prior to the adult stage to regulate CB differentiation.

As RhoA, Rac1 and cdc42 are highly expressed and active in the larval gonad and were not required in the adult germaria for CB differentiation, we asked if these proteins are required earlier in development. To test this we used an IC driver traffic-jam-GAL4 (tj-GAL4). Tj is expressed only in ICs in the late larval gonad, therefore tj-GAL4 is a more restricted driver for ICs [11,29]. We depleted dsh, DAAM1, RhoA, Rac1, cdc42 and expressed RhoA^DN^ and cdc42^DN^ versions in the ICs, in the late larval stages using tj-GAL4. Because we posited that RhoA, Rac1 and cdc42 act downstream of dWnt4, we also monitored the late larval gonads of dWnt4 mutants for defects, by staining for Vasa, Tj and 1B1. We found that compared to dWnt4 heterozygotes, dWnt4 mutants and dsh, DAAM1, RhoA, RhoA^DN^, Rac1, cdc42 and cdc42^DN^ mutants showed loss of intermingling of ICs with PGCs (n = 25) (Fig 5A–5H) (S10A–S10C Fig). Although loss of Rac1 in ICs led to loss of intermingling, the phenotype was weaker than dWnt4 mutants, RhoA and cdc42 depleted ICs. Additionally, depletion of Rac1 in the ICs resulted in a decrease in the size of the larval gonad. It has been shown Rac1, Ras-related C3 botulinum toxin substrate 2 (Rac2) and Mig-2-like (Mtl) act redundantly to regulate PCP [115]. For this reason, we also depleted Mtl and Rac2 specifically in the ICs and found that Mtl depleted ICs resulted in a strong loss of ICs intermingling but Rac2 depleted ICs did not exhibit any intermingling defect (Fig 5I and 5J). Although we find that depletion of Rac2 does not lead to a differentiation defect, we do not know if this is due to a feeble RNAi mediated depletion, or in fact Rac2 does not play a role. These results suggest that dsh, DAAM1, RhoA, Rac1, Mtl and cdc42 regulate intermingling of ICs in the larval gonad.

In order to further validate our results obtained by RNAi and expression of dominant negative forms of RhoA and cdc42, we also analyzed larval gonads of dsh^1^ mutants, cdc42^2^, a hypomorphic allele that has been previously shown to exhibit mild PCP defects, cdc42 (cdc42^2^/cdc42^2^) hypomorphic mutant and heterozygous larval gonads of flies having one genetically reduced copy of all three Rac genes [115]. We found that these mutants and the heterozygous larval gonad for flies that remove all three Rac genes also exhibited loss of ICs intermingling (69% dsh^1^ mutants, n = 20; 55% cdc42^5^ mutants, n = 20; 80% cdc42 (cdc42^2^/cdc42^2^) mutants, n = 20 and 60% heterozygous Rac1, Rac2 and Mtl larval gonads, n = 10) (S10D–S10H Fig). Together, these results suggest that RhoA, Rac genes (Rac1 and Mtl) and cdc42 regulate intermingling of ICs.

**Components of the Wnt non-canonical pathway regulate IC number**

As RhoA, Rac1 and cdc42 are known to play a role in cell division, we wondered if there were additional defects apart from loss of intermingling in these mutants. We found that compared to dWnt4 heterozygotes and tj-GAL4, dWnt4 mutants and RhoA, Rac1, Mtl and cdc42 mutant larval gonads had significantly fewer ICs (151 ± 18 for dWnt4/CyO; 84 ± 8 for dWnt4 mutants, P-value = 6.74402E-05; 213 ± 36 for tj-GAL4; 36 ± 13 for tj-GAL4>Rhoa RNAi, P-value = 5.16854E-06; 67 ± 14 for tj-GAL4>Rac1 RNAi, P-value = 9.83637E-05; 111 ± 10 for tj-GAL4>Mtl RNAi, P-value = 6.74402E-05; 111 ± 10 for tj-GAL4>Mtl RNAi, P-value = 6.74402E-05).
RNAi, P-value = 0.000231 and 61 ± 17 for tj-GAL4>;cdc42 RNAi, P-value = 2.07541E-05 (for all n = 5). It is possible that the decrease in the number of ICs is either due to decrease in their division rate or due to death of ICs. To test if loss of dWnt4, RhoA, Rac1, Mtl and cdc42 affects the division rate of the ICs, we stained these mutants with a mitotic marker, Phospho-Histone 3 (PH3), Tj and 1B1. We observed that RhoA, Rac1, Mtl and cdc42 depleted IC mutants show a decrease in the number of ICs expressing PH3, suggesting that RhoA, Rac1, Mtl and cdc42 also regulate division of the ICs. However, dWnt4 mutants did not exhibit this defect compared to heterozygous, but displayed a defect compared to wild type (WT) control (12.5 ± 2 for WT control; 4.5 ± 2 for dWnt4/Cyo, P-value = 1.72943E-08; 3 ± 2 for dWnt4 mutants, P-value (WT) = 7.42348E-09, P-value (dWnt4/Cyo) = 0.104558; 9 ± 3 for tj-GAL4, n = 10; 4 ± 2 for tj-GAL4>;RhoA RNAi, n = 10, P-value = 0.00010; 3 ± 2 for tj-GAL4>;Rac1 RNAi, n = 10, P-value = 2.56584E-05; 7 ± 2 for tj-GAL4>;Mtl RNAi, n = 10, P-value = 0.035212, and 5 ± 3 for tj-GAL4>;cdc42 RNAi, n = 10, P-value = 0.003773) (S11A–S11H Fig). To determine if the ICs in these mutants show increased cell death, we stained them for a death marker, cleaved Caspase3, along with Tj and 1B1. We observed that only ICs of RhoA mutants showed Caspase3 staining.
Compared to 0% of Tj positive cells in control, 3% of Tj positive cells in RhoA depleted IC mutants exhibited Caspase3 staining indicating that in addition to its role in cell division, RhoA also regulates IC survival (n = 5) (S11I–S11O Fig). Together, these results show that dWnt4 and the downstream components of the Wnt non-canonical pathway, RhoA, Rac1, Mll and cdc42 play a critical role to regulate intermingling and cell division of ICs in the larval gonad. Additionally, as RhoA depletion exhibited a stronger defect than dWnt4 mutants it is likely that it has roles independent of dWnt4 signaling.

dWnt4 mutants and RhoA, Rac1 and cdc42 depleted escort cell germaria exhibit reduced number of escort cells and loss of CB encapsulation [46]. Additionally, depletion of RhoA, Rac1 and cdc42 in the ICs resulted in reduced number of ICs. We asked if reduction in the escort cell number and loss of encapsulation of CB in the adults is a consequence of reduced number of ICs in the larval gonad. To answer this question, we counted the number of Tj positive escort cells in the adult flies, of 29°C-18°C temperature shift escort cell depleted RhoA, Rac1 and cdc42 mutants. We found that these mutants exhibited significantly lower number of escort cells (15.4 ± 2.6 for c587-GAL4>RhoA RNAi, P-value = 0.00048; 10.6 ± 2 for c587-GAL4>Rac1 RNAi, P-value = 0.00068 and 6.9 ± 1.6 for c587-GAL4>cdc42 RNAi, P-value = 8.1045E-08 (for all n = 10)). These results suggest that RhoA, Rac1 and cdc42 act in the ICs to regulate the escort cell number and therefore, proper GSC differentiation in the adults.

Wnt canonical pathway and PCP regulators do not regulate the intermingling of ICs in the larval gonad

To determine if PCP core components regulate intermingling in the larval gonad we depleted vang/stbm, ds, fz and ft in ICs and found that depletion of these genes did not exhibit any intermingling defects (S12A–S12E Fig). To ascertain that the canonical pathway is not required in the larval gonad, we depleted the canonical specific co-receptor, arr in ICs and found that these mutants also did not exhibit any intermingling defect (S12A and S12F Fig). We used arr as β-catenin has roles outside of Wnt signaling and loss of arr in the adult escort cells results in CB differentiation defects [45,116]. These results suggest that the Wnt canonical pathway and vang/stbm, ds, fz and ft, the core components and Ds/Ft systems that define PCP, do not play a critical role in the ICs.

dWnt4 regulates RhoA, Rac and cdc42 activity in the larval gonad

Similar to dWnt4 mutants, RhoA, Rac1 and cdc42 depletion in the ICs showed a defect in intermingling of ICs. To determine if RhoA, Rac1 and cdc42 act downstream of dWnt4, we analyzed the expression and activity of RhoA, Rac1 and cdc42 with the help of the GFP tagged lines, in larval gonads of both control and dWnt4 mutants. We found that compared to the control, expression of RhoA, Rac1 and cdc42 was not altered in the ICs of dWnt4 mutants, suggesting that dWnt4 does not regulate the protein levels of RhoA, Rac1 and cdc42 (P-value = 0.64182 for RhoAGFP, P-value = 0.79908 for Rac1GFP, and P-value = 0.63675 for cdc42GFP in dWnt4 mutants) (S13A–S13G Fig). The activity level of RhoA/Rac and cdc42 was determined using biosensors in dWnt4 mutants. To ensure that there is no background, we also imaged larval gonads, without the biosensors (S13H–S13I1 Fig). We found that RhoA/Rac and cdc42 activity was significantly downregulated in ICs (P-value = 0.02067, n = 25 for RhoA/Rac activity and P-value = 0.01369, n = 25 for cdc42 activity) (Fig 6A–6F). We also observed cdc42 activity in the terminal filaments and found that it was not altered in dWnt4 mutants (P-value = 0.35901, n = 3) (S13J Fig). These results suggest that dWnt4 regulates the activity of RhoA, Rac and cdc42 in the ICs to regulate intermingling of ICs with PGCs.
Here, we find that *dWnt4* regulates CB differentiation through the downstream non-canonical pathway components. We show that RhoA, Rac1 and cdc42 are expressed at high levels in the ICs of the larval gonad and are active while the expression of Fz3RFP, a β-catenin-dependent transcriptional reporter, is not detectable. Conversely, we find low levels of RhoA, Rac1 and cdc42 in the adults but the β-catenin-dependent canonical reporter is expressed at high levels. Consistent with this, we find a role for RhoA, Rac1 and cdc42 but not the β-catenin-dependent canonical pathway in the larval gonad for promoting intermingling of ICs (Fig 7). Our results, in conjunction with Mottier-Pavie *et al*. and Wang *et al.*, that show the canonical pathway is required in the adult, point towards a switch from utilizing Wnt non-canonical components to utilizing a canonical pathway to regulate the formation of the differentiation niche [44,45].
Our findings that the depletion of vang/stbm, ds, fz and ft in either ICs or escort cells does not lead to differentiation defects, suggests that PCP does not play a critical role in regulating CB differentiation. These results are consistent with previous literature, where Cohen et al. showed that mutations in, fz, fmi, dgo, vang/stbm and pk, genes required for PCP of a cell, do not lead to significant disruption of ovarian morphology [82]. We show that the other PCP system components, ds and ft, are also not required to regulate differentiation. Together, Cohen et al. and our results suggest that it is not the PCP system, but the downstream Wnt non-canonical components that play a critical role in the differentiation niche. We propose that dWnt4 uses the downstream Wnt/Fz non-canonical pathway to regulate formation of the differentiation niche.

In the larval gonad, dWnt4 uses RhoA, Rac1, Mtl and cdc42 to regulate intermingling, division and number of the ICs while in the adult, dWnt4 uses the β-catenin dependent pathway to regulate the cytoplasmic protrusions.

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Our findings that the depletion of vang/stbm, ds, fz and ft in either ICs or escort cells does not lead to differentiation defects, suggests that PCP does not play a critical role in regulating CB differentiation. These results are consistent with previous literature, where Cohen et al. showed that mutations in, fz, fmi, dgo, vang/stbm and pk, genes required for PCP of a cell, do not lead to significant disruption of ovarian morphology [82]. We show that the other PCP system components, ds and ft, are also not required to regulate differentiation. Together, Cohen et al. and our results suggest that it is not the PCP system, but the downstream Wnt non-canonical components that play a critical role in the differentiation niche. We propose that dWnt4 uses the downstream Wnt/Fz non-canonical pathway to regulate formation of the differentiation niche.

We find that RhoA, Rac1 and cdc42 regulate intermingling of ICs in the larval gonad. In addition, we find that loss of RhoA leads to death of ICs and loss of Rac1 leads to decrease in the larval gonad size. dWnt4 mutants do not show either of these outcomes. Together, these results suggest that RhoA and Rac1 have roles in the larval gonad independent of Wnt signaling. It has been extensively shown that Rac1, Rac2 and Mtl have redundant roles in axon growth, guidance and PCP in the eye and the wing [93,95,115]. Indeed, we found that Rac1 depleted ICs resulted in loss of intermingling but this phenotype was not as strong as dWnt4, RhoA and cdc42 depleted ICs. Rac2 depleted ICs resulted in no intermingling defect. However, depletion of Mtl in the ICs resulted in a strong loss of intermingling phenotype. This suggests that small GTPases play a critical role in ICs function, both independently and in coordination with Wnt signaling.
In the adult, escort cells extend dynamic cytoplasmic protrusions to encapsulate the GSCs, the differentiating daughter and differentiating cysts, a function necessary for proper differentiation [28,39–41,102,117]. It was recently shown that in adult female germaria, stat regulates the formation of these protrusions through cdc42. Loss of stat or woc, a component of the stat pathway, leads to formation of smaller protrusions instead of longer stable protrusions. Loss of woc also leads to an accumulation of undifferentiated cells. Overexpression of cdc42 in woc mutants rescued the small protrusion defect and partially rescued the phenotype. However, Banisch et al report that expression of dominant active or dominant negative cdc42 in the escort cells only mildly affects protrusions and does not lead to an accumulation of undifferentiated cells. Our temperature shift experiments also suggest that expression of dominant negative cdc42 exclusively in the adults does not result in an accumulation of undifferentiated cells, while its expression prior to eclosion does (S9C and S9D and S9E and S9F Fig). Moreover, Banisch et al. also show that active cdc42 is expressed in the escort cells and active RhoA could not be detected in most of the escort cells using Dia-RBD-GFP but could be detected mainly in the escort cell body and weakly in the escort cell protrusion using Capu-RBD-GFP [117]. We think that cdc42 may have a role in the adults by controlling protrusive activity of escort cells, but it does not affect CB differentiation in a biologically meaningful way. This suggests that cdc42 and Rho activity in the adult is attenuated and their activity is independent of dWnt4.

Cell movement requires dynamic actin-myosin polymerization at the leading edge, driven by Rac1 and cdc42, and actin-myosin contraction at the lagging end, driven by RhoA [118,119]. In our study, we find that both RhoA/Rac and cdc42 are present and active in the ICS. Loss of RhoA, Rac1 and cdc42 leads to loss of intermingling of ICS, suggesting that these proteins drive ICS cell movement and intermingling in the larval gonad. In contrast, only cdc42, but not RhoA, is active in the adult escort cells, consistent with the fact that these cells are stationary [41,117]. Thus the switch in the module used by Wnt signaling to regulate the activity of RhoA/Rac and cdc42 parallels the developmental demands of these cells to first move between germ cells in the larval gonad, then to stop and create long, stable protrusions that promote differentiation in the pupal and adult gonad.

Cancer metastasis requires cell migration. The invasion of metastatic cancer cells requires cells to lose epithelial properties and gain mesenchymal properties. During this process, known as Epithelial-mesenchymal transition (EMT), the epithelial cells, which are otherwise polarized, non-motile and have strong cell-cell interaction, subsequently lose polarity, cell-cell adhesion and become motile [120,121]. RhoA, Rac1 and cdc42 are conserved from lower eukaryotes to mammals and are key players that affect cell division, cytoskeletal rearrangement, cell polarity, and cell motility [76,77,79,80,91,106,118]. It has been previously discovered that during cancer, these proteins are upregulated and therefore help initiate metastasis [122]. We have discovered that RhoA, Rac1 and cdc42 are turned on in the ICS of the larval gonad and switched off in the adult escort cells. This is fascinating because the ICS and the escort cells are essentially the same cells at different developmental time points. A better understanding of how this switch is mediated may give us an insight into cancer and aid in developing mechanisms to block metastasis.

Materials and methods

Fly stocks

The following fly stocks were used in the study: c587-Gal4, trafficJam (tj)-Gal4, dWnt4C1/CyO (6651), bamGFP, arrow RNAi (31313), Rho1 RNAi (Bloomington 32383), UASRho1DN (Bloomington 7327), Rac1 RNAi (Bloomington 28985 and 34910), Mtl RNAi (Bloomington 51932), Rac2 RNAi (v28926), cdc42 RNAi (Bloomington 35756 and 37477), UAScdc42DN (Bloomington
6288), dDAAM1 RNAi (Bloomington 39058, V24885), dsh RNAi (Bloomington 34321), van gogh RNAi (Bloomington v7376), daschous RNAi (Bloomington 28008), fat RNAi (Bloomington 29566), y¹ w'; Rho1¹72F/Cyo (Bloomington 7326), y¹ w67c3; P{EPgy2}Rac¹FM564/TM6B, Tb¹ (Bloomington 15461), y¹[w¹]; Rac1¹J10; Rac2[Δ]; P[w¹+m.w.hs] = FRT[w¹hs]2A Mit[Δ]/TM6B, Tb¹ (Bloomington 6679), y¹ w¹ Cdc42¹/FM6 (Bloomington 7337), y¹ w¹ Cdc42² P{neoFRT¹9A (Bloomington 9105), y¹ w¹ Cdc42³ P{neoFRT¹9A (Bloomington 52298), y¹ Mi{MIC}DAAM¹MI04569 w¹1118/FM7b (Bloomington 38567), w¹ dsh¹ (Bloomington 5298), RhoAGFP (V318439), Rac1GFP (Bloomington 52285), cdc42GFP (V218151), w² P{sqh-Pkn.RBD.G58A-eGFP}312a P{sqh-Pkn.RBD.G58A-eGFP}312b (Bloomington 52298), w² P{sqh-WASp.RBD-GFP}378a P{sqh-WASp.RBD-GFP}378b (Bloomington 56746), UASmCD8GFP (Bloomington 32184), faxGFP, Sco/Cyo;MKRS/TM6 (Lehmann Lab), Sco/Cyo;Nos-Gal4;VP16, Vasa-KO/TM6 (Lehmann Lab); Sco/Cyo;Fz3RFP (Bach Lab).

Collection and Fixation of tissues
3–4 day old fly ovaries were dissected in PBS, fixed for 30 min in PBS plus 5% formaldehyde, incubated for 1 h in PBST (0.2% Tween 20 (Sigma) in PBS) supplemented with 1% Triton X-100 (Sigma), followed by incubation for 2 h in BBT (PBST supplemented with 1% (w/v) bovine serum albumin (BSA; Sigma)). Primary antibodies were added in BBT and incubation was carried out overnight at 4 °C. The following day, ovaries were washed four times for 10 min, 20 min, 30 min in BBT and for 30 min in BBT supplemented with 2% (w/v) donkey serum (Sigma). Secondary antibodies were added in BBT supplemented with 4% (w/v) donkey serum (Sigma) and incubated for 2 h followed by five washes, 10 min each in PBST. VECTASHIELD (Vector Laboratories) with DAPI was added prior to mounting. Fixation and staining of larval gonads was carried out as previously described [123].

Temperature shift experiment
In order to express a specific gene only in the adults, the flies were kept at 18 °C until they eclosed. Once eclosed, the young flies were shifted to 29 °C and kept at this temperature for 7 days. Staining was followed to determine any differentiation defects.

In order to express a specific transgene only in the larval stage, the flies were kept at 29 °C until they eclosed. Once eclosed, the young flies were shifted to 18 °C and kept at this temperature for 7 days. These flies were then dissected and stained for observation.

Unless specified, all experiments that utilized the UAS-GAL4 system were performed constitutively at 29 °C.

Antibodies
Immunostaining of the ovaries and larval gonad was carried out with the following primary antibodies: Mo 1B1 (1:20, DSHB), Rb Vasa (1:5000, Rangan Lab), Ch Vasa (1:500, Rangan Lab), GP Traffic Jam (1:5000, Godt Lab), Rb GFP (1:2000, ab6556), Rb pMAD (1:200, abcam AB52903), Mo BamC (1:200, DSHB), Rb Bruno (1:500, Lehmann Lab), Rb PH3 (1:200, Cell Signaling 97015), and Rb Caspase 3 (1:300, Cell Signaling 96615).

Alexa 488 (Molecular Probes), Cy3 and Cy5 (Jackson Labs) conjugated secondary antibodies were used at a concentration of 1:500.

Fluorescence imaging
The tissues were visualized under 10X, 20X, 40X and 63X objective lenses. The images were acquired using a Zeiss LSM-710 confocal microscope under 20x, 40X and 63x objective.
Live imaging

The tissues were dissected in Schneider’s media and mounted. These were then visualized under 20X objective lens. The images were acquired using a Zeiss LSM-710 confocal microscope under 20x objective.

Quantification analysis

A.U of protein levels. In order to calculate intensities for Fz3RFP and GFP in ICs or escort cells, images for both, control and mutant larval gonad and adult germaria were taken using the same confocal settings. Z stack planes were obtained for all images. Specific planes showing Tj positive ICs or escort cells were chosen, the area of Tj positive cell was outlined and analyzed using the ‘analyze’ tool in ImageJ. The mean and area of the specified region was obtained. An average of all the ratios (Mean/Area), for Fz3RFP, GFP and Tj, per image was calculated for both, control and mutants. The average ratio for Fz3RFP and GFP was normalized with average ratio for Tj. These are the arbitrary units (A.U) for Tj, GFP and RFP for this region. A minimum of 7 germaria and 7 larval gonads (minimum 5 Tj positive cells per image) were considered for the each quantification.

A.U of activity reporters. In order to calculate intensities of activity reporters, images with Z stacks were obtained for both, control and mutant larval gonads using the same confocal settings. For ICs, an area right besides a PGC was outlined and for terminal filament an area in the terminal filament was outlined. All these areas were analyzed using the ‘analyze’ tool in ImageJ. The mean and area of the specified region was obtained (25 cells were analyzed per image). An average of all the ratios (Mean/Area), per image was calculated for control and mutants. The average ratio for activity was subtracted with average ratio for background. The results are the arbitrary units (A.U) for activity reporters for this region. 25 cells of 2 larval gonads each were quantified for RhoA/Rac and cdc42.

In order to calculate intensities of activity reporters between larval gonads and adult germaria, A.U of GFP was normalized to Vasa-KO. 5 cells of 3 larval gonads and adult germaria each were quantitated for RhoA/Rac and cdc42.

The number of Tj and Fz3RFP positive cells were calculated manually in all stacks for 10 germaria in ImageJ.

Statistical analysis

P-values were determined by two-tailed equal variance t test in mutants vs. wild type strains. Z-score was determined by a two-tailed test for data represented in percentage.

Materials and reagents

Fly food was created using the procedures from the Ruth Lehmann lab at NYU (summer/winter mix), and used to fill narrow vials to approximately 12mL.

Supporting information

S1 Fig. Planar cell polarity (PCP) regulators do not play a critical role in the formation of the differentiation niche. (A) Quantification of spectrosomes in c587-GAL4, dsh, DAAM1, RhoA, Rac1 and cdc42 depleted escort cells showing a significant difference in mutants (n = 50). (B) A schematic showing proteins that regulate the PCP of a cell. (C-G) Germaria of c587-GAL4 (control), vang, ds, fz and ft depleted escort cells stained with 1B1 (red) and Vasa (blue) showing no accumulation of undifferentiated cells in vang, ds, fz and ft depleted escort cells (yellow line). (H) Percentage of the germaria with >3 spectrosomes in c587-GAL4, vang,
ds, fz and ft depleted escort cells showing no difference in vang, ds, fz and ft depleted escort cells (n = 50). (I) Quantification of spectrosomes in c587-GAL4, vang, ds, fz and ft depleted escort cells showing no significant difference (n = 50). Scale bar for all images is 20μm.

(TIF)

**S2 Fig. The Wnt downstream non-canonical pathway components are required in the escort cells for proper germline stem cell differentiation.** (A-C) dsh\(^{1/+}\), RhoA/CyO, and Rac1/TM6 stained with 1B1 (red) and Vasa (blue) showing no accumulation of undifferentiated cells (yellow line). (D) Percentage of the germaria with >3 spectrosomes in dsh\(^{1/+}\), RhoA/CyO, Rac1/TM6, dsh\(^{1}\) mutants, dsh\(^{1}\)/RhoA trans-heterozygote and dsh\(^{1}\)/Rac1 trans-heterozygote showing a significant difference in differentiation defects in the trans-heterozygotes (n = 50). (E) Quantification of the number of spectrosomes in dsh\(^{1/+}\), RhoA/CyO, Rac1/TM6, dsh\(^{1}\) mutants, dsh\(^{1}\)/RhoA trans-heterozygote and dsh\(^{1}\)/Rac1 trans-heterozygote showing a significant difference in the number of spectrosomes in the trans-heterozygotes (n = 50). (F-H) dsh\(^{1}\) mutants, dsh\(^{1}\)/RhoA trans-heterozygote and dsh\(^{1}\)/Rac1 trans-heterozygote stained with 1B1 (red) and Vasa (blue) showing an accumulation of >3 undifferentiated cells in the trans-heterozygotes (yellow line). Scale bar for all images is 20μm.

(TIF)

**S3 Fig. The Wnt downstream non-canonical pathway components are required in the escort cells for proper cystoblast differentiation.** (A-D2) Germaria of c587-GAL4 (control), RhoA, Rac1 and cdc42 depleted escort cells stained with pMAD (red), GFP (green) and Vasa (blue) showing an accumulation of >3 pMAD and Bam negative cells in RhoA, Rac1 and cdc42 mutants (yellow line). pMAD channel is shown in A1, B1, C1 and D1; GFP channel is shown in A2, B2, C2 and D2. (E) Quantification of number of pMAD and Bam negative cells in c587-GAL4, RhoA, Rac1 and cdc42 depleted escort cells showing a significant increase in pMAD and Bam negative cells in RhoA, Rac1 and cdc42 mutants (n = 20). Scale bar for all images is 20μm.

(TIF)

**S4 Fig. The Wnt downstream non-canonical pathway components are required in the escort cells for proper BamC expression.** (A-D2) Germaria of c587-GAL4 (control), RhoA, Rac1 and cdc42 depleted escort cells stained with pMAD (red), BamC (green) and Vasa (blue) showing an accumulation of >3 pMAD and BamC negative cells in RhoA, Rac1 and cdc42 mutants (yellow line). pMAD channel is shown in A1, B1, C1 and D1; BamC channel is shown in A2, B2, C2 and D2. Scale bar for all images is 20μm.

(TIF)

**S5 Fig. RhoA, Rac1 and cdc42 act upstream of Bam in the escort cells to regulate cystoblast differentiation.** (A) c587-GAL4 (control) carrying a hs-bam transgene stained with 1B1 (red) and Vasa (blue) without heat-shock. (B-E) c587-GAL4 (control), RhoA, Rac1 and cdc42 depleted escort cell mutants carrying a hs-bam transgene stained with 1B1 (red) and Vasa (blue) showing differentiating cysts marked by fusomes (red) (white arrow) and a lack of undifferentiated cells post heat-shock. (F-I1) c587-GAL4 (control), RhoA, Rac1 and cdc42 depleted escort cell mutants carrying a hs-bam transgene, without heat-shock stained with 1B1 (red), Bruno (green) and Vasa (blue) showing accumulation of undifferentiated cells (yellow line) marked by the presence of spectrosomes (red) and low Bruno expression in the undifferentiated cells and early cysts, while high Bruno expression in 16-cell cyst and onwards. Bruno channel is shown in F1, G1, H1 and I1. (J-M1) c587-GAL4 (control), RhoA, Rac1 and cdc42 depleted escort cell mutants carrying a hs-bam transgene stained with 1B1 (red), Bruno (green) and Vasa (blue), post heat-shock showing Bruno expression in the differentiating cysts.
(white arrow) marked by fusomes (red) and a lack of undifferentiated cells post heat-shock. Cysts in post-heat-shock Rho mutants showed weak Bruno staining. Bruno channel is shown in J1, K1, L1, and M1. Scale bar for all images is 20\(\mu\)m.

(TIF)

S6 Fig. The downstream Wnt non-canonical pathway components, RhoA, Rac1, cdc42 are temporally regulated in the ICs and adult escort cells. (A-D1) Late larval gonad of transgenic flies with GFP tagged to Rac1 and cdc42, respectively stained for Tj (red), Vasa (blue) and GFP (green) showing high expression of Rac1 and cdc42 in the ICs (white arrow). 63x is shown in C-C1 and D-D1. GFP channel is shown in A1, B1, C1 and D1. E-G1) Late pupal germaria of transgenic flies with GFP tagged to RhoA, Rac1 and cdc42, respectively stained for Tj (red), GFP (green) and Vasa (blue) showing low expression of RhoA, Rac1, and cdc42 in pupal escort cells (white arrow). GFP channel is shown in E1, F1 and G1. (H-I1) Adult germaria of transgenic flies with GFP tagged to Rac1 and cdc42, respectively stained for Tj (red), GFP (green) and Vasa (blue) showing low expression of Rac1 and cdc42 in the adult escort cells (white arrow). GFP channel is shown in H1 and I1. Scale bar for C1 and D1 is 10\(\mu\)m. Scale bar for all other images is 20\(\mu\)m.

(TIF)

S7 Fig. Escort cells express activated forms of cdc42 but not of RhoA/Rac. (A-B1) Adult germaria of transgenic flies that report active form of RhoA/Rac and cdc42. While RhoA/Rac is not expressed in the escort cells, it was expressed in the follicle cells (white arrow). cdc42 is active in the adult escort cells. Germline is marked by Vasa-KO. GFP channels are shown in A1 and B1. C) Quantification (n = 3) of GFP in the ICs and adult escort cells showing that while active RhoA/Rac is not expressed in the adult escort cells, active form of cdc42 is highly expressed in the ICs in comparison to the adult escort cells. (D-D1) Late pupal gerarium of fly carrying the Wnt canonical reporter, Frizzled3 (Fz3) stained for RFP (red), Tj (green) and 1B1 (blue) showing high expression of Fz3 in the pupal escort cells (white arrow). RFP channel is shown in D1. Scale bar for all images is 20\(\mu\)m.

(TIF)

S8 Fig. c587 is expressed in ICs of the larval gonad and escort cells of the adult germaria. (A-B1) Late larval gonad of flies with mCD8GFP under the control of c587 stained for Tj (red), GFP (green) and Vasa (blue) showing GFP expression in the ICs (white arrows). 63x is shown in B-B1. GFP channel is shown in A1 and B1. (C-C1) Adult germaria of flies with mCD8GFP under the control of c587 stained for Tj (red), GFP (green) and Vasa (blue) showing GFP expression in the escort cells (white arrow). GFP channel is shown in C1. Scale bar for B and B1 is 10\(\mu\)m. Scale bar for all other images is 20\(\mu\)m.

(TIF)

S9 Fig. cdc42 is required prior to eclosion to regulate germline stem cell differentiation. (A-B) Germaria of c587-GAL4 (control) and 18 °C to 29 °C temperature shift cdc42 depleted escort cells using cdc42\textsuperscript{DN} stained with 1B1 (red) and Vasa (blue) showing no differentiation defects in cdc42 mutants (yellow line). (C-D) Germaria of c587-GAL4 (control) and 29 °C to 18 °C temperature shift cdc42 depleted escort cells using cdc42\textsuperscript{DN} stained with 1B1 (red) and Vasa (blue) showing an accumulation of >3 undifferentiated cells in cdc42 mutants (yellow line). (E) Percentage of the germaria with >3 spectrosomes in 18 °C to 29 °C and 29 °C to 18 °C temperature shift flies, in c587-GAL4 and cdc42\textsuperscript{DN} mutants showing a significant difference between c587-GAL4 and cdc42\textsuperscript{DN} expressed escort cells in 29 °C to 18 °C but not in 18 °C to 29 °C (n = 50). (F) Quantification of the number of spectrosomes in 18 °C to 29 °C and 29 °C to 18 °C temperature shift flies in c587-GAL4, cdc42\textsuperscript{DN} expressed escort cells showing a significant
difference between c587-GAL4 and cdc42 mutants in 29 °C to 18 °C but not in 18 °C to 29 °C (n = 50). (G) Quantification of the number of spectrosomes in 18 °C to 29 °C temperature shift flies in c587-GAL4, dWnt4, RhoA, Rac1 and cdc42 depleted escort cells showing a significant difference between c587-GAL4 and dWnt4 RNAi mutants, but not between c587-GAL4 and RhoA, Rac1 and cdc42 mutants (n = 50). (H) Quantification of the number of spectrosomes in 29 °C to 18 °C temperature shift flies, in c587-GAL4, dWnt4, RhoA, Rac1 and cdc42 depleted escort cells showing a significant difference between c587-GAL4 and dWnt4, RhoA, Rac1 and cdc42 depleted escort cells (n = 50). Scale bar for all images is 20μm.

**S10 Fig. Downstream Wnt non-canonical components regulate intermingling of ICs.**

(A-C) Larval gonads of tj-GAL4 (control) and mutants expressing dominant negative Rho and cdc42 in the ICs stained with Tj (red), 1B1 (blue) and Vasa (green) showing loss of intermingling of ICs in the mutants. (D-H) Larval gonads of control, dsh, cdc42, dWnt4, Rac1, Mtl heterozygotes stained with Tj (red), 1B1 (blue) and Vasa (green) showing loss of intermingling of ICs in the mutants and Rac heterozygotes. Scale bar for all images is 20μm.

**S11 Fig. Downstream Wnt non-canonical components, RhoA, Rac1, Mtl and cdc42 regulate IC cell division and RhoA regulates cell survival.** (A-H) Larval gonads of control, dWnt4 heterozygous, dWnt4 mutant, tj-GAL4 (control), RhoA, Rac1, Mtl and cdc42 depleted ICs stained with Tj (red), PH3 (green) and 1B1 (blue) showing a significant difference in the division rate of the ICs between control and dWnt4 mutants and, tj-GAL4 and RhoA, Rac1, Mtl and cdc42 mutants (n = 10). (I-O) Larval gonads of dWnt4 heterozygous, dWnt4 mutants, tj-GAL4 (control), RhoA, Rac1, Mtl and cdc42 depleted ICs stained with Tj (red), Caspase3 (green) and 1B1 (blue) showing Caspase3 positive staining in gonads with RhoA depleted ICs (white arrows) (n = 5). Scale bar for all images is 20μm.

**S12 Fig. PCP regulators and Wnt canonical pathway component do not regulate the intermingling of ICs.** (A-F) Larval gonads of tj-GAL4 (control), vang, ds, fz, ft and arr depleted ICs stained with Tj (red), 1B1 (blue) and Vasa (green) showing no intermingling defects in vang, ds, fz, ft and arr depleted ICs (n = 25). Scale bar for all images is 20μm.

**S13 Fig. dWnt4 does not regulate the protein levels of RhoA, Rac1 and cdc42 in the ICs.** (A-F1) Larval gonad of dWnt4 heterozygous and dWnt4 mutants that carry GFP tagged to RhoA, Rac1 or cdc42 stained for Tj (red), GFP (green) and Vasa (blue) showing similar expression of RhoA, Rac1 or cdc42 in the ICs of heterozygotes and mutants. GFP channel is shown in A1, B1, C1, D1, E1 and F1. (G) Quantification (n = 3) of GFP in the ICs showing that the protein levels of RhoA, Rac1 and cdc42 are not altered between dWnt4 heterozygotes and dWnt4 mutants. (H-H1) Larval gonad of dWnt4 heterozygous lacking RhoA/Rac GTPase reporter, imaged under the same confocal settings (Fig 6A–6B1) showing no GFP. GFP channel is shown in H1. (I-I1) Larval gonad of dWnt4 heterozygous lacking cdc42 GTPase reporter, imaged under the same confocal settings (Fig 6C–6D1) showing no GFP. GFP channel is shown in I1. (J) Quantification (n = 3) of GFP in the terminal filament showing that cdc42 activity is not altered in dWnt4 mutants. Scale bar for all images is 20μm.
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References

1. Boyle M, Wong C, Rocha M, Jones DL. Decline in self-renewal factors contributes to aging of the stem cell niche in the Drosophila testis. Cell Stem Cell. 2007; 1: 470–478. https://doi.org/10.1016/j.stem.2007.08.002 PMID: 18371382

2. Pan L, Chen S, Weng C, Call G, Zhu D, Tang H, et al. Stem Cell Aging Is Controlled Both Intrinsically and Extrinsically in the Drosophila Ovary. Cell Stem Cell. 2007; 1: 458–469. https://doi.org/10.1016/j.stem.2007.09.010 PMID: 18371381

3. Hanahan D, Weinberg RA. Hallmarks of Cancer: The Next Generation. Cell. Elsevier; 2011; 144: 646–674. https://doi.org/10.1016/j.cell.2011.02.013 PMID: 21376230

4. Spradling AC, de Cuevas M, Drummond-Barbosa D, Keyes L, Lilly M, Pepling M, et al. The Drosophila germarium: stem cells, germ line cysts, and oocytes. Cold Spring Harb Symp Quant Biol. 1997; 62: 25–34. PMID: 9598333

5. Spradling A, Fuller MT, Braun RE, Yoshida S. Germline stem cells. Cold Spring Harb Perspect Biol. Cold Spring Harbor Laboratory; 2011; 3: a002642–a002642. https://doi.org/10.1101/cshperspect.a002642 PMID: 21791699

6. Spradling AC, Nystul T, Lighthouse D, Morris L, Fox D, Cox R, et al. Stem cells and their niches: integrated units that maintain Drosophila tissues. Cold Spring Harb Symp Quant Biol. Cold Spring Harbor Laboratory Press; 2008; 73: 49–57. https://doi.org/10.1101/sqb.2008.73.023 PMID: 19022764

7. Lehmann R. Germline stem cells: origin and destiny. Cell Stem Cell. Elsevier; 2012; 10: 729–739. https://doi.org/10.1016/j.stem.2012.05.016 PMID: 22704513

8. Morrison SJ, Spradling AC. Stem Cells and Niches: Mechanisms That Promote Stem Cell Maintenance throughout Life. Cell. 2008; 132: 596–611. https://doi.org/10.1016/j.cell.2008.01.036 PMID: 18295578

9. Li L, Xie T. STEM CELL NICHE: Structure and Function. Annual Review of Cell and Developmental Biology. Annual Reviews; 2005; 21: 605–631. https://doi.org/10.1146/annurev.cellbio.21.012704.131528 PMID: 16212509

10. Xie. Germline stem cell niches. StemBook. 2008. https://doi.org/10.3824/stembook.1.23.1 PMID: 20614617

11. Gilboa L, Lehmann R. Soma–germline interactions coordinate homeostasis and growth in the Drosophila gonad. Nature. Nature Publishing Group; 2006; 443: 97–100. https://doi.org/10.1038/nature05068 PMID: 16936717
12. Dansereau DA, Lasko P. The development of germline stem cells in Drosophila. Methods Mol Biol. Totowa, NJ: Humana Press; 2008; 450: 3–26. https://doi.org/10.1007/978-1-60327-214-8_1 PMID: 18370048

13. Gott D, Laski FA. Mechanisms of cell rearrangement and cell recruitment in Drosophila ovary morphogenesis and the requirement of bic & brac. Development. 1995; 121: 173–187. PMID: 7867498

14. Sahut-Barnola I, Gott D, Laski FA, Couderc J-L. Drosophila Ovary Morphogenesis: Analysis of Terminal Filament Formation and Identification of a Gene Required for This Process. Developmental Biology. 1995; 170: 127–135. https://doi.org/10.1006/dbio.1995.1201 PMID: 7601303

15. Zhu C-H, Xie T. Clonal expansion of ovarian germline stem cells during niche formation in Drosophila. Development. 2003; 130: 2579–2588. PMID: 12736203

16. Ehrman L. Ovarian Development in Drosophila melanogaster. Robert C. King. The Quarterly Review of Biology. Stony Brook Foundation, Inc; 1971; 46: 79–79. https://doi.org/10.1086/406775

17. Asaoka M, Lin H. Germline stem cells in the Drosophila ovary descend from pole cells in the anterior region of the embryonic gonad. Development. 2004; 131: 5079–5089. https://doi.org/10.1242/dev.01391 PMID: 15459101

18. Gilboa L, Lehmann R. Repression of primordial germ cell differentiation parallels germ line stem cell maintenance. Curr Biol. 2004; 14: 981–986. https://doi.org/10.1016/j.cub.2004.05.049 PMID: 15182671

19. Lin H, Spradling AC. Germline Stem Cell Division and Egg Chamber Development in Transplanted Drosophila Germaria. Developmental Biology. 1993; 159: 140–152. https://doi.org/10.1006/dbio.1993.1228 PMID: 8365558

20. Chen D, McKearin D. Dpp Signaling Silences bam Transcription Directly to Establish Asymmetric Divisions of Germline Stem Cells. Current Biology. 2003; 13: 1786–1791. https://doi.org/10.1016/j.cub.2003.09.033 PMID: 14561403

21. Ohlstein B, McKearin D. Ectopic expression of the Drosophila Bam protein eliminates oogenic germ-line stem cells. Development. 1997; 124: 3651–3662. PMID: 9342057

22. McKearin DM, Spradling AC. bag-of-marbles: a Drosophila gene required to initiate both male and female gametogenesis. Genes & Development. Cold Spring Harbor Lab; 1990; 4: 2242–2251. https://doi.org/10.1101/gad.4.12b.2242

23. McKearin D, Ohlstein B. A role for the Drosophila bag-of-marbles protein in the differentiation of cystoblasts from germline stem cells. Development. 1995; 121: 2937–2947. PMID: 7555720

24. Song X. Bmp signals from niche cells directly repress transcription of a differentiation-promoting gene, bag of marbles, in germline stem cells in the Drosophila ovary. Development. 2004; 131: 1353–1364. https://doi.org/10.1242/dev.01026 PMID: 14973291

25. Fuchs E, Tumbar T, Guasch G. Socializing with the neighbors: stem cells and their niche. Cell. 2004; 116: 769–778. PMID: 15035980

26. Xie T, Spradling AC. A niche maintaining germ line stem cells in the Drosophila ovary. Science. American Association for the Advancement of Science; 2000; 290: 328–330. https://doi.org/10.1126/science.290.5490.328 PMID: 11030649

27. Xie T, Song X, Jin Z, Pan L, Weng C, Chen S, et al. Interactions between stem cells and their niche in the Drosophila ovary. Cold Spring Harb Symp Quant Biol. 2008; 73: 39–47. https://doi.org/10.1101/sqb.2008.73.014 PMID: 19022749

28. Maimon I, Popliker M, Gilboa L. Without children is required for Stat-mediated zfh1 transcription and for germline stem cell differentiation. Development. Oxford University Press for The Company of Biologists Limited; 2014; 141: 2602–2610. https://doi.org/10.1242/dev.109611 PMID: 24903753

29. Li MA, Alls JD, Avancini RM, Koo K, Gott D. The large Maf factor Traffic Jam controls gonad morphogenesis in Drosophila. Nat Cell Biol. 2003; 5: 994–1000. https://doi.org/10.1038/ncb1058 PMID: 14578908

30. Ting X. Control of germline stem cell self-renewal and differentiation in the Drosophilaovary: concerted actions of niche signals and intrinsic factors. Wiley Interdiscip Rev Dev Biol. John Wiley & Sons, Inc; 2012; 2: 261–273. https://doi.org/10.1002/wdev.69 PMID: 24090936

31. Xie T, Spradling AC. decapentaplegic Is Essential for the Maintenance and Division of Germline Stem Cells in the Drosophila Ovary. Cell. 1998; 94: 251–260. https://doi.org/10.1016/S0092-8674(00)81424-5 PMID: 9695953

32. Song X, Call GB, Kirilly D, Xie T. Notch signaling controls germline stem cell niche formation in the Drosophila ovary. Development. The Company of Biologists Ltd; 2007; 134: 1071–1080. https://doi.org/10.1242/dev.003392 PMID: 17287246

33. Ward EJ, Shcherbata HR, Reynolds SH, Fischer KA, Hatfield SD, Ruohola-Baker H. Stem cells signal to the niche through the Notch pathway in the Drosophila ovary. Curr Biol. Elsevier; 2006; 16: 2352–2358. https://doi.org/10.1016/j.cub.2006.10.022 PMID: 17070683
34. King FJ, Lin H. Somatic signaling mediated by fs(1)Yb is essential for germline stem cell maintenance during Drosophila oogenesis. Development. 1999; 126: 1833–1844. PMID: 10101118

35. Cox DN, Chao A, Lin H. piwi encodes a nucleoplasmic factor whose activity modulates the number and division rate of germline stem cells. Development. 2000; 127: 503–514. PMID: 10631171

36. King FJ, Szakmary A, Cox DN, Lin H. Yb Modulates the Divisions of Both Germline and Somatic Stem Cells through piwi- and hh-Mediated Mechanisms in the Drosophila Ovary. Molecular Cell. 2001; 7: 497–508. https://doi.org/10.1016/S1097-2765(01)00197-6 PMID: 11463375

37. Song X. Germline Stem Cells Anchored by Adherens Junctions in the Drosophila Ovary Niches. Science. American Association for the Advancement of Science; 2002; 296: 1855–1857. https://doi.org/10.1126/science.1069871 PMID: 12052957

38. Song X, Xie T. DE-cadherin-mediated cell adhesion is essential for maintaining somatic stem cells in the Drosophila ovary. Proceedings of the National Academy of Sciences. National Acad Sciences; 2002; 99: 14813–14818. https://doi.org/10.1073/pnas.232389399 PMID: 12393817

39. Decotto E, Spradling AC. The Drosophila ovarian and testis stem cell niches: similar somatic stem cells and signals. Developmental Cell. Elsevier; 2005; 9: 501–510. https://doi.org/10.1016/j.devcel.2005.08.012 PMID: 16198292

40. Kirilly D, Wang S, Xie T. Self-maintained escort cells form a germline stem cell differentiation niche. Development. Oxford University Press for The Company of Biologists Limited; 2011; 138: 5087–5097. https://doi.org/10.1242/dev.067850 PMID: 22031542

41. Morris LX, Spradling AC. Long-term live imaging provides new insight into stem cell regulation and germline-soma coordination in the Drosophila ovary. Development. Oxford University Press for The Company of Biologists Limited; 2011; 138: 2207–2215. https://doi.org/10.1242/dev.065508 PMID: 21558370

42. König A, Scherbertha HR. Soma influences GSC progeny differentiation via the cell adhesion-mediated steroid-let-7-Wingless signaling cascade that regulates chromatin dynamics. Biology Open. The Company of Biologists Ltd; 2015; 4: 285–300. https://doi.org/10.1242/bio.201410553 PMID: 25661868

43. Morris LX, Spradling AC. Steroid signaling within Drosophila ovarian epithelial cells sex-specifically modulates early germ cell development and meiotic entry. Bökel C, editor. PLoS ONE. Public Library of Science; 2012; 7: e46109. https://doi.org/10.1371/journal.pone.0046109 PMID: 23056242

44. Wang S, Gao Y, Song X, Ma X, Zhu X, Mao Y, et al. Wnt signaling-mediated redox regulation maintains the germ line stem cell differentiation niche. eLife. eLife Sciences Publications Limited; 2015; 4: 408. https://doi.org/10.7554/eLife.08174 PMID: 26452202

45. Mottier-Pavie VI, Palacios V, Eliazer S, Scoggin S, Buszczak M. The Wnt pathway limits BMP signaling outside of the germline stem cell niche in Drosophila ovaries. Developmental Biology. 2016; 417: 50–62. https://doi.org/10.1016/j.ydbio.2016.06.038 PMID: 27364467

46. Upadhyay M, Martino Cortez Y, Wong-Deyrup S, Tavares L, Schwalter S, Flora P, et al. Transposon Dysregulation Modulates dWnt4 Signaling to Control Germline Stem Cell Differentiation in Drosophila. Molecular Cell. The Company of Biologists Limited; 2015; 4: 285–300. https://doi.org/10.1242/bio.201410553 PMID: 25661868

47. Komiya Y, Habas R. Wnt signal transduction pathways. Organogenesis. Landes Bioscience; 2008; 4: 68–75. PMID: 19279717

48. Bhanot P, Brink M, Samos CH, Hsieh J-C, Wang Y, Macke JP, et al. A new member of the frizzled family from Drosophila functions as a Wingless receptor. Nature. Nature Publishing Group; 1996; 382: 225–230. https://doi.org/10.1038/382225a0 PMID: 8717036

49. MacDonald BT, Tamai K, He X. Wnt/beta-catenin signaling: components, mechanisms, and diseases. Developmental Cell. Elsevier; 2009; 17: 9–26. https://doi.org/10.1016/j.devcel.2009.06.016 PMID: 19619488

50. Wehrli M, Dougan ST, Caldwell K, O’Keefe L, Schwartz S, Vaizel-Ohayon D, et al. arrow encodes an LDL-receptor-related protein essential for Wingless signalling. Nature. 2000; 407: 527–530. https://doi.org/10.1038/35035110 PMID: 11029006

51. Orsulic S, Peifer M. An in vivo structure-function study of armadillo, the beta-catenin homologue, reveals both separate and overlapping regions of the protein required for cell adhesion and for wingless signalling. The Journal of Cell Biology. The Rockefeller University Press; 1996; 134: 1283–1300. PMID: 8794868

52. van de Watering M, Cavallo R, Dooijes D, van Beest M, van Es J, Loureiro J, et al. Armadillo coactivates transcription driven by the product of the Drosophila segment polarity gene dTCF. Cell. 1997; 88: 789–799. PMID: 9118222

53. Peifer M, Orsulic S, Sweeten D, Wieschaus E. A role for the Drosophila segment polarity gene armadillo in cell adhesion and cytoskeletal integrity during oogenesis. Development. 1993; 118: 1191–1207. PMID: 8269848
54. Huber O, Korn R, McLaughlin J, Ohsugi M, Herrmann BG, Kemler R. Nuclear localization of beta-catenin by interaction with transcription factor LEF-1. Mech Dev. 1996; 59: 3–10. PMID: 8892228
55. Behrens J, Kries von JP, Kühler M, Bruhn L, Wedlich D, Grosschedl R, et al. Functional interaction of beta-catenin with the transcription factor LEF-1. Nature. Nature Publishing Group; 1996; 382: 638–642. https://doi.org/10.1038/382638a0 PMID: 8757136
56. Nusse R. Wnt signaling in disease and in development. Cell Res. Nature Publishing Group; 2005; 15: 28–32. https://doi.org/10.1038/sj.cr.7290260 PMID: 15686623
57. Olson ER, Pancratov R, Chatterjee SS, Pervaiz Z, DasGupta R, Yan, an ETS-domain transcription factor, negatively modulates the Wingless pathway in the Drosophila eye. EMBO Rep. EMBO Press; 2011; 12: 1047–1054. https://doi.org/10.1038/embor.2011.159 PMID: 21869817
58. Wu J, Roman A-C, Carvajal-Gonzalez JM, Mlodzik M. Wg and Wnt4 provide long-range directional input to planar cell polarity orientation in Drosophila. Nat Cell Biol. Nature Research; 2013; 15: 1045–1055. https://doi.org/10.1038/ncb2806 PMID: 23912125
59. Carvajal-Gonzalez JM, Mlodzik M. Mechanisms of planar cell polarity establishment in Drosophila. F1000Prime Rep. 2014; 6: 98. https://doi.org/10.12703/P6-98 PMID: 25580252
60. Singh J, Mlodzik M. Planar cell polarity signaling: coordination of cellular orientation across tissues. Wiley Interdiscip Rev Dev Biol. John Wiley & Sons, Inc; 2012; 1: 479–499. https://doi.org/10.1002/wdev.32 PMID: 23066429
61. Mlodzik M. Planar cell polarization: do the same mechanisms regulate Drosophila tissue polarity and vertebrate gastrulation? Trends Genet. 2002; 18: 564–571. PMID: 12414186
62. Strutt D, Warrington SJ. Planar polarity genes in the Drosophila wing regulate the localisation of the FH3-domain protein Multiple Wing Hairs to control the site of hair production. Development. The Company of Biologists Ltd; 2008; 135: 3103–3111. https://doi.org/10.1242/dev.025205 PMID: 18701542
63. Zallen JA. Planar polarity and tissue morphogenesis. Cell. 2007; 129: 1051–1063. https://doi.org/10.1016/j.cell.2007.05.050 PMID: 17574020
64. Simons M, Mlodzik M. Planar cell polarity signaling: from fly development to human disease. Annu Rev Genet. 2008; 42: 517–540. https://doi.org/10.1146/annurev.genet.42.110807.091432 PMID: 18710302
65. Donoughe S, DiNardo S. dachsous and frizzled contribute separately to planar polarity in the Drosophila ventral epidermis. Development. Oxford University Press for The Company of Biologists Limited; 2011; 138: 2751–2759. https://doi.org/10.1242/dev.063024 PMID: 21613320
66. Casal J, Lawrence PA, Struhl G. Two separate molecular systems, Dachso us/Fat and Starry night/Frizzled, act independently to confer planar cell polarity. Development. The Company of Biologists Ltd; 2006; 133: 4561–4572. https://doi.org/10.1242/dev.02641 PMID: 17075008
67. Devenport D. The cell biology of planar cell polarity. The Journal of Cell Biology. 2014; 207: 171–179. https://doi.org/10.1083/jcb.201408039 PMID: 25349257
68. Boutros M, Paricio N, Strutt DI, Mlodzik M. Dishevelled activates JNK and discriminates between JNK pathways in planar polarity and wingless signaling. Cell. 1998; 94: 109–118. PMID: 9674432
69. Eaton S, Wepf R, Simons K. Roles for Rac1 and Cdc42 in planar polarization and hair outgrowth in the wing of Drosophila. The Journal of Cell Biology. The Rockefeller University Press; 1996; 135: 1277–1289. PMID: 8947551
70. Winter CG, Wang B, Baliew A, Royou A, Karess R, Axelrod JD, et al. Drosophila Rho-associated kinase (Drok) links Frizzled-mediated planar cell polarity signaling to the actin cytoskeleton. Cell. 2001; 105: 81–91. PMID: 11301004
71. Hale R, Strutt D. Conservation of Planar Polarity Pathway Function Across the Animal Kingdom. Annu Rev Genet. Annual Reviews; 2015; 49: 529–551. https://doi.org/10.1146/annurev-genet-110807.091432 PMID: 26390326
72. Habas R, Dawid IB, He X. Coactivation of Rac and Rho by Wnt/Frizzled signaling is required for vertebrate gastrulation. Genes & Development. Cold Spring Harbor Lab; 2003; 17: 295–309. https://doi.org/10.1101/gad.102203 PMID: 12533515
73. Habas R, Kato Y, He X. Wnt/Frizzled activation of Rho regulates vertebrate gastrulation and requires a novel Formin homology protein Daam1. Cell. 2001; 107: 843–854. PMID: 11779461
74. Schlessinger K, Hall A, Tolwinski N. Wnt signaling pathways meet Rho GTPases. Genes & Development. Cold Spring Harbor Lab; 2009; 23: 265–277. https://doi.org/10.1101/gad.176080 PMID: 19204114
75. Veeman MT, Axelrod JD, Moon RT. A second canon. Functions and mechanisms of beta-catenin-independent Wnt signaling. Developmental Cell. 2003; 5: 367–377. PMID: 12967557
76. Ridley AJ. Rho GTPases and cell migration. Journal of Cell Science. 2001; 114: 2713–2722. PMID: 11683406
77. Jaffe AB, Hall A. Rho GTPases: biochemistry and biology. Annual Review of Cell and Developmental Biology. Annual Reviews; 2005; 21: 247–269. https://doi.org/10.1146/annurev.cellbio.21.020604.150721 PMID: 16212495
78. Nobes CD, Hall A. Rho GTPases control polarity, protrusion, and adhesion during cell movement. The Journal of Cell Biology. The Rockefeller University Press; 1999; 144: 1235–1244. PMID: 10087266
79. Svit T, Manser E. Rho GTPases and their role in organizing the actin cytoskeleton. Journal of Cell Science. 2011; 124: 679–683. https://doi.org/10.1242/jcs.064964 PMID: 21321325
80. Nobes CD, Hall A. Rac, and cdc42 GTPases regulate the assembly of multimolecular focal complexes associated with actin stress fibers, lamellipodia, and filopodia. Cell. 1995; 81: 53–62. PMID: 7536630
81. Cau J. Cdc42 controls the polarity of the actin and microtubule cytoskeletons through two distinct signal transduction pathways. Journal of Cell Science. The Company of Biologists Ltd; 2005; 118: 2579–2587. https://doi.org/10.1242/jcs.02385 PMID: 15928049
82. Cohen ED, Maril M-C, Wallace RMH, Weyers J, Kamberov YG, Pradel J, et al. DWnt4 regulates cell migration and focal adhesion kinase during Drosophila ovarian morphogenesis. Developmental Cell. 2002; 2: 437–448. https://doi.org/10.1016/S1534-5807(02)00142-9 PMID: 11970894
83. Ma X, Wang S, Do T, Song X, Inaba M, Nishimoto Y, et al. Piwi is required in multiple cell types to control germline stem cell lineage development in the Drosophila ovary. Bökel C, editor. PLoS ONE. Public Library of Science; 2014; 9: e90267. https://doi.org/10.1371/journal.pone.0090267 PMID: 24658126
84. Eliazer S, Palacios V, Wang Z, Kollipara RK, Kitler R, Buszczak M. Lsd1 restricts the number of germline stem cells by regulating multiple targets in escort cells. Perrimon N, editor. PLoS Genetics. Public Library of Science; 2014; 10: e1004200. https://doi.org/10.1371/journal.pgen.1004200 PMID: 24625679
85. de Cuevas M, Lee JK, Spradling AC. alpha-spectrin is required for germline cell division and differentiation in the Drosophila ovary. Development. 1996; 122: 3959–3968. PMID: 9012516
86. Lin H, Yue L, Spradling AC. The Drosophila fusome, a germline-specific organelle, contains membrane skeletal proteins and functions in cyst formation. Development. 1994; 120: 947–956. PMID: 7600970
87. Lasko PF, Ashburner M. Posterior localization of vasa protein correlates with, but is not sufficient for, pole cell development. Genes & Development. 1990; 4: 905–921. https://doi.org/10.1101/gad.4.6.905
88. Hogan J, Valentine M, Cox C, Doyle K, Collier S. Two frizzled planar cell polarity signals in the Drosophila wing are differentially organized by the Fat/Dachsous pathway. Rulifson E, editor. PLoS Genetics. Public Library of Science; 2011; 7: e1001305. https://doi.org/10.1371/journal.pgen.1001305 PMID: 21379328
89. Lin C, Kataeva VL. Kermit interacts with Gαo, Vang, and motor proteins in Drosophila planar cell polarity. Moreno E, editor. PLoS ONE. 2013; 8: e76885. https://doi.org/10.1371/journal.pone.0076885 PMID: 24204696
90. Ng J. Wnt/PCP proteins regulate stereotyped axon branch extension in Drosophila. Development. Oxford University Press for The Company of Biologists Limited; 2011; 139: 165–177. https://doi.org/10.1242/dev.068668 PMID: 22147954
91. Raftopoulou M, Hall A. Cell migration: Rho GTPases lead the way. Developmental Biology. 2004; 265: 23–32. https://doi.org/10.1016/j.ydbio.2003.06.003 PMID: 14697350
92. Fehon RG, Oren T, LaJeunesse DR, Melby TE, McCartney BM. Isolation of mutations in the Drosophila homologues of the human Neurofibromatosis 2 and yeast CDC42 genes using a simple and efficient reverse-genetic method. Genetics. Genetics Society of America; 1997; 146: 245–252. PMID: 9136014
93. Ng J, Nardine T, Harms M, Tzu J, Goldstein A, Sun Y, et al. Rac GTPases control axon growth, guidance and branching. Nature. Nature Publishing Group; 2002; 416: 442–447. https://doi.org/10.1038/416442a PMID: 11919635
94. Matussek T, Djiane A, Jankovics F, Brunner D, Mlodzik M, Mihały J. The Drosophila formin DAAM regulates the tracheal cuticle pattern through organizing the actin cytoskeleton. Development. The Company of Biologists Ltd; 2006; 133: 957–966. https://doi.org/10.1242/dev.02266 PMID: 16469972
95. Hakeda-Suzuki S, Ng J, Tzu J, Dietzl G, Sun Y, Harms M, et al. Rac function and regulation during Drosophila development. Nature. Nature Publishing Group; 2002; 416: 438–442. https://doi.org/10.1038/416438a PMID: 11919634
96. Strutt DI, Weber U, Mlodzik M. The role of RhoA in tissue polarity and Frizzled signalling. Nature. Nature Publishing Group; 1997; 387: 292–295. https://doi.org/10.1038/387292a0 PMID: 9153394

97. Habas R, Dawid IB. Dishevelled and Wnt signalling: is the nucleus the final frontier? J Biol. BioMed Central; 2005; 4: 2. https://doi.org/10.1186/jbiol22 PMID: 15720723

98. Axelrod JD, Miller JR, Shulman JM, Moon RT, Perrimon N. Differential recruitment of Dishevelled provides signaling specifity in the planar cell polarity and Wingless signaling pathways. Genes & Development. Cold Spring Harbor Laboratory Press; 1998; 12: 2610–2622.

99. Chen D. A discrete transcriptional silencer in the bam gene determines asymmetric division of the Drosophila germline stem cell. Development. The Company of Biologists Ltd; 2003; 130: 1159–1170. https://doi.org/10.1242/dev.00325 PMID: 12571107

100. Parisi MJ, Deng W, Wang Z, Lin H. The arrest gene is required for germline cyst formation during Drosophila oogenesis. Genesis. 2001; 29: 196–209. PMID: 11309853

101. Sugimura I, Lilly MA. Bruno inhibits the expression of mitotic cyclins during the prophase I meiotic arrest of Drosophila oocytes. Developmental Cell. 2006; 10: 127–135. https://doi.org/10.1016/j.devcel.2005.10.018 PMID: 16399084

102. Schulz C, Wood CG, Jones DL, Tazuke SI, Fuller MT. Signaling from germ cells mediated by the rhomboid homolog stet organizes encapsulation by somatic support cells. Development. 2002; 129: 4523–4534. PMID: 12223409

103. Sarov M, Barz C, Jambor H, Heim MY, Schmied C, Suchold D, et al. A genome-wide resource for the analysis of protein localisation in Drosophila. eLife. eLife Sciences Publications Limited; 2016; 5: e12068. https://doi.org/10.7554/eLife.12068 PMID: 26896675

104. Van Aelst L, D’Souza-Schorey C. Rho GTPases and signaling networks. Genes & Development. 1997; 11: 2295–2322.

105. Eliazer S, Shalaby NA, Buszczak M. Loss of lysine-specific demethylase 1 nonautonomously causes stem cell tumors in the Drosophila ovary. Proc Natl Acad Sci USA. National Acad Sciences; 2011; 108: 7064–7069. https://doi.org/10.1073/pnas.1015874108 PMID: 21482791

106. Aktary Z, Bertrand JU, Larue L. The WNT-less wonder: WNT-independent β-catenin signaling. Pigment Cell Melanoma Res. 2016; 29: 524–540. https://doi.org/10.1011/pcmr.12501 PMID: 27311806

107. Banisch TU, Maimon I, Dadosh T, Gilboa L. Escort cells generate a dynamic compartment for germline stem cell differentiation via combined Stat and Erk signalling. Development. 2017; 144: 1937–1947. https://doi.org/10.1242/dev.143727 PMID: 28659239
118. Raftopoulou M, Hall A. Cell migration: Rho GTPases lead the way. Developmental Biology. 2004; 265: 23–32. PMID: 14697350
119. Ridley AJ. Rho GTPase signalling in cell migration. Curr Opin Cell Biol. 2015; 36: 103–112. https://doi.org/10.1016/j.ceb.2015.08.005 PMID: 26363959
120. Friedl P, Wolf K. Tumour-cell invasion and migration: diversity and escape mechanisms. Nat Rev Cancer. 2003; 3: 362–374. https://doi.org/10.1038/nrc1075 PMID: 12724734
121. Eger A, Mikulits W. Models of epithelial–mesenchymal transition. Drug Discovery Today: Disease Models. 2005; 2: 57–63. https://doi.org/10.1016/j.ddmod.2005.04.001
122. Sahai E, Marshall CJ. RHO-GTPases and cancer. Nat Rev Cancer. 2002; 2: 133–142. https://doi.org/10.1038/nrc725 PMID: 12635176
123. Maimon I, Gilboa L. Dissection and staining of Drosophila larval ovaries. J Vis Exp. 2011; e2537–e2537. https://doi.org/10.3791/2537 PMID: 21610675