Piwi in the stem cell niche regulates nurse cell number and oocyte specification

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Abstract

Figure 1. Knockdown of piwi expression in terminal filament and cap cells, but not posterior escort cells, results in GSC proliferation and differentiation defects:
Description

In the *Drosophila* ovary, the proliferation of germline cells and differentiation of the oocyte is informed by cues from a variety of ovarian somatic cell (OSC) types. In particular, the stem cell niche, composed of cap cells, terminal filament cells, and escort cells, directly signals to germline stem cells (GSCs) to promote their self-renewal and differentiation into cystoblasts. Cap cells express Decapentaplegic (Dpp), which signals to the germline to promote proliferation and maintenance of GSCs (Xie and Spradling 1998, Chen and McKearin 2003, Song et al. 2004). Subsequently, the differentiation of cystoblasts and their four synchronous divisions with incomplete cytokinesis to form 16-cell cysts depends on the repression of Dpp by Wnt, EGF, and Hedgehog signaling from escort cells (Liu et al. 2010, Luo et al. 2015, Mottier-Pavie et al. 2016, Huang et al. 2017). The germline cyst then becomes encased in somatic follicle cells to form an egg chamber, in which one germinal cell is the oocyte and the other 15 are polyplid nurse cells. Follicle cells impact multiple aspects of germline differentiation throughout the rest of oogenesis, including polarization of the oocyte guided by JAK/STAT and Notch signaling (reviewed in Riechmann and Ephrussi 2001).

Early studies aimed at identifying genes involved in the balance of GSC maintenance and differentiation uncovered *piwi* (Cox et al. 1998, Cox et al. 2000). Clonal analysis showed that *piwi* expression in somatic, but not germline, cells is required for GSC maintenance (Cox et al. 1998). This led to the suggestion that Piwi mainly functions on GSCs through the stem cell niche. Mechanistic studies into Piwi and its associated small noncoding Piwi-interacting RNAs (piRNA), which guide Piwi to target sequences, have since focused on their role in transposon suppression. There is also a growing body of evidence that Piwi regulates some non-transposon gene expression programs during development (reviewed in Ozata et al. 2019).

Tissue-specific knockdown using the UASp-GAL4 system now provides an opportunity to dissect the function of Piwi in specific cell types of the ovary. Early studies that used this approach focused on the proliferation and differentiation of germline cells within the gerarium, and revealed that depletion of *piwi* in escort cells results in an over-proliferation of undifferentiated GSCs into “GSC tumors” (Jin et al. 2013, Ma et al. 2014). In our study focusing on mid-stage egg chambers, after depleting *piwi* in all OSCs using traffic jam-GAL4 (“*piwi-sKD*”), we observed defects in nurse cell number, oocyte number, and oocyte specification (Gonzalez et al. 2020). Because the aberrant egg chambers often contained improper numbers of nurse cells (but rarely in multiples of 15) and could contain from 0 to 3 “oocytes” (determined by morphology and localization of the oocyte marker protein Orb), we concluded that this phenotype was more likely due to improper germline proliferation and differentiation than improper encapsulation of the germline by follicle cells. Here, we further investigate whether these defects are due to loss of Piwi activity in terminal filament and cap cells (using *bab1-GAL4*, Figure 1A–A”), and/or in escort cells (using 13C06-GAL4, Figure 1G–G”).

Depletion of *piwi* in terminal filament and cap cells via *bab1-GAL4* (herein referred to as *piwi–bab1KD*, Figure 1B–1F”) resulted in egg chambers with defects in nurse cell number and oocyte specification, partially phenocopying *piwi-sKD* driven by *tj-GAL4*, which depletes *piwi* in all OSCs. 96.3% of the GFP-*bab1KD* egg chambers contained the expected 15 nurse cells, while only 82.8% and 29.8% of egg chambers in *piwi–bab1KD* #1 and *piwi–bab1KD* #2, respectively, contained exactly 15 nurse cells (Figure 1M). However, unlike depletion of *piwi* in all somatic cells, very few egg
chambers contained either tumorous (Figure 1M) or degenerate (Figure 1N) germline cells. Thus, piwi depletion in terminal filament and cap cells explains some, but not all, of the nurse cell number defects caused by piwi-sKD.

We also observed defects in oocyte specification upon piwi-bab1KD (Figure 1B-1F”). Only 84.1% and 30.9% of piwi–bab1KD #1 and piwi–bab1KD #2 egg chambers, respectively, contained egg chambers with a single oocyte at the posterior of the egg chamber and displaying Orb accumulation (Figure 1O). Within egg chambers that contained one presumptive oocyte (96.6% and 65.8% of egg chambers in piwi–bab1KD #1 and piwi–bab1KD #2, respectively), the most common abnormality in oocyte phenotype was a dispersal of Orb throughout the egg chamber (Figure 1O), indicating that oocyte specification is impaired upon piwi–bab1KD.

Notably, piwi-bab1KD #1 had a milder effect on both nurse cell number and oocyte specification than piwi–bab1KD #2. We had previously shown that piwi-RNAi #2 more strongly depletes piwi mRNA levels compared to piwi-RNAi #1 when driven in all somatic cells (Gonzalez et al. 2020), so this may reflect that a threshold level of piwi is necessary in the stem cell niche for this function.

We then investigated the effect of depleting piwi in posterior escort cells, follicle stem cells, and pre-follicle cells using 13C06-GAL4 (Figure 1G-1L”). Previous studies had shown a major role for Piwi in escort cells in regulating cystocyte differentiation (Jin et al. 2013, Ma et al. 2014), so we were surprised to observe relatively few egg chambers with defects in nurse cell number and/or oocyte specification following piwi-13C06KD (Figure 1M-O). Our result indicates that Piwi activity in posterior escort cells and follicle stem cells may not regulate nurse cell number and oocyte specification, and suggests that the GSC tumors described by previous studies may not be linked to the mid-stage germline defects we have described. However, we depleted piwi using 13C06-GAL4, which is expressed weakly in anterior escort cells but strongly in posterior escort cells (Sahai-Hernandez and Nystul 2013), while previous studies used c587-GAL4, which expresses more strongly in anterior escort cells (Song et al. 2004), so it is possible that Piwi’s escort cell function is primarily in anterior escort cells.

Altogether, these results indicate that Piwi functions within the stem cell niche, and perhaps mostly within terminal filament and cap cells, to regulate germline proliferation and differentiation within the gerarium, and that this has long-lasting developmental effects on egg chambers throughout oogenesis. The proliferation of germline cells and specification of the oocyte are temporally and spatially separate from the GSC-niche interaction, so the observation that gene expression in terminal filament and cap cells can influence these processes is striking. It suggests that the niche not only regulates the self-renewal of GSCs and their immediate differentiation into cystoblasts, but also modulates their developmental potential for processes that occur later in oogenesis. Alternatively, because bab1-GAL4 is also active in somatic cells of the larval gonad (Cho et al. 2018), our results may reflect a requirement for somatic piwi during gonadogenesis for the subsequent organization, proliferation, and specification of germline cells in the adult ovary.

Piwi regulates gene expression at both the transcriptional level (Brower-Toland et al. 2007, Wang & Elgin 2011, Sienski et al. 2012) and the post-transcriptional level (Robine et al. 2009, Saito et al. 2009, Klein et al. 2016), so further investigations should seek to understand which gene expression programs Piwi directly regulates to influence these oogenic processes. The observation that piwi depletion in somatic cells impacts the proliferation and specification of germline cells suggests that Piwi regulates the expression of genes involved in soma-to-germline signaling. Piwi has previously been shown to mediate Dpp signaling to regulate GSC maintenance (Jin et al. 2013, Ma et al. 2014), and the piRNA biogenesis factor Yb has been suggested to interact with the Notch pathway to regulate very similar nurse cell and oocyte phenotypes to those we have described upon piwi-sKD (Johnson et al. 1995). Future studies should investigate whether Piwi in the stem cell niche regulates the expression of genes in the Dpp, Notch, or other signaling pathways. These new findings would add to the growing evidence that the biological function of the PIWI/piRNA pathway extends well beyond the suppression of transposons.

Methods
Request a detailed protocol

Drosophila husbandry and genetics

All Drosophila stocks were raised on standard agar/molasses medium and raised at 25°C for experiments. Bab1-GAL4 (BDSC stock #6802) was used to express UASp constructs in all cap cells and terminal filament cells throughout development, and 13C06-GAL4 (BDSC stock #47860) was used to express UASp constructs in posterior escort cells, follicle stem cells, and prefollicle cells. UASp-GFP (BDSC #1521) was used to verify the expression pattern of GAL4 lines. Two anti-Piwi RNAi lines (“piwi RNAi #1” and “piwi-RNAi #2” in this study) were used to knock down piwi expression. Piwi-siRNA #1 targets exon 2 of the piwi mRNA and is BDSC stock #37483; piwi siRNA #2 targets exon 3 of the piwi mRNA and was a gift from T. Xie, Stowers Institute For Medical Research, Kansas City, MO. GFP-RNAi (BDSC stock #41550) was used as a negative control. To generate GAL4/UASp flies for analysis, two males carrying the GAL4 driver were crossed with three virgin females carrying the UASp construct. GAL4/UASp females were aged for two to three days in a ratio of 2:1 with w1118 males prior to ovary dissection.
Immunostaining

Ovaries from 2-3 day old females were dissected in 1X PBS and fixed with 200 μL of the following fixing solution (v/v%): PBS (89.5%), 10% Nonidet P-40 (5%), 37% formaldehyde (5.5%). The fixed ovaries were then washed 3 times for 15 minutes each in PBST (PBS and 0.2% Triton X-100). Ovaries were then blocked overnight in 5% NGS at 4°C, followed by incubation overnight at 4°C in primary antibody diluted in PBST + 2% NGS. Samples were washed three times in PBST and incubated in secondary antibodies overnight at 4°C. Samples were washed three times with PBST, stained with DAPI (1:500) and Phalloidin (1:200, ThermoFisher, #R415) for 15 minutes and mounted in Vectashield mounting media (Vector Labs, #H1000).

We used mouse anti-Orb 4H8 (1:300, DSHB) to visualize Orb localization. The following conjugated secondary antibodies were used, all at 1:500 dilutions: The Alexa 488-conjugated goat anti-mouse antibody, the Alexa 555-conjugated goat anti-mouse. Orb, Phalloidin, and DAPI were used qualitatively and quantitatively characterize the observed phenotypes of the egg chambers from all crosses.

Microscopy and phenotypic characterization

To observe the GAL4 expression pattern, confocal images of DAPI stained bab1-GAL4>UASp-GFP and 13C06-GAL4>UASp-GFP ovaries were taken using Leica TCS SP5 Confocal Laser Scanning Microscope. Piwi-knockdown samples were analyzed on the ZEISS Axio Imager2 for quantitative and qualitative characterizations.

DAPI and Phalloidin staining was used to identify nurse cells and oocytes. To more accurately quantify the effect of piwi–bab1KD and piwi-13C06KD on the number of nurse cells, we created four categories to describe how many nurse cells were in an egg chamber: 15 normal nurse cells; 0-14 normal nurse cells; 16-29 normal nurse cells; and ≥30 normal nurse cells, which we referred to as a tumorous number. Degenerate nurse cells were identified by the presence of bright pyknotic nuclei. Any large area of cytoplasm without a polyploid nucleus was considered a presumptive oocyte. We quantified how many egg chambers had zero, one, two, or three presumptive oocytes. We also characterized the localization of the Orb staining based on its relative position with the presumptive oocyte. We created four categories of Orb’s localization pattern: absent, oocyte-localized, mislocalized, and dispersed. If the egg chamber had no trace of Orb staining, the pattern was considered absent. If the Orb staining localized with the presumptive oocyte(s) it was designated oocyte-localized; but if the Orb staining was not localized to the presumptive oocyte and formed discrete patches, it was considered mislocalized. If the Orb staining filled the entire egg chamber, its pattern was considered dispersed.

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