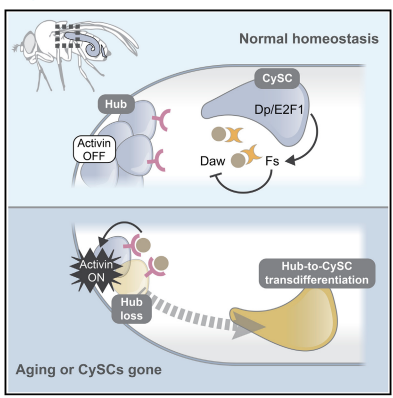
# Proliferative stem cells maintain quiescence of their niche by secreting the Activin inhibitor Follistatin

#### **Graphical abstract**



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#### In brief

How stem cells support niches remains largely unknown. Herrera et al. show that testicular stem cells maintain niche quiescence by secreting Follistatin, an Activin antagonist. Loss of Follistatin in stem cells or gain of Activin signaling in niche cells causes niche-to-stem-cell transdifferentiation, and this process underlies niche decay during aging.

#### **Highlights**

- Dp/E2f1 is required in stem cells to non-autonomously maintain niche quiescence
- Dp/E2f1 promotes niche quiescence through Fs, an Activin antagonist
- Activin signaling in niche cells causes transdifferentiation into functional stem cells
- Fs in stem cells regulates the physiological decay of the niche with age







#### **Article**

# Proliferative stem cells maintain quiescence of their niche by secreting the Activin inhibitor Follistatin

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#### **SUMMARY**

Aging causes stem cell dysfunction as a result of extrinsic and intrinsic changes. Decreased function of the stem cell niche is an important contributor to this dysfunction. We use the *Drosophila* testis to investigate what factors maintain niche cells. The testis niche comprises quiescent "hub" cells and supports two mitotic stem cell pools: germline stem cells and somatic cyst stem cells (CySCs). We identify the cell-cycle-responsive Dp/E2f1 transcription factor as a crucial non-autonomous regulator required in CySCs to maintain hub cell quiescence. Dp/E2f1 inhibits local Activin ligands through production of the Activin antagonist Follistatin (Fs). Inactivation of Dp/E2f1 or Fs in CySCs or promoting Activin receptor signaling in hub cells causes transdifferentiation of hub cells into fully functional CySCs. This Activin-dependent communication between CySCs and hub regulates the physiological decay of the niche with age and demonstrates that hub cell quiescence results from signals from surrounding stem cells.

#### **INTRODUCTION**

The niche creates a distinct microenvironment for stem cells and secretes short range self-renewal cues that promote "stemness" in the resident population (Morrison and Spradling, 2008). Decreased stem cell function with age can at least in part be attributed to both decreased niche function and niche cell numbers (Oh et al., 2014). These observations raise the question of what factors maintain niche cells. The Drosophila testis provides an ideal system to address this important issue. The testis niche comprises approximately twelve quiescent somatic hub cells that support two mitotic stem cell pools: germline stem cells (GSCs) that ultimately produce sperm and somatic cyst stem cells (CySCs) that support GSCs and produce somatic support cells. The hub and CySCs share a common lineage during development: somatic cells are initially all equivalent somatic gonadal precursors, but during embryonic stages, a subset of these are specified to become hub cells (Okegbe and DiNardo, 2011; Dinardo et al., 2011; Le Bras and Van Doren, 2006; Anllo et al., 2019; Kitadate and Kobayashi, 2010). The remaining somatic precursors become CySCs and their offspring, cyst cells. Intriguingly, despite their common origin, CySCs are the only somatic cells in the testis to proliferate, and both hub and cyst cells

are post-mitotic. Since they cease proliferating, hub cells need to be maintained during adulthood; previous work has identified several factors that act autonomously within the hub to maintain quiescence or survival (Hétié et al., 2014; Voog et al., 2014; Resende et al., 2013; Greenspan and Matunis, 2018). These include the transcription factor Escargot (Esg), as well as the cell-cycleinhibitor retinoblastoma homolog, Rbf. Most intriguingly, prior work has shown that genetic ablation of all CySCs causes hub cells to exit quiescence and transdifferentiate into CySCs, suggesting the existence of unidentified CySC-derived factors that non-autonomously maintain hub cells (Hétié et al., 2014). Finally, during normal aging, the ability of the hub to support stem cells declines. This is due both to a reduction in hub cell numbers (Sreejith et al., 2019; Wallenfang et al., 2006; Lee et al., 2016), as well as lower production of self-renewal ligands to support surrounding stem cells (Toledano et al., 2012; Boyle et al., 2007). Yet how the mechanisms maintaining hub cell quiescence are affected during aging is still unknown.

Here, we identify how CySCs maintain hub cell quiescence. We find that hub cells are lost following depletion of the transcription factor that regulates S-phase gene expression, which is a complex of the activator E2f, called E2f1 in *Drosophila*, and the sole dimerization partner homolog, Dp, in CySCs. This





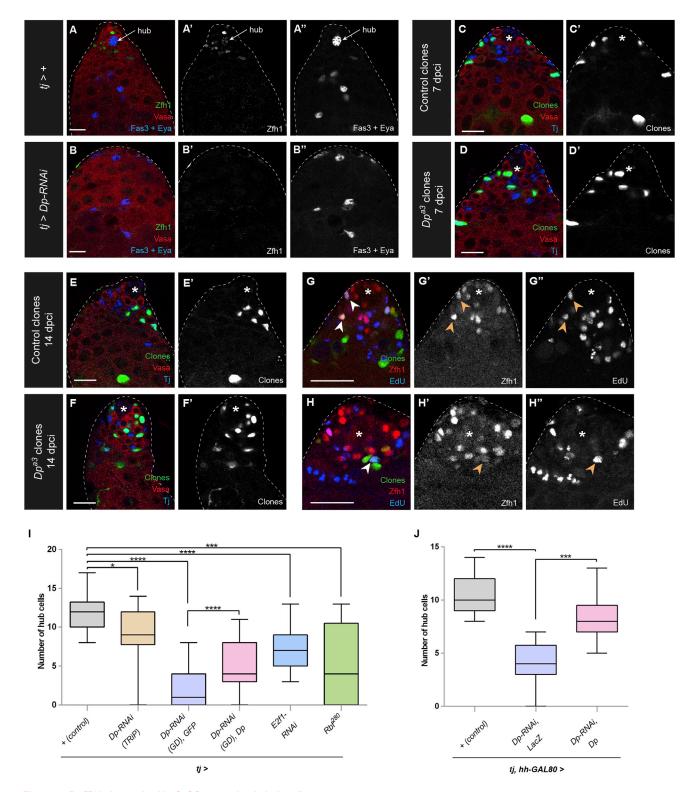


Figure 1. Dp/E2f1 is required in CySCs to maintain hub cells

(A and B) A control *tj-GAL4* (labeled *tj>+*) adult testis with hub cells (A, arrow) surrounded by both GSCs and CySCs. A *tj>Dp-RNAi* (B) adult testis lacking CySCs, GSCs, and hub cells. Both testes were isolated after 10 days at 29°C to induce maximal GAL4 activity. Zfh1 (green) labels CySCs, Vasa (red) marks the germline, Fas3 (blue) marks the hub cell membranes, and Eya (blue) labels the nucleus of differentiating cyst cells.

(C–H) GFP-positive FRT<sup>42D</sup> control clones (C, E, and G) or FRT<sup>42D</sup> Dp<sup>a3</sup> mutant clones (D, F, and H). Both types of CySC clones can be recovered at 7 days post clone induction (dpci) (C and D) and 14 dpci (E and F) and both incorporate EdU (blue, G and H), indicating that they can undergo S phase. Clones are marked by GFP (green), Vasa (red, C–F) marks the germline, and Tj (blue, C–F) marks CySCs and early cyst cells. Zfh1 (blue, G and H) marks CySCs.



signaling between cycling CySCs and the hub is mediated through production of Follistatin (Fs), an antagonist of Activin signaling. Finally, we show that Activin autonomously promotes hub cells to transdifferentiate into CySCs, leading to hub cell loss, and that increased activity of this pathway is responsible for age-dependent loss of hub cells.

#### **RESULTS**

### Dp/E2f1 functions in CySCs to non-autonomously maintain hub cells

Our prior work focused on how cell-cycle progression in CySCs, the only mitotic somatic cells in the testis, influenced cell fate (Amoyel et al., 2014). Here, we focused on the transcription factor that regulates S-phase gene expression, which is a complex of E2f1 and Dp, and which we previously reported to be active in CySCs (Amoyel et al., 2014). Knockdown of Dp by RNAi using the cyst lineage driver traffic jam (tj)-GAL4 resulted in a complete loss of CySCs, as identified by Zfh1 expression, and only Eya-positive differentiated cyst cells were visible (Leatherman and Dinardo, 2008; Fabrizio et al., 2003) (Figures 1A and 1B). In most cases (12/ 20), testes with somatic Dp depletion lacked the entire stem and early-differentiated cell compartment for both somatic and germ lineages and contained only spermatocytes or spermatid fibers. This suggested that Dp was required for CySC self-renewal and that its loss resulted in ectopic and premature differentiation. To confirm this, we used mitotic recombination to generate clones mutant for Dp. Surprisingly, Dp-mutant CySC clones, which we identified as Tj-positive cells adjacent to the hub, were recovered at similar rates to control clones, both 7 days post clone induction (dpci) and 14 dpci (Figures 1C-1F, S1A, and S1B), indicating that they had no autonomous self-renewal defect. We confirmed this result using an independent Dp null allele and used an antibody against Dp to verify that the mutant clones lacked Dp protein (Figures S1A-S1C). Similarly, CySC clones expressing the same Dp-RNAi as in Figure 1B were also recovered at 7 dpci (Figure S1D). Dp null mutant clones resulting from single-clonal-induction events contained many cells, indicating that they had proliferated over the course of the experiment (Figure 1D). Indeed, Dp-mutant CySCs were found to incorporate the nucleotide analog 5-ethynyl-2'-deoxyuridine (EdU), demonstrating that cells lacking Dp could undergo DNA replication (Figures 1G, 1H, S1A, and S1E, arrowheads). Consistently, we recovered clones mutant for E2f1, the sole activator E2f in Drosophila, at similar rates to control clones at 7 dpci (Figures S1F and S1G). Although surprising, this result concurs with recent work showing that Dp is dispensable for most larval proliferation in Drosophila and that viable adults lacking Dp in all but muscle tissues can be obtained (Zappia and Frolov, 2016). Similarly in mouse, proliferation still occurs in the absence of all activating E2fs (Chen et al., 2009).

The fact that Dp was not required autonomously for CySC selfrenewal, but its depletion in all CySCs led to loss of the entire stem cell population suggested that Dp may be required in a non-autonomous manner to maintain CySCs. Indeed, we noticed that in addition to loss of CySCs, hub cells were also absent in testes in which Dp was knocked down with tj-GAL4 (Figures 1A, 1B, and 1I). Whereas in controls the hub is composed of  $11.8 \pm 0.5$  cells, when Dp was knocked down in the cyst lineage, there were many fewer cells (2.1  $\pm$  0.4). To verify this observation, we used the markers upd-LacZ and hh-LacZ to label hub cells and found a reduction of labeled cells in the Dp knockdown and, in many cases, no cells expressing these markers (Tulina and Matunis, 2001; Forbes et al., 1996; Amoyel et al., 2013) (Figures S2A-S2D). To confirm that this phenotype was due to loss of Dp and not an off-target effect of the RNAi, we used three approaches: first, we expressed full-length Dp together with Dp-RNAi and observed a partial rescue of hub cell numbers compared with co-expressing GFP as a titration control (Figure 1I, compare purple with pink bar); second, we used an independent RNAi line targeting Dp, which gave a similar, albeit weaker, loss of hub cells (Figure 1I, compare brown with gray bar); third, we inactivated the Dp/E2f1 transcription factor by knocking down E2f1 or overexpressing a constitutively active form of the Dp/E2f1 inhibitor Rbf, called Rbf<sup>280</sup>, which recapitulated the loss of hub cells observed in Dp knockdowns in both cases (Figure 1I, compare blue and green bars with gray bar).

We then ruled out the possibility of hub loss being due to a developmental defect of the Dp-RNAi by using a temperaturesensitive form of GAL80 (GAL80<sup>TS</sup>) to only induce transgene expression in adult stages (McGuire et al., 2004). We analyzed flies at eclosion (0 days) after raising them at the permissive temperature for GAL80<sup>TS</sup> (18°C) and observed no significant differences in hub cell numbers between control flies expressing tj-GAL4 and GAL80<sup>TS</sup>, referred to as tj<sup>TS</sup>, and flies also carrying Dp or E2f RNAi transgenes (Figure S2E). After shifting to the restrictive temperature, we observed no significant change in hub cells in control  $tj^{TS}$  flies over 10 days, although we see a slight but not significant decrease in 20-day-old flies. By contrast, Dp knockdown, E2f1 knockdown, or Rbf<sup>280</sup> overexpression in CySCs led to a progressive loss of hub cells and an almost complete loss (0.03 ± 0.03 in the Dp knock down) by 20 days of adulthood (Figure S2E and Table S1). Altogether, these results indicate that the hub cell loss observed in Dp knockdowns is due to a progressive defect in maintenance of hub cells, not in their establishment.

We sought to establish that Dp/E2f1 function was indeed required, specifically in CySCs, to maintain hub cells non-autonomously. Since *tj-GAL4* is occasionally expressed in hub cells (Fairchild et al., 2016), we generated a strain expressing *GAL80* under the control of the endogenous *hh* locus, which is

(I) Graph showing the average number of hub cells after 10 days in  $29^{\circ}$ C using tj-GAL4 in control (+, gray bar, n = 18), Dp-RNAi (brown and purple bars, n = 14 and n = 31, respectively), Dp depletion plus exogenous Dp (pink bar, n = 15), E2f1 depletion (blue bar, n = 17), and overexpression of Rbf<sup>280</sup> (green bar, n = 12). (I and J) Graphs showing the average number of hub cells after 10 days in  $29^{\circ}$ C using tj-GAL4 and hh-GAL80, which inhibits GAL4 activity in the hub, limiting expression of UAS-dependent constructs to CySCs in control (+, gray bar, n = 27), Dp-RNAi (purple bar, n = 8), and Dp depletion plus exogenous Dp (pink bar, n = 33).

An asterisk marks the hub.

Error bars represent the data range. \*\*\*\*p < 0.0001; \*\*\*p < 0.001; \*p < 0.05 as assessed by Student's t test.

See also Tables S1 and S2; Figures S1 and S2. Scale bar, 20  $\mu\text{M}.$ 

#### **Article**



expressed exclusively in hub cells (see STAR Methods) (Amoyel et al., 2013; Michel et al., 2012). Importantly, knockdown of Dp using tj-GAL4 led to significant loss of hub cells in a hh-GAL80 background (4.0  $\pm$  0.8 hub cells in *Dp*-RNAi versus 10.6  $\pm$  0.3 in control, Figure 1J, purple bar). As expected, hub cell loss was suppressed by co-expressing Dp in non-hub somatic cells  $(8.4 \pm 0.3 \text{ hub cells versus } 4.0 \pm 0.8 \text{ in controls expressing})$ LacZ as a titration control, n = 33 and 8, respectively, p < 0.001, Figure 1J, pink bar). Finally, to rule out any possibility of ectopic hub expression of the RNAi, we used alternative GAL4 lines, which drive expression in CySCs and cyst cells and which were previously shown to have sporadic or negligible expression in hub cells: eyaA3-GAL4 (Fairchild et al., 2016), fringe (fng)-GAL4 (Dinardo et al., 2011), and C587-GAL4 (Fairchild et al., 2016; Hétié et al., 2014). Knockdown of Dp using all three drivers resulted in a loss of hub cells (Figure S2F). In the case of fng-GAL4, we used GAL80<sup>TS</sup> to show progressive hub cell loss in adult flies when Dp or E2f1 were knocked down (Figure S2F and Table S1). In sum, our results indicate that activity of the Dp/E2f1 transcription factor is dispensable within individual CySCs for cell-cycle progression but that its loss in all adult CySCs results in a progressive and non-autonomous loss of hub cells. These observations suggest a model in which Dp/ E2f1 activity in CySCs causes them to signal to hub cells to promote their maintenance in adult testes.

### **Dp/E2f1** regulates *follistatin* expression to maintain the hub non-autonomously

To determine the signals downstream of Dp/E2f1 that acted on the hub non-autonomously, we performed an RNAi screen. We used tj-GAL4 to deplete from CySCs secreted factors that are enriched in testicular stem cells and then assessed hub cell number (Terry et al., 2006; Kurusu et al., 2008) (Figure S3A). Knockdown of Fs, encoding a conserved repressor of TGFβ/Activin ligands (Figure 2A) (Pentek et al., 2009), recapitulated the phenotypes observed with Dp or E2f1 knockdown. After 14 days of RNAi expression, tj>Fs-RNAi testes contained 1.1 ± 0.3 hub cells, whereas control tj>LacZ testes had 9.7  $\pm$  0.5 hub cells (Figures 2B-2D). Frequently, tj>Fs-RNAi testes contained 0 hub cells. In these testes, both stem cell populations were lost, and we frequently observed only differentiated spermatids, indicating that all the more undifferentiated cell types had been lost to ectopic differentiation (Figures S3B and S3C). The specificity of the Fs-RNAi transgene was confirmed by qRT-PCR analysis showing a significant decrease in Fs transcripts in tj>Fs-RNAi testes (Figures S3D and S3E) and by a significant rescue of hub cells when full-length Fs was concomitantly overexpressed with Fs-RNAi (Figure 2D). Overexpression of wildtype Fs alone using tj-GAL4 did not affect the number of hub cells (Figure 2D). No phenotype was observed when Fs was depleted from the germline using nanos (nos)-GAL4 (Figure S3F).

To confirm these results, we generated a  $Fs^{null}$  allele lacking the sequence spanning coding exons 1 and 4 (Figure S3D, see STAR Methods). The deletion was confirmed by sequencing and by RT-qPCR (Figure S3E).  $Fs^{null}$  flies were adult viable and showed progressive hub cell loss with age: from 9.4  $\pm$  0.9 hub cells in 0-day-old adult flies to 2.2  $\pm$  0.6 hub cells in 28-day-old flies (Figures 2E and S3G). Hub cell loss in the  $Fs^{null}$  mutant was significantly rescued by overexpressing wild-type Fs in either somatic cells

(fng-GAL4) or hub cells (upd<sup>TS</sup>-GAL4) (Figures S3H and S3I). Furthermore, we observed similar hub cell loss in testes from trans-heterozygous combinations of other Fs alleles (Figure S3J). As expected, 28-day-old Fs<sup>null</sup> males were significantly less fertile than age-matched control flies (Figure 2F). Using a Fs-GAL4 transcriptional reporter, we found that Fs was expressed in hub cells, CySCs and early somatic cells (Figure 2G).

Given the remarkable similarity in the non-autonomous hub cell phenotypes observed upon Dp/E2f1 or Fs loss in CySCs, we asked whether Fs functioned downstream of the Dp/E2f1 complex in CySCs. To test this, we examined expression of the Fs-GAL4 transcriptional reporter in control testes or in testes somatically depleted for Dp. Fs transcription in the somatic lineage, as assessed by UAS-GFP expression, was significantly reduced when Dp was depleted compared with controls (Figures 2G-2I). We observed a similar reduction in the levels of an Fs-GFP protein trap upon E2f1 knockdown (Figures S4A-S4C). We reasoned that if Dp/E2f1 activity in CySCs non-autonomously maintained the hub by inducing Fs expression, then hub cell loss caused by Dp depletion should be prevented when Fs was concomitantly supplied. Indeed, exogenous Fs completely rescued hub cell loss compared with Dp-RNAi alone (Figure 2J; Table S1). Thus, Fs mediates the non-autonomous effects of Dp/E2f1 on hub cell maintenance.

### Activin signaling triggers hub-to-CySC transdifferentiation

Since Fs is an extracellular antagonist of Activin ligands, we hypothesized that sustained autonomous activation of the Activin pathway within hub cells would have a similar effect on hub cell maintenance. In Drosophila, three Activin ligands (Dawdle [Daw], Activinß [Actß], and Myoglianin [Myo]) stimulate the type I Activin receptor Baboon (Babo) (Upadhyay et al., 2017). This results in the activation of transcription factor Smox, the SMAD3 homolog, which then alters target gene transcription. Fs binds to Activin ligands and prevents them from binding to and activating Babo (Figure 2A).

First, we determined that hub cells can indeed respond to Activin ligands, by examining the distribution of the receptor. A protein trap for Babo was present at the surface of many cells in the testis, and importantly, was found decorating the membrane of Fas3positive hub cells (Figure 3A). Next, we tested what effect Activin pathway signaling had within hub cells. We used upd-GAL4 and GAL80<sup>TS</sup> (termed upd<sup>TS</sup>) to overexpress a constitutively active form of Babo (Babo QD) (Brummel et al., 1999) in hub cells and assessed hub cell number at 0, 7, 14, 21, and 28 days of adulthood. Freshly eclosed flies that were raised at the permissive temperature for GAL80<sup>TS</sup> had no significant differences in hub cell numbers, whether or not they carried the  $\textit{UAS-babo}^{\textit{QD}}$  transgene (Figure 3B). After shifting to the restrictive temperature, autonomous activation of the Activin pathway (upd<sup>TS</sup>>babo<sup>QD</sup>) in hub cells induced their progressive and complete loss by 28 days, whereas controls showed a modest decrease in hub cell number over this period (Figures 3B and 3C). We then asked what the fate of lost hub cells with ectopic Activin signaling could be and hypothesized that, since hub cells have the potential, under certain experimental conditions, to transdifferentiate into CySCs (Hétié et al., 2014; Voog et al., 2014; Greenspan and Matunis, 2018), Activin signaling may induce this identity switch. To test this



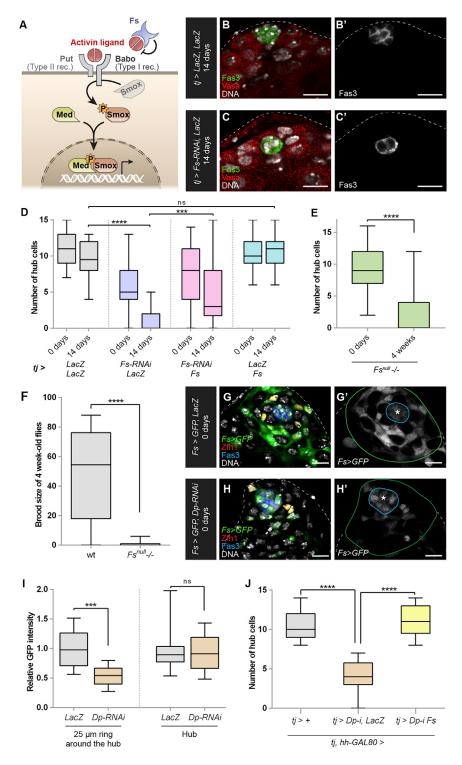


Figure 2. Follistatin acts downstream of Dp/ E2f1 in CySCs to maintain hub cells

(A) Model of the Activin pathway. Fs (purple) inhibits Activin ligands (red) binding to Activin receptors, Baboon (Babo, type I receptor) and Punt (Put, type II receptor). Receptor activation causes phosphorylation (orange P star) of the SMAD3 homolog Smox (inactive Smox is gray; active Smox is brown). Active Smox associates with the Co-SMAD Medea (Med, yellow) at regulatory sites of target genes to alter transcription.

(B and C) A control tj-GAL4 (labeled tj>LacZ, LacZ) adult testis has a normal number of hub cells (B). whereas a ti>Fs-RNAi, LacZ (C) adult testis has only 2 hub cells. Both testes were isolated after 14 days at 29°C to induce maximal GAL4 activity. Fas3 (green) labels hub cells, Vasa (red) marks germ cells, and DNA marked by DAPI is white.

(D) Graph showing the average number of hub cells at 0 and 14 days at 29°C in tj>LacZ, LacZ (gray bars, n = 49 and n = 22, respectively), tj > Fs - RNAi, LacZ (purple bars, n = 32 and n = 28, respectively), tj>Fs-RNAi, UAS-Fs (pink bars, n = 31 and n = 34, respectively), or tj>LacZ, UAS-Fs (blue bars, n = 37 and n = 31, respectively). See Table S1 for n values. (E) Graph showing the average number of hub cells at 0 days and 4 weeks in Fs<sup>null</sup> mutant (n = 18 and n = 31, respectively).

(F) Graph showing the fertility (brood size) at 4 weeks in a control and  $Fs^{null}$  mutant (n = 50 in

(G and H) Expression at 0 days of adulthood of Fs-GAL4 in a control testis (Fs>GFP, LacZ) or a testis in which Dp was depleted throughout development (Fs>GFP, Dp-RNAi). Fs-GAL4 is expressed strongly in CySCs and early cyst cells and weakly in hub cells (G) but its expression is substantially reduced when Dp is depleted (H). GFP (green) labels Fs-GAL4-expressing cells, Zfh1 (red) marks CySCs, Fas3 (blue) marks hub cells, and DNA marked by DAPI is white. Blue line surrounds hub cells and green line marks 25  $\mu M$  away from the

(I) Graph of relative GFP intensity in Fs>GFP, LacZ testes (labeled LacZ, gray bars, n = 13) or Fs>GFP, Dp-RNAi testes (labeled Dp-RNAi, brown bars, n = 13) in CySCs and early cyst cells (i.e., area between blue and green lines in (G and H) (labeled "25  $\mu$ M ring around the hub") and in hub cells (i.e., the area within the blue line, labeled "Hub").

(J) Graph of the number of hub cells in testes from control tj-GAL4, hh-GAL80 males (gray bar, labeled "tj>+," n = 27) or these males expressing Dp-RNAi and LacZ (brown bar, n = 8) or expressing Dp-RNAi and UAS-Fs (yellow bar, n = 17). UAS-Fs significantly rescues hub cell number compared with Dp-RNAi alone.

Error bars represent the data range. \*\*\*\* p < 0.0001; \*\*\* p < 0.001. ns = not significant as assessed by

See also Tables S1 and S2; Figures S3 and S4. Scale bar, 10 µM.

possibility, we permanently labeled the hub cell lineage with GFP (see STAR Methods, Figures 3D-3F). In testes where babo QD was overexpressed using upd<sup>TS</sup>, 67% of CySCs and early differentiating cyst cells (labeled by Zfh1) were GFP-positive after

14 days, indicating that they originated from hub cell transdifferentiation (Figures 3E and 3F). In contrast, in control testes, only 9% of Zfh1-positive CySCs and their immediate daughter cells were GFP-positive (p < 0.0001) (Figures 3D and 3F).





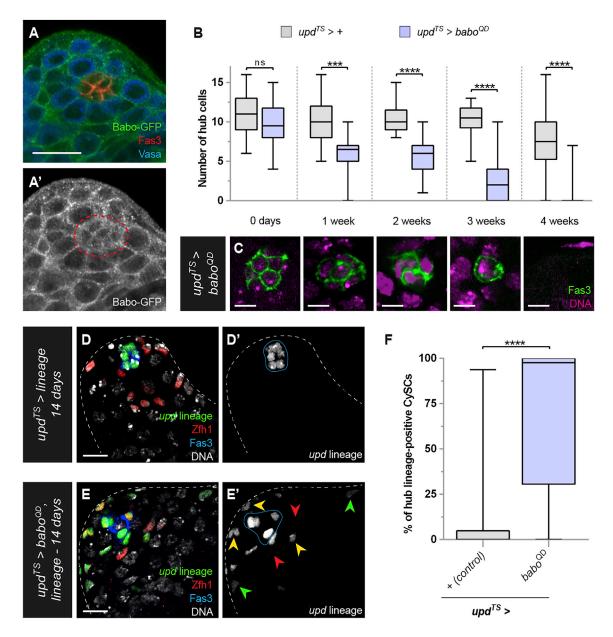


Figure 3. Autonomous Activin signaling disrupts hub cell quiescence

(A) Expression of Babo-GFP fusion protein (green, single channel in A') is detected in the testis apex. In particular, Babo-GFP expression is visible in hub cells (labeled with Fas3, red), outlined with a red dashed line in (A'). Vasa (blue) labels the germline.

(B) Graph showing the number of hub cells in control testes ( $upd^{TS}$ )+, gray bars, n = 23 for 0 days, n = 15 for 1 week, n = 13 for 2 weeks, n = 8 for 3 weeks, n =4 weeks) or those with sustained Activin signaling in hub cells (upd<sup>TS</sup>>babo<sup>QD</sup>, purple bars, n = 16 for 0 days, n = 16 for 1 week, n = 28 for 2 weeks, n = 35 for 3 weeks, n = 30 for 4 weeks) at the indicated time points. Note the progressive loss of hub cells in  $upd^{TS}$ > $babo^{QD}$ .

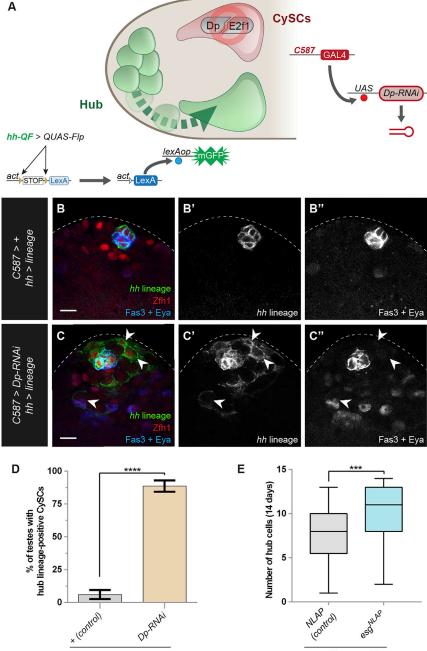
(C) Confocal sections of upd<sup>TS</sup>>babo<sup>QD</sup> testes at the indicated time points. Fas3 (green) marks hub cells and TO-PRO (magenta) marks DNA.

(D and E) Lineage tracing hub cells in control upd<sup>TS</sup>>+ (D) and upd<sup>TS</sup>>babo<sup>QD</sup> testes (E). Note in (E) the presence of hub-lineage-positive (green) cells outside of the niche that express the CySC marker Zfh1 (red). Fas3 (blue) marks hub cells and TO-PRO (DNA) is white. In (E'), yellow and red arrowheads indicate hub-lineagepositive CySCs and hub-lineage-negative CySCs, respectively, and the green arrowheads indicate differentiating cyst cells descended from a hub-lineage CySC. Blue lines in (C and D) indicate Fas3-positive hub cells.

(F) Graph indicating the percentage of CySCs positive for hub lineage in upd<sup>TS</sup>>+ (gray bar, n = 34) and upd<sup>TS</sup>>babo<sup>QD</sup> (purple bar, n = 26) testes. Error bars represent the data range. \*\*\*\*p < 0.0001; \*\*\*p < 0.001. ns, not significant as assessed by Student's t test. See also Tables S1 and S2.

Scale bar, 5  $\mu M$  in (C) and 10  $\mu M$  in all other panels.





#### Figure 4. CySC depletion of Dp results in non-autonomous hub-to-CySC transdiffer-

(A) Model indicating experimental design to deplete Dp from CySCs while concomitantly tracing hub cells using independent binary expression systems. To trace the lineage of hub cells, we used hh-QF, which is expressed only in hub cells, to induce QUAS-FLP. In turn, FLP recombines FRT sites in the act>STOP>LexA transgene. This leads to the production of LexA. Then LexA induces expression of lexAop-GFP, hereby exclusively labeling hub cells and their lineage with membrane GFP. In the same animal. C587-GAL4 drives expression of a Dp-RNAi transgene, which depletes Dp from CySCs but not from hub cells

(B and C) There are GFP-positive cells expressing 7fh1 (C. arrowheads) outside the cluster of hub cells in a C587>Dp-RNAi; hh>lineage testis but not in a control C587>+; hh>lineage testis (B). Hub lineage is in green, Zfh1 (red) labels CySCs, Fas3 (blue) labels hub cell membranes, and Eya (blue) labels the nuclei of differentiating cyst cells.

(D) Graph showing the percentage of testes in which hub-lineage-positive CySCs were present in C587>+ (control) (gray bar, n = 53) or C587>Dp-RNAi (brown bar, n = 49).

(E) Graph of the number of hub cells in testes in which baboQD was mis-expressed in hub cells (upd<sup>TS</sup>>) with either escargot (esg-NLAP, gray bar, n = 41) or the control transgene (NLAP, blue bar, n = 38).

Error bars represent the data range. \*\*\*\* p < 0.0001; \*\*\* p < 0.001 as assessed by Fisher's exact test (D) or Student's t test (E).

See also Tables S1 and S2, Figure S5. Scale bar, 10 uM.

C587 >  $upd^{TS} > babo^{QD}$ Since autonomous activation of Activin signaling resulted in (Figures S5E and S5F), a marker of CySC identity. Additionally, hub-to-CySC transdifferentiation, we asked whether knocking lineage-positive cells many cell diameters away from the nichte down Dp in CySCs also resulted in ectopic transdifferentiation of presented long membrane extensions and expressed the differeintiation marker Eya (Figures 4C and S5G), suggesting that transdifhub cells, leading to their eventual complete loss. We designed a strategy to trace the hub cell lineage, while simultaneously knockferentiated hub cells are functional CySCs, capable of proliferating ing down Dp in CySCs by employing orthogonal binary expression and differentiating. Taken together, our data demonstrate that strategies (Figures 4A and S5A, see STAR Methods). In control autonomously increasing Activin signaling in hub cells or nongenotypes, where Dp was not depleted in CySCs, hub-lineageautonomously depleting Dp from CySCs lead to transdifferentia-

ve CySCs were detected in 87%-89% of testes when Dp was depleted from all CySCs, a significant increase compared with controls (Figures 4C, 4D, S5B, S5D, S5F, and S5G). These cells lacked the hub cell marker Fas3 but expressed high levels of the CySC marker Zfh1, suggesting that they had adopted CySC identity (Leatherman and Dinardo, 2008) (Figures 4C, S5D, S5F, and S5G). Importantly, hub-lineage-expressing cells could incorporate the S-phase marker EdU

tion of hub cells into CySCs and that these new hub-derived

CySCs display all the hallmarks of stem cell behavior, namely

self-renewal and differentiation.

derived cells were detected infrequently outside the hub, in

6%-14% of testes (Figures 4B, 4D, S5B, S5C, and S5E) consistent

with prior results (Voog et al., 2014). By contrast, hub-lineage-pos-



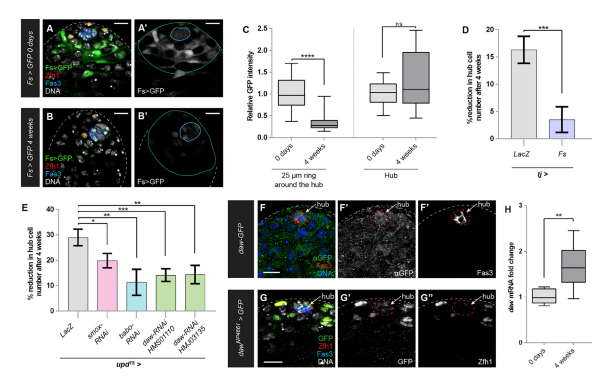


Figure 5. Increased Activin signaling is responsible for the decline of hub cell numbers during normal aging

(A-C) Expression at 0 days (A) and 4 weeks (B) of adulthood of Fs-GAL4, UAS-GFP in a control testis (Fs>GFP). Fs-GAL4 is expressed strongly in CySCs and early cyst cells and weakly in hub cells (A), and its expression is substantially reduced at 4 weeks of age (B). GFP (green) labels Fs-GAL4-expressing cells, Zfh1 (red) marks CySCs, Fas3 (blue) marks hub cells, and DNA labeled with DAPI is white. Blue line surrounds hub cells and green line marks a 25 μM ring from the hub. (C) Graph of relative GFP intensity in Fs>GFP testes at 0 days (light gray bars, n = 15) or 4 weeks (dark gray bars, n = 17) in CySCs and early cyst cells (i.e., area between blue and green lines in (A) and (B) (labeled "25 µM ring around the hub") and in hub cells (i.e., the area within the blue line, labeled "Hub"). (D) Graph showing the reduction in hub cell number after 4 weeks relative to newly eclosed flies in control tj>LacZ (gray bar, n = 49 at 0 days and n = 66 at 4 weeks) and in tj>Fs (purple bar, n=37 at 0 days and n=42 at 4 weeks).

(E) Graph showing the reduction in hub cell number after 4 weeks relative to newly eclosed flies in control upd>lacZ (gray bar, n = 45 at 0 days and n = 84 at 4 weeks); upd>Smox-RNAi (pink bar, n = 19 at 0 days and n = 47 at 4 weeks); upd>babo-RNAi (blue bar, n = 38 at 0 days and n = 26 at 4 weeks); upd>days RNAi-HMS01110 (first green bar, n = 33 at 0 days and n = 45 at 4 weeks); upd>daw RNAi-HMJ03135 (second green bar, n = 29 at 0 days and n = 46 at 4 weeks). (F and G) Expression of daw in the testis tip, as detected with a protein trap (F) or enhancer trap (G). Both reporters show weak expression in the hub (marked by Fas3, red in F, blue in G, and indicated with a dotted line). Daw-GFP protein is also detected outside the hub, and the enhancer trap reveals expression in Zfh1positive CySCs (red, G).

(H) Expression of daw mRNA increases with age in wild-type testes, as measured by qPCR (n = 8).

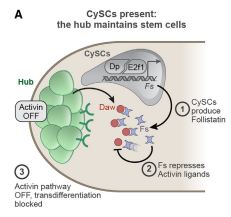
Error bars represent the data range. \*\*\*\* p < 0.0001; \*\*\* p < 0.001; \*\* p < 0.01; \* p < 0.05; ns = not significant, as assessed by Student's t test. See also Tables S1 and S2, Figure S6. Scale bar, 10  $\mu$ M.

We sought to define the autonomous relationship between Activin signaling and Esg, a Snail family transcriptional repressor required in hub cells to prevent their transdifferentiation into CySCs (Voog et al., 2014). We reasoned that excess Esg might be able to counteract the hub cell loss caused by Babo QD. To test this, we mis-expressed Esg while simultaneously inducing Activin signaling and found that sustained Esg expression suppressed hub cell loss induced by Babo QD (Figure 4E). This result indicates that Esg functions downstream of or in parallel to Activin signaling in hub cells to maintain hub cell identity.

#### The physiological age-dependent loss of hub cells depends on Activin signaling

Our experiments show that loss of Dp/E2f1 or Fs from all CySCs or increased Activin signaling in all hub cells results in loss of hub cells, indicating that CySCs signal to maintain quiescence of hub cells. Since lineage-wide insults are unlikely in normal physiological conditions, we asked what the functional significance of this CySC-to-hub cell signaling could be. We hypothesized that increased Activin signaling may be responsible for the decrease in the number of hub cells shown to occur during normal aging (Wallenfang et al., 2006; Boyle et al., 2007). Consistent with this, expression of the Fs>GFP transcriptional reporter significantly declined in CySCs in testes from 4-week-old flies compared with those from 0-day-old flies (Figures 5A-5C). Importantly, overexpressing Fs in CySCs blocked the loss of hub cells in 4-week-old males (Figure 5D). Although age had a significant effect on hub cell number (p = 0.0049), it displayed a strong interaction with genotype (p = 0.0067), indicating that the number of hub cells in Fs-overexpressing testes declined significantly less with age. These data implied that in older animals, reduced Fs levels could result in increased availability of Activin ligands to induce pathway signaling in hub cells. To test this model, we knocked down the Activin receptor babo or the Activin-dependent transcription factor Smox in hub cells. In both conditions, the decline in hub cell number was significantly





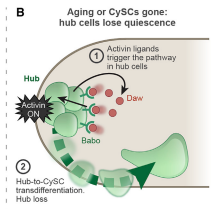


Figure 6. Model of the communication between CySCs and hub cells that underlies hub cell maintenance

(A) In a young wild-type testis, hub cells are quiescent. (1) Fs is produced by CySCs downstream of Dp/E2f1 either directly or indirectly. (2) Daw is produced by hub cells and CySCs. Daw is inactivated by Fs and does not activate its receptor Babo on hub cells. (3) In quiescent hub cells, Activin signaling is not activated.

(B) In an old testis or a testis lacking CySCs, (1) Fs expression declines, whereas increased Daw can now activate Babo on hub cells. (2) Activin signaling disrupts hub cell guiescence, leading to hub-to-CySC transdifferentiation.

lower after 4 weeks than in controls (Figure 5E). Finally, we asked which of the Activin ligands was responsible for the age-dependent loss of hub cells. After surveying transcriptional reporters and endogenously tagged Activin ligand lines (Figure S6), we found that only daw was expressed in the testis stem cell niche, as both a Daw protein trap (Figure 5F) and two daw-GAL4 transcriptional reporters (Figures 5G and S6A) were expressed in hub cells and in CySCs. The transcription of daw increased with age, suggesting that it could be partly responsible for the age-related decline of hub cell numbers (Figure 5H). Indeed, knockdown of daw in hub cells significantly suppressed their loss in aged testes (Figure 5E). In sum, our data demonstrate that the natural decline in hub cell number during aging is caused, at least in part, by increased Activin signaling in the hub leading to transdifferentiation of hub cells into CySCs.

#### **DISCUSSION**

We uncover a previously unknown aspect of the relationship between stem cells and their niches. Although traditionally thought of as unidirectional and top-down, with niche cells secreting selfrenewal factors essential for stem cells, prior work has shown that stem cells can induce their niche, both in normal development, and in malignant situations (Song et al., 2007; Ward et al., 2006; Patel et al., 2015). Here, we show that signals from the stem cells are required continually throughout the lifetime of the animal to maintain their niche. We find that the hub surveils signals from resident stem cells and acts as a reserve pool of stem cells if Activin increases (Figures 6A and 6B). Linking Fs production to Dp/E2f1 activity in CySCs ensures that the hub cells maintain their quiescence when surrounded with proliferating stem cells.

We show that expression of both Fs and daw in CySCs changes with age and demonstrate that reducing Activin signaling in hub cells significantly ameliorates age-dependent hub cell loss. However, it is interesting that Fs expression in hub cells does not decrease with age. Since hub-derived Fs can rescue the Fs<sup>null</sup> phenotype when overexpressed, suggesting that it is functional, it is likely that hub-derived Fs alone is not sufficient to fully inhibit Activin signaling. Similarly, both hub cells and CySCs express daw. We show that Daw produced by the hub is relevant to physiological aging and that its knock-

down produces a similar rescue to inhibiting Activin signaling in hub cells through babo or Smox knockdown. Thus, it is likely that hub-produced Daw is responsible for all the effects on hub cell numbers with age; however, it would be intriguing to test what the role of Daw produced by CySCs is, both in normal homeostasis and in age-dependent dysfunction.

In mammals, the Fs homolog FST is also involved in maintaining fertility via roles in somatic cells (Fullerton et al., 2017), although it has not yet been implicated in regulating the germline stem cell niche. Additionally, similar to our result that Activin signaling can force hub cells to exit quiescence, mammalian TGFβ/Activin signaling can reverse dormancy in disseminated tumor cells in the bone marrow, leading to tumor growth and metastases (Bragado et al., 2013; Ghajar et al., 2013).

Intriguingly, loss of retinoblastoma, the negative regulator of E2f/Dp transcription, resulted in perturbed interactions between hematopoietic cells and their niche (Walkley et al., 2007). Thus, cells may utilize the E2f/Dp transcription factor in many different contexts to transmit information about their cell-cycle state and allow effective monitoring of stem cells by the niche. Gaining a better understanding of the mechanisms that maintain niches in other stem cell models will enable targeting of the interactions between cancer stem cells and their supportive niches and provide new avenues for therapy (Plaks et al., 2015).

#### **Limitations of study**

Although our study showed no effect of Dp loss on CySC function, we cannot rule out that there are defects we failed to detect. Clones mutant for Dp persist and proliferate for up to 2 weeks after induction, but they may display defects when aged for longer periods. We showed decreased Fs expression upon knockdown of E2f1 or Dp. This could be due to direct or indirect regulation of the Fs locus by the E2f1/Dp transcription factor, which would need to be assessed by testing for occupancy of E2f1/Dp at the Fs locus. Finally, we show that ectopic Activin signaling promotes hub-to-CySC transdifferentiation and that changes in signaling with age are relevant to physiological hub cell loss. Both Fs and daw transcript levels change with age, but the mechanisms controlling these changes are unknown. It will be important in future studies to establish what directly regulates expression of these factors and how these regulators change during aging.

#### **Article**



#### **STAR**\*METHODS

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#### SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j. devcel.2021.07.010.

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#### **AUTHOR CONTRIBUTIONS**

Conceptualization, M.A. and E.A.B.; methodology, S.C.H., M.A., and E.A.B.; validation, M.A. and E.A.B.; formal analysis, S.C.H., M.A., and E.A.B.; investigation, S.C.H., M.A., D.S.d.I.M., L.G., S.M., R.P., and M.B.; resources, M.O., M.A., and E.A.B.; data curation, S.C.H., M.A., and E.A.B.; writing-original draft, S.C.H., M.A., and E.A.B.; writing-review and editing, S.C.H., M.A., and E.A.B.; visualization, S.C.H.; supervision, M.A. and E.A.B.; project administration, M.A. and E.A.B.; funding acquisition, M.A. and E.A.B.

#### **DECLARATION OF INTERESTS**

The authors declare no competing interests.

#### **INCLUSION AND DIVERSITY**

One or more of the authors of this paper self-identifies as an underrepresented ethnic minority in science.

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#### **STAR**\***METHODS**

#### **KEY RESOURCES TABLE**

Reagent or Resource	SOURCE	IDENTIFIER
Antibodies		
Mouse monoclonal anti-Fasciclin III (Fas3) (1:50)	DSHB	Cat# 7G10 anti-Fasciclin III; RRID: AB_528238
Rabbit polyclonal anti-GFP (1:500)	Invitrogen	Cat# A-6455; RRID: AB_221570
Goat polyclonal anti-Vasa (1:200)	Santa Cruz	Cat# sc26877; RRID: AB_793877
Chicken polyclonal anti-GFP (1:500)	Abcam	Cat# ab13970; RRID: AB_300798
Rat monoclonal anti-N Cadherin (1:20)	DSHB	Cat# DN-ex #8; RRID: AB_528121
Rat monoclonal anti DE-Cadherin (1:20)	DSHB	Cat# DCad2; RRID: AB_528120
Mouse monoclonal anti-Eyes absent Eya) (1:20)	DSHB	Cat# eya10H6; RRID: AB_528232
Rat monoclonal anti-Vasa	DSHB	Cat# anti-vasa; RRID: AB_760351
Mouse monoclonal anti- β-galactosidase (1;500)	Promega	Cat# Z3781; RRID: AB_430877
Mouse monoclonal anti-Dp (1:5)	N. Dyson (MGH Charlestown, USA	N/A
Rabbit polyclonal anti-Zfh1 (1:1000)	K. White (University of Chicago, USA)	N/A
Guinea pig polyclonal anti-Traffic jam (1:3000)	D. Godt (University of Toronto, Canada)	N/A
Cy3-AffiniPure Donkey Anti-Mouse IgG (1:400)	Jackson ImmunoResearch Labs	Cat# 715-165-150; RRID: AB_2340813
Alexa Fluor 488-AffiniPure Donkey Anti-Rabbit IgG (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 711-545-152; RRID: AB_2313584
Cy3-AffiniPure Donkey Anti-Rabbit gG (H+L)(1:400)	Jackson ImmunoResearch Labs	Cat# 711-165-152; RRID: AB_2307443
Cy5-AffiniPure Donkey Anti-Rabbit gG (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 711-175-152; RRID: AB_2340607
Alexa Fluor 488-AffiniPure Donkey Anti-Rat IgG (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 712-545-150; RRID: AB_2340683
Cy3-AffiniPure Donkey Anti-Rat gG (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 712-165-150; RRID: AB_2340666
Cy5-AffiniPure Donkey Anti-Rat gG (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 712-175-150; RRID: AB_2340671
Alexa Fluor 488 AffiniPure Donkey Anti-Chicken IgY (IgG) (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 703-545-155; RRID: AB_2340375
Cy3-AffiniPure Donkey Anti- Chicken IgY (IgG) (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 703-165-155; RRID: AB_2340363
Cy5-AffiniPure Donkey Anti- Chicken IgY (IgG) (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 703-175-155; RRID: AB_2340365
Cy3-AffiniPure Donkey Anti-Guinea Pig IgG (1:400)	Jackson ImmunoResearch Labs	Cat# 706-165-148; RRID: AB_2340460
Cy5-AffiniPure Donkey Anti-Guinea Pig IgG (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 706-175-148; RRID: AB_2340462
Cy3-AffiniPure Donkey Anti-Goat gG (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 705-165-003; RRID: AB_2340411
Alexa Fluor 647 AffiniPure Donkey Anti-Goat IgG (H+L) (1:400)	Jackson ImmunoResearch Labs	Cat# 705-605-003; RRID: AB_2340436
Chemicals, peptides, and recombinant proteins		
/ECTASHIELD Mounting Medium	Vector Laboratories	Cat# H-1000; RRID: AB_2336789
/ECTASHIELD Mounting Medium with DAPI	Vector Laboratories	Cat# H-1200; RRID: AB_2336790
Paraformaldehyde, 16% w/v aq. soln., methanol free	Thermo Fisher Scientific	Cat# 43368-9L
ΓΟ-PRO-3 iodide (1 μM)	Invitrogen	Cat# T3605
5-ethynyl-2'-deoxyuridine (EdU)	Invitrogen	Cat# A10044

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Continued		
Reagent or Resource	SOURCE	IDENTIFIER
AF405 picolylazide	Click chemistry tools	Cat# 1308-5
Tris(3-Hydroxyproyltriazolylmethyl)Amine	Sigma Aldrich	Cat# 762342
Sodium Ascorbate	Sigma Aldrich	Cat# PHR1279
Copper(II) sulfate	Fisher Scientific	Cat# 15617730
Molasses	Labscientific	Cat# FLY-8008-16
Agar	Mooragar	Cat# 41004
Cornmeal	LabScientific	Cat# FLY-8010-20
Yeast	LabScientific	Cat#FLY-8040-20F
Tegosept	Sigma	Cat# H3647-1KG
Reagent alcohol	Fisher	Cat# A962P4
Propionic acid	Fisher	Cat# A258500
Experimental models: organisms/strains		
D. melanogaster; Oregon <sup>R</sup>	Bach lab	N/A
D. melanogaster; P[GawB]NP1624/CyO (tj-GAL4)	Bach lab	Kyoto Stock Center: 104055 Flybase: FBst0302922
D. melanogaster; nos-GAL4-VP16	Ruth Lehmann (Whitehead Institute, USA)	N/A
D. melanogaster; C587-GAL4	Ruth Lehmann (Whitehead Institute, USA)	N/A
D. melanogaster; fng-GAL4	Stephen Dinardo (Perelman School of Medicine, University of Pennsylvania, USA)	N/A
D. melanogaster; eyaA3-GAL4 (denoted eya-GAL4)	Stephen Dinardo (Perelman School of Medicine, University of Pennsylvania, USA)	N/A
D. melanogaster; upd-GAL4	Bach lab	N/A
D. melanogaster; tub-GAL80 <sup>TS</sup>	Bloomington Drosophila stock Center (BDSC)	BDSC_7017 FlyBase: FBst0007017
D. melanogaster; hh-LacZ	Bach lab	BDSC_5530 FlyBase: FBst0005530
D. melanogaster; upd-LacZ	Bach lab	N/A
D. melanogaster; hh-GAL80	This paper	N/A
D. melanogaster; hh-QF	This paper	N/A
D. melanogaster; w <sup>1118</sup> ; P[w[+mC]=UAS-lacZ.NZ]J312. Insertion on III	BDSC	BDSC_3956 FlyBase: FBst003956
D. melanogaster; w <sup>1118</sup> ; P[w[+mC]=UAS- lacZ.NZ]20b.Insertion on II	BDSC	BDSC_3955 FlyBase: FBst003955
D. melanogaster; UAS-GFP; Ubi- p63E(FRT.STOP)Stinger	BDSC	BDSC_32251
D. melanogaster; w <sup>1118</sup> ; P[GD4444]v12722 (Dp-RNAi)	Vienna <i>Drosophila</i> Resource Center (VDRC)	VDRC_v12722 FlyBase: FBst0450633
D. melanogaster; FRT <sup>42D</sup> , Dp <sup>a3</sup> /CyO	M. Frolov, University of Illinois at Chicago, USA	Flybase: FBgn0011763
D. melanogaster; FRT <sup>42D</sup> , Dp <sup>a4</sup> /CyO	M. Frolov, University of Illinois at Chicago, USA	Flybase: FBgn0011763
D. melanogaster; UAS-Dp	M. Frolov, University of Illinois at Chicago, USA	Flybase: FBgn0011763
D. melanogaster; FRT <sup>e2B</sup> E2f1 <sup>729</sup>	M. Frolov, University of Illinois at Chicago, USA	FlyBase: FBgn0011766
D. melanogaster; y¹, v¹; P[TRiP.JF02519] attP2/TM3, Sb¹ (Dp-RNAi)	BDSC	BDSC_30515 FlyBase: FBst0030515

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Continued		
Reagent or Resource	SOURCE	IDENTIFIER
D. melanogaster, w <sup>1118</sup> ; P[GD4448] v15886 (E2f1-RNAi)	VDRC	VDRC: v15886 FlyBase: FBst0452055
D. melanogaster, UAS-Rbf <sup>280</sup> Insertion on III	W. Deng (Tulane University, USA)	BDSC_50748 Flybase: FBst0050748
D. melanogaster, act>y[+]>LHV2- 86Fb,13XlexAop2-myr::GFP (FLEX-AMP)	C. Desplan (New York University, USA)	N/A
D. melanogaster; UAS-FLP	BDSC	BDSC_4539
D. melanogaster; QUAS-FLP	BDSC	BDSC_30126
D. melanogaster; w <sup>1118</sup> ; P[GD15843]v46260 (Fs-RNAi)	VDRC	VDRC v46260 FlyBase: FBst0466595
D. melanogaster; y <sup>1</sup> v <sup>1</sup> ;P[TRiP.HMJ03135] attP40 (daw-RNAi)	BDSC	BDSC_50911 Flybase: FBst0050911
D. melanogaster; P[KK110248]VIE-260B (daw-RNAî)	VDRC	VDRC #v105309 Flybase: FBst0477137
D. melanogaster; y¹sc* v¹ sev²¹; P[TRiP.HMS01110]attP2 (daw-RNAi)	BDSC	BDSC_34974 Flybase: FBst0034974
D. melanogaster; y <sup>1</sup> v <sup>1</sup> ; P[TRiP.GL01476]attP2 (smox-RNAi)	BDSC	BDSC_43138 Flybase: FBst0043138
D. melanogaster; y <sup>1</sup> w*; Mi[MIC]Fs <sup>MI01433</sup> (Fs <sup>MI01433</sup> )	BDSC	BDSC_33121 Flybase: FBst0033121
D. melanogaster; y <sup>1</sup> w*; Mi[MIC]Fs <sup>MI11350</sup> (Fs <sup>MI11350</sup> )	BDSC	BDSC_56310 Flybase: FBst0056310
D. melanogaster; y <sup>1</sup> w ;Fs <sup>MI04308-TG4.1</sup> CG8079 <sup>MI04308-TG4.1-X</sup> (Fs <sup>TJ4.1</sup> )	BDSC	BDSC_66838 (lost during the pandemic)
D. melanogaster; y <sup>1</sup> w <sup>*</sup> ; Mi[PT-GFSTF.1] Fs <sup>MI04308-GFSTF.1</sup> CG8079 <sup>MI04308-GFSTF.1-X</sup> /CyO (Fs <sup>GFSTF.1</sup> or Fs-GFP)	BDSC	BDSC_65327 Flybase: FBst0065327
D. melanogaster; Fs <sup>null</sup>	This paper	N/A
D. melanogaster; UAS-Fs	O'Connor lab	N/A
D. melanogaster; Fs-GAL4 (Fs <sup>MI</sup> -GAL4)	This paper	N/A
D. melanogaster; UAS-esgNLAP	L. Jones (UCSF, USA)	N/A
D. melanogaster; UAS-NLAP	L. Jones (UCSF, USA)	N/A
D. melanogaster; Actβ-GAL4	O'Connor lab	N/A
D. melanogaster; myo-GAL4	O'Connor lab	N/A
D. melanogaster; y <sup>1</sup> w <sup>*</sup> ; Mi[MIC]daw <sup>MI05383</sup> (daw-GFP)	BDSC	BDSC_43001 Flybase: FBst0043001
D. melanogaster; UAS-babo-RNAi.	O'Connor lab	N/A
D. melanogaster; UAS-babo <sup>QD</sup>	O'Connor lab	N/A
D. melanogaster; PBac[fTRG00506. sfGFP-TVPTBF]VK00033 (Actβ <sup>fTRG00506.sfGFP-TVPTBF</sup> )	VDRC	VDRC #v318136 Flybase: FBst0491562
D. melanogaster; PBac[fTRG00161.sfGFP-TVPTBF]VK00033 (myo <sup>fTRG00161.sfGFP-TVPTBF</sup> )	VDRC	VDRC #v318065 Flybase: FBst0491390
D. melanogaster; PBac[fTRG00444.sfGFP-TVPTBF]VK00033 [babo <sup>fTRG00444.sfGFP-TVPTBF</sup> )	VDRC	VDRC #v318433 Flybase: FBst0491516
D. melanogaster; w <sup>*</sup> ; P[GawB] daw <sup>NP4661</sup> / CyO (daw <sup>NP4661</sup> -GAL4)	Kyoto Stock Center	Kyoto #113490 Flybase: FBst0316217
D. melanogaster; y <sup>*</sup> w <sup>*</sup> ; P[GawB] daw <sup>NP6274</sup> / CyO (daw <sup>NP6274</sup> -GAL4)	Kyoto Stock Center	Kyoto #105179 Flybase: FBst0304038

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Continued		
Reagent or Resource	SOURCE	IDENTIFIER
Oligonucleotides		
gRNA Fs 1	5' CCGGTTGCATCATGTATCTTGGC 3'	IDTDNA
gRNA Fs 2	5' CCAACTGGAGGTCGCCTATCGGG 3'	IDTDNA
qPCR primer Fs-fwd:	5'-AGTGTCATATATACTCTCCGCATGT-3'	IDTDNA
qPCR primer Fs-rev:	5'-ACAGCAACTGCTTTTTAACTATGCC-3'	IDTDNA
qPCR primer α-tub84B-fwd:	5'-TCGTTTTACGTTTGTCAAGCCTC-3'	IDTDNA
qPCR primer α-tub84B-rev:	5'-GAGATACATTCACGCATATTGAGTT-3'	IDTDNA
qPCR primer daw-fwd	5'-CCCATCTTCGACGGGATGAC-3'	IDTDNA
qPCR primer <i>daw-rev</i>	5'-TTGCACTCGACCTCCTCT-3'	IDTDNA
Software and algorithms		
lmageJ/Fiji	Fiji	http://fiji.sc/
Photoshop/Illustrator	Adobe	https://www.adobe.com/products/
Prism	GraphPad	https://www.graphpad.com
ZEN	Zeiss	https://www.zeiss.com/microscopy/us/ products/microscope-software/zen.html
Excel	Microsoft	https://products.office.com/en-us/excel
Imaris	Oxford Instruments	https://imaris.oxinst.com/

#### RESOURCE AVAILABILITY

#### **Lead contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Erika Bach (erika.bach@nyu.edu).

#### **Materials availability**

All Drosophila stocks generated in this study are available from the Lead Contact without restriction.

#### Data and code availability

- All data reported in this paper will be shared by the Lead Contact upon request.
- This paper does not report original code
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request

#### **EXPERIMENTAL MODEL AND SUBJECT DETAILS**

#### **Drosophila stocks and maintenance**

Drosophila melanogaster strains used in this study are listed in the key resources table. Drosophila were reared on food made with these ingredients: 1800mL molasses (LabScientific, Catalog no. FLY-8008-16), 266 g agar (Mooragar, Catalog no. 41004), 1800 g cornmeal (LabScientific, Catalog no. FLY-8010-20), 744g Yeast (LabScientific, Catalog no. FLY-8040-20F), 47 L water, 56 g Tegosept (Sigma no. H3647-1KG), 560mL reagent alcohol (Fisher no. A962P4), and 190mL propionic acid (Fisher no. A258500).

Flies were kept at 25°C, except crosses with GAL80<sup>TS</sup>, which were maintained at 18°C until eclosion, and the adult flies were transferred to 29°C. Clonal experiments were analyzed at 2 dpci to assess clone induction and 7 and 14 dpci for maintenance as is standard in the field (Amoyel et al., 2014). For transgene expression in CySCs, we examined hub cell number at 0, 3, 7, 10 and 20 days of adulthood. Our reason for selecting these time points is that Dp knockdown by tj-GAL4 led to an almost complete loss of hub cells by 10 days, but since other genotypes exhibited a slower loss of hub cells, we also examined hub cells at 20 days of adulthood. We looked at earlier time points (3 and 7 days) to examine hubs before their complete loss. For the aging experiments, we examined hub cell number at 1, 2, 3, or 4 weeks (or 7, 14, 21, 28 days) of adulthood.

Males were aged separately from females and provided with fresh food every other day. For adult-onset overexpression and RNAidepletion, we used the appropriate driver combined with the temperature-sensitive repressor tub-GAL80<sup>TS</sup>. Flies were reared at the permissive temperature (18°C) and adult males were collected and shifted to the restrictive temperature of 29°C to allow GAL4 activity. Experiments without GAL80 were raised at room temperature until eclosion and shifted to 29°C for the specified period to ensure maximum GAL4 activity.

#### **Article**



We used the following fly stocks: Oregon-R; tj-GAL4 (Kyoto Stock Center #NP1624); nos-GAL4-VP16 (gift of Ruth Lehmann, Whitehead Institute, USA; upd-LacZ (Tsai and Sun, 2004); hh-LacZ; tub-GAL80<sup>TS</sup> (McGuire et al., 2004); hh-GAL80 (this study); hh-QF (this study); UAS-LacZ (Bloomington Drosophila Stock center (BDSC) #3955 and 3956); UAS-GFP; Ubi-p63E(FRT.STOP) Stinger (also known as GTRACE, BDSC # 32251); UAS-Dp RNAi (Vienna Drosophila Resource Center (VDRC) #v12722); FRT<sup>42D</sup> Dp<sup>a3</sup>andFRT<sup>42D</sup> Dp<sup>a4</sup> (both gifts of M. Frolov, University of Illinois at Chicago, USA); FRT<sup>62B</sup> E2f1<sup>729</sup> (gift of M. Frolov); UAS-Dp RNAi (BDSC #30515); UAS-Dp (Zappia and Frolov, 2016) (gift of M. Frolov); UAS-E2f1 RNAi (VDRC #v15886); UAS-Rbf<sup>280</sup> (BSDC #50748); act>y[+]>LHV2-86Fb,13XlexAop2-myr::GFP (also known as FLEXAMP (Bertet et al., 2014)) where LHV2-86Fb encodes an optimized version of LEXA-VP16 with reduced toxicity; UAS-FLP (BDSC #4539); QUAS-FLP (BDSC #30126); UAS-Fs-RNAi (GD15843, VDRC #v46260); UAS-daw-RNAi (HMJ03135 BDSC #50911; HMS01110 BDSC #34974; and KK110248 VDRC #v105309); UAS-smox-RNAi (GL01476 BDSC #43138); Fs<sup>MI01433</sup> (BDSC #33121); Fs<sup>MI11350</sup> (BDSC #56310); Fs<sup>MI04308-TG4.1</sup>  $CG8079^{MI04308-TG4.1-X}$  (referred to as  $Fs^{TJ4.1}$ , BDSC #66838);  $Fs^{MI04308-GFSTF.1}$   $CG8079^{MI04308-GFSTF.1-X}$  (referred to as  $Fs^{GFSTF.1}$  or Fs-GFP) BDSC #65327); Fs<sup>null</sup> (this study); UAS-Fs (Pentek et al., 2009); Fs-GAL4 (referred to also as Fs<sup>MI</sup>-GAL4) (this study); UAS-esgNLAP and UAS-NLAP (both gifts from Leanne Jones, UCLA, USA) (Voog et al., 2014); Actβ-GAL4 (Song et al., 2017); myo-GAL4 (Awasaki et al., 2011); daw MI05383 (referred to as daw-GFP, BDSC #43001); UAS-babo-RNAi (Peterson and O'Connor, 2013); UAS-babo<sup>QD</sup>;  $Act\beta^{FRG00506.sfGFP-TVPTBF}$  (Sarov et al., 2016), VDRC #v318136);  $myo^{fRG00161.sfGFP-TVPTBF}$  (Sarov et al., 2016), VDRC #v318065); babo<sup>fTRG00444.sfGFP-TVPTBF</sup> (Sarov et al., 2016), VDRC #v318433); punt (put)<sup>fTRG00910.sfGFP-TVPTBF</sup> (Sarov et al., 2016). 2016), VDRC #v318264); daw<sup>NP4661</sup>-GAL4 and daw<sup>NP6274</sup>-GAL4, UAS-LacZ (Kyoto Stock Center #113490 and #105179, respectively).

#### **METHOD DETAILS**

#### Generation of transgenic Drosophila lines

The Fs-GAL4 line used to monitor Fs transcriptional activity was created by recombination mediated cassette exchange (RMCE) to insert GAL4 in the MiMIC line Fs<sup>MIO1433</sup> (Venken et al., 2011). The Fs<sup>null</sup> mutant was created by GenetiVision using CRISPR/Cas9 to delete 3 kb of the Fs locus, corresponding to the first 4 coding exons of both predicted Fs isoforms (Figure S3D). This genomic region was replaced by a splicing acceptor site and stop codons in all the 3 frames, followed by a 3xP3-GFP cassette. We used CCGGTTGCATCATGTATCTTGGC and CCAACTGGAGGTCGCCTATCGGG as gRNAs. We validated the mutant by sequencing the PCR product using the genomic DNA from the Fs<sup>null</sup> mutant as a template and the following primers:

fwd 5' CGGTGCATAATGCGCCAAACC 3' rev 5' CTTGCAGTGCACTGGATATGG 3'

hh-GAL80 and hh-QF were generated by RMCE by injection into line MI10526 (BDSC #53865), carrying a MiMIC transposon in the hh locus, using plasmids from the Drosophila Genomics Resource Center (DGRC) #1390 (Diao et al., 2015) and #1296 (Venken et al., 2011), respectively.

#### **Drosophila** genetics

To label the lineage of hub cells, we used *upd-GAL4* and *tub-GAL80<sup>TS</sup>* to drive expression of the *UAS-FLP* recombinase, which excises a stop from the *ubiP63E(FRT.STOP)Stinger* G-TRACE cassette (Evans et al., 2009). After the excision of FRT sites following recombination, the resulting GFP serves an indelible and persistent marker to identify CySCs derived from the hub lineage. For tracing the hub lineage while manipulating expression in the cyst lineage, we used *C587-GAL4* to knock down *Dp* in CySCs, and an orthogonal expression system, *hh-QF*, to drive *QUAS-FLP* expression in hub cells. The lineage was permanently marked upon FLP expression using either the GTRACE cassette, as above, or the FLEXAMP cassette, in which a *yellow*<sup>+</sup> transgene is excised by FLP, leading to permanent expression of LexA under the *actin* promoter. LexA labels cells through expression of myr-GFP, driven by multimerized *LexAop* binding sites.

For mutant clones, crosses were raised at 25°C until eclosion. Males were collected for up to 2 days, then heat-shocked at 37°C for 1 h and returned to 25°C until dissection, 2, 7 or 14 days later.

#### **Antibodies and immunofluorescence**

The following primary antibodies were used: goat anti-Vasa (1:200; Santa Cruz), mouse anti-Fasciclin3 (Fas3) (1:50; Developmental Studies Hybridoma Bank (DHSB), mouse anti-Eya (1:20, DSHB), rat anti-Vasa (1:20, DSHB), rat anti-N-Cadherin (1:20, DSHB), rabbit anti-GFP (1:500; Invitrogen), chicken anti-GFP (1:500, Aves Labs), mouse anti- $\beta$ -Galactosidase (1:500, Promega), guinea pig anti-Tj (1:3000, a gift of Dorothea Godt, University of Toronto, Canada), rabbit anti-Zfh1 (1:1000), mouse anti-Dp (1:5, a gift of Nick Dyson, MGH Charlestown, USA), TO-PRO-3 iodide (1  $\mu$ M; Molecular Probes); 4′,6-diamidino-2-phenylindole (DAPI) (1:500, Invitrogen). Donkey secondary antisera were used 1:400 (Jackson ImmunoResearch). Dissections and staining were carried out as previously described (Flaherty et al., 2010). In brief, testes were dissected in 1x phosphate buffered saline (PBS), fixed for 15 min in 4% formaldehyde in 1xPBS, washed for 1 h at 25°C in 1xPBS with 0.5% Triton X-100, and blocked in PBTB (1xPBS 0.2% Triton X-100 and 1% bovine serum albumin) for 1 h at 25°C. Primary antibodies were incubated overnight at 4°C. They were washed two times for 30 min in PBTB and incubated 2 h in secondary antibody in PBTB at 25°C and then washed two times for 30 min in 1x PBS with 0.2% Triton X-100. They were mounted in Vectashield or Vectashield + DAPI (Vector Laboratories). 5-ethynyl-2′-deoxyuridine (EdU) labeling was achieved by incubating dissected samples in 10 $\mu$ M EdU (Life Technologies) in Schneider's medium prior to fixation. Following



primary and secondary antibody incubations and washes, samples were incubated for 30 min in buffer containing 0.1M THPTA, 2mM sodium ascorbate, 1mM CuSO<sub>4</sub> and 2.5µM picolylazide conjugated to a fluorophore (Alexa 405, 488, 568 or 643, Click Chemistry Tools). Samples were then washed and mounted for imaging. Confocal images were captured using Zeiss LSM 510, LSM 700, LSM 880 and a Leica Sp8confocal microscopeswith a 63x objective.

#### **Quantitative RT-PCR**

Whole testes (n=20-25) were isolated and mRNA was extracted with PicoPure RNA Isolation system (Applied Biosystems) following the manufacturer's instructions. Reverse transcription was performed using Maxima Reverse Transcriptase (ThermoFisher) as per manufacturer's instructions and using 0.5µg of RNA as template. gRT-PCR was performed using SYBR Green PCR Master Mix (ThermoFisher) and a QuantStudio 5 Real-Time PCR System machine (Applied Biosystems). 8 replicates were performed for the experiments in Figures 5H and 4 replicates for each experiment shown in Figure S3E. To detect Fsisoform B (FlyBase code FBtr0339996), we used the following primers and normalized expression levels to the control gene α-tub84B:

Fs-fwd: 5'-AGTGTCATATATACTCTCCGCATGT-3' Fs-rev: 5'-ACAGCAACTGCTTTTTAACTATGCC-3' α-tub84B-fwd: 5'-TCGTTTTACGTTTGTCAAGCCTC-3' α-tub84B-rev: 5'-GAGATACATTCACGCATATTGAGTT-3' daw-fwd: 5'-CCCATCTTCGACGGGATGAC-3' daw-rev: 5'-TTGCACTCGACCTCCTCT-3'

We failed to detect transcripts of Fs isoform A in the testis (FlyBase FBtr0087398), using the following primers and normalized expression levels to the control gene  $\alpha$ -tub84B:

Fs-A-fwd:5'-GAACGGACCGCGCTAAAAAT-3' Fs-A-rev:5'-GGCAAACGCACTGGTTTCAT-3'

#### **Fertility tests**

Mutant males were crossed individually with two Oregon<sup>R</sup> females. After 48 h of mating and egg laying, all adults were removed from the vials. The size of the brood of each male was scored ten days later.

#### **QUANTIFICATION AND STATISTICAL ANALYSIS**

#### **Data analysis and statistics**

Hub cells were quantified by counting nuclei (labelled with DAPI, or low levels of Ti or Zfh1) in cells that were also Fas3-positive. For GFP intensity quantifications, we dissected control flies and experiments on the same day and processed them simultaneously. Images were acquired with the same microscope settings. Plots show mean fluorescence intensity normalized to the average value of the controls.

The percentage of transdifferentiated CySCs was calculated as the ratio of Zfh1-positive cells simultaneously positive for the hub cell-lineage labeling.

Image processing, quantifications, and figure preparation were performed with Fiji-ImageJ (Schindelin et al., 2012), Adobe Photoshop and Adobe Illustrator software. Statistical tests were performed using GraphPad Prism8 or JMP software. To test for an interaction between genotype and aging in Figures 5D and 5E, we used a linear model featuring full factorial design and tested for the effect of both age and genotype on hub cell number and their interaction. Categorical data analysis in Figures 4D and S1B, S1G, and S5B was performed as described in (Xu et al., 2010) using Fisher's Exact tests. Other data were analyzed with Student's t tests because they were pairwise comparisons between genotypes. Data were analyzed and plotted with GraphPad Prism8. In all graphs, whiskers indicate the entire range of data, the boxes show the second and third quartiles and the line shows the median.

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### **Supplemental information**

Proliferative stem cells maintain quiescence of their niche by secreting the Activin inhibitor Follistatin

Salvador C. Herrera, Diego Sainz de la Maza, Lydia Grmai, Shally Margolis, Rebecca Plessel, Michael Burel, Michael O'Connor, Marc Amoyel, and Erika A. Bach

Table S1: Hub cell number in the analyzed genotypes, Related to Figures 1-5 and STAR methods

#	Figure	Genotype	Treatment	Average	SE	n	р
1	11	<i>tj</i> > +	10 days	11.83	±0.53	18	
2	11	tj > Dp-RNAi (Vienna), GFP	10 days	2.10	±0.43	31	1e <sup>-16</sup> (vs #1)
3	11	tj > Dp-RNAi (TRiP)	10 days	9.14	±0.96	14	0.0232 (vs #1)
4	11	tj > Dp-RNAi, Dp	10 days	4.93	±0.86	15	0.0075 (vs #2)
5	11	tj > E2f1-RNAi	10 days	7.29	±0.64	17	5e <sup>-6</sup> (vs #1)
6	11	$tj > Rbf^{280}$	10 days	5.00	±1.50	12	0.0008 (vs #1)
7	1J	$tj$ , $hh$ - $GAL80^{TS} > +$	10 days	10.56	±0.32	27	` ,
8	1J	tj, hh-GAL80 <sup>⊤S</sup> > Dp- RNAi, LacZ	10 days	4.00	±0.76	8	2e <sup>-5</sup> (vs #7)
9	1J	tj, hh-GAL80 <sup>™</sup> > Dp- RNAi, Dp	10 days	8.36	±0.35	33	0.0004 (vs #8)
10	2D	tj > LacZ, LacZ	0 days	10.92	±0.30	49	
11	2D	tj > LacZ, LacZ	14 days	9.69	±0.54	22	
12	2D	tj > Fs-RNAi, LacZ	0 days	6.06	±0.48	32	
13	2D	tj > Fs-RNAi, LacZ	14 days	1.11	±0.28	28	2e <sup>-15</sup> (vs #11)
14	2D	tj > Fs-RNAi, Fs	0 days	7.52	±0.71	31	,
15	2D	tj > Fs-RNAi, Fs	14 days	4.56	±0.68	34	3e <sup>-5</sup> (vs #13)
16	2D	tj > LacZ, Fs	0 days	10.49	±0.30	37	,
17	2D	tj > LacZ, Fs	14 days	10.58	±0.43	31	0.20 (vs #11)
18	2E	Fs <sup>null</sup> -/-	0 days	9.39	±0.85	18	,
19	2E	Fs <sup>null</sup> -/-	4 weeks	2.16	±0.63	31	6e <sup>-8</sup> (vs #18)
20	2J	$tj$ , $hh$ - $GAL80^{TS} > +$	10 days	10.56	±0.32	27	,
21	2J	tj, hh-GAL80 <sup>⊤S</sup> > Dp- RNAi, LacZ	10 days	4.00	±0.76	8	2e <sup>-5</sup> (vs #20)
22	2J	tj, hh-GAL80 <sup>⊤s</sup> > Dp- RNAi, Fs	10 days	10.88	±0.45	17	4e <sup>-6</sup> (vs #21)
23	3B	$upd^{TS} > +$	0 days	11.17	±0.74	23	
24	3B	$upd^{TS} > +$	1 week	10.33	±0.73	15	
25	3B	$upd^{TS} > +$	2 weeks	10.62	±0.58	13	
26	3B	upd <sup>TS</sup> > +	3 weeks	10.13	±0.85	8	
27	3B	upd <sup>TS</sup> > +	4 weeks	7.68	±0.38	84	
28	3B	upd <sup>TS</sup> > babo <sup>QD</sup>	0 days	9.88	±0.68	16	0.2032 (vs #23)
29	3B	upd <sup>TS</sup> > babo <sup>QD</sup>	1 week	6.25	±0.62	16	0.0002 (vs #24)
30	3B	upd <sup>TS</sup> > babo <sup>QD</sup>	2 weeks	5.18	±0.47	28	8e <sup>-8</sup> (vs #25)
31	3B	upd <sup>TS</sup> > babo <sup>QD</sup>	3 weeks	2.69	±0.47	35	7e <sup>-6</sup> (vs #26)
32	3B	upd <sup>TS</sup> > babo <sup>QD</sup>	4 weeks	0.60	±0.28	30	7e <sup>-28</sup> (vs #27)
33	4E	upd TS > NLAP	4 weeks	7.68	±0.48	41	,
34	4E	$upd^{TS} > esg^{NLAP}$	4 weeks	10.24	±0.50	38	0.0004 (vs #33)
41	5D	tj > LacZ	0 days	10.92	±0.30	49	,
42	5D	tj > LacZ	4 weeks	9.14	±0.27	66	
43	5D	tj > Fs	0 days	10.49	±0.30	37	
44	5D	tj > Fs	4 weeks	10.45	±0.38	42	0.0058 (vs #42)

4 -	FF	un alTS > 1 7	O al	10.04	10.40	4.5	
45	5E	upd <sup>TS</sup> > LacZ	0 days	10.64	±0.42	45	
46	5E	upd <sup>TS</sup> > LacZ	4 weeks	7.56	±0.34	84	
47	5E	upd <sup>TS</sup> > smox-RNAi	0 days	11.79	±0.40	19	
48	5E	upd <sup>™</sup> > smox-RNAi	4 weeks	9.45	±0.33	47	0.0007 (vs
49	5E	upd <sup>TS</sup> > babo-RNAi	0 days	11.45	±0.45	38	#46)
50	5E	upd <sup>TS</sup> > babo-RNAi	4 weeks	10.15	±0.43	26	0.0042 (vs
50		•	4 WEEKS	10.15	±0.56		#46)
51	5E	upd <sup>⊤s</sup> > daw-RNAi- HMS01110	0 days	12.97	±0.41	33	
52	5E	upd <sup>⊤s</sup> > daw-RNAi- HMS01110	4 weeks	11.13	±0.32	45	2e <sup>-34</sup> (vs #46)
53	5E	upd <sup>⊤s</sup> > daw-RNAi- HMJ03135	0 days	12.24	±0.34	29	
54	5E	upd <sup>⊤s</sup> > daw-RNAi- HMJ03135	4 weeks	10.48	±0.45	46	0.0010 (vs #46)
55	S2E	$tj^{TS} > +$	0 days	11.88	±0.33	17	,
56	S2E	$ti^{TS} > +$	3 days	12.05	±0.33	21	
57	S2E	$ti^{TS} > +$	7 days	11.63	±0.50	8	
58	S2E	$tj^{TS} > +$	10 days	11.69	±0.29	35	
59	S2E	$tj^{TS} > +$	20 days	10.76	±0.47	17	
60	S2E	$tj^{TS} > Dp-RNAi$	0 days	11.18	±0.38	28	0.1686 (vs #55)
61	S2E	$tj^{TS} > Dp-RNAi$	3 days	9.79	±0.68	14	0.0076 (vs #56)
62	S2E	$tj^{TS} > Dp-RNAi$	7 days	5.00	±1.59	6	0.0074 (vs #57)
63	S2E	tj™ > Dp-RNAi	10 days	2.31	±0.47	39	6e <sup>-20</sup> (vs #58)
64	S2E	tj <sup>TS</sup> > Dp-RNAi	20 days	0.03	±0.03	30	1e <sup>-13</sup> (vs #59)
65	S2E	tj <sup>⊤S</sup> > E2f1-RNAi	0 days	10.61	±0.78	18	0.2699 (vs #55)
66	S2E	$tj^{TS} > E2f1-RNAi$	3 days	9.58	±0.55	17	0.0083 (vs #56)
67	S2E	tj <sup>™</sup> > E2f1-RNAi	7 days	7.00	±1.04	11	0.0013 (vs #57)
68	S2E	$tj^{TS} > E2f1-RNAi$	10 days	4.24	±0.36	29	6e <sup>-16</sup> (vs #58)
69	S2E	$ti^{TS} > E2f1-RNAi$	20 days	0.50	±0.33	22	2e <sup>-17</sup> (vs #59)
70	S2F	eyaA3 > +	10 days	11.25	±0.33	32	(***********************************
71	S2F	eyaA3 > Dp-RNAi	10 days	1.88	±0.33	43	7e <sup>-31</sup> (vs #70)
72	S2F	C587 > +	10 days	11.82	±0.27	35	(10 (10 1110)
73	S2F	C587 > Dp-RNAi	10 days	0.46	±0.17	52	9e <sup>-42</sup> (vs #72)
74	S2F	$fng^{TS} > +$	0 days	12.50	±0.17	12	30 (V3 #12)
75	S2F	$fng^{TS} > +$	10 days	12.57	±0.72	7	
76	S2F	$fng^{TS} > +$	20 days	12.62	±0.72	37	
77	S2F	fng <sup>TS</sup> > Dp-RNAi	0 days	11.93	±0.64	14	0.6416 (vs
78	S2F	$fng^{TS} > Dp-RNAi$	10 days	7.40	±0.32	9	#74) 0.0001 (vs #75)
79	S2F	$fng^{TS} > Dp-RNAi$	20 days	4.59	±0.60	32	9e <sup>-16</sup> (vs #76)
80	S2F	$fng^{TS} > E2f1-RNAi$	10 days	9.00	±0.42	16	0.0015 (vs #75)

81	S2F	fng <sup>⊤S</sup> > E2f1-RNAi	20 days	7.33	±0.68	12	3e <sup>-6</sup> (vs #76)
82	S3F	nos > +	0 days	11.70	±0.56	10	
83	S3F	nos > +	10 days	12.83	±1.28	12	
84	S3F	nos > Fs-RNAi	0 days	11.76	±0.61	17	0.9382 (vs #82)
85	S3F	nos > Fs-RNAi	10 days	12.25	±0.57	24	0.9793 (vs #83)
86	S3H	upd TS > +, Fs <sup>null</sup> / +	0 days	10.71	±0.41	21	,
87	S3H	upd TS > +, Fs <sup>null</sup> / +	2 weeks	10.20	±0.47	30	
88	S3H	upd TS > +, Fs <sup>null</sup> / +	4 weeks	9.40	±0.96	10	
89	S3H	upd TS > Fs, Fs <sup>null</sup> / +	0 days	10.24	±0.38	21	
90	S3H	$upd^{TS} > Fs$ , $Fs^{null} / +$	2 weeks	9.92	±0.58	12	
91	S3H	upd TS > Fs, Fs <sup>null</sup> / +	4 weeks	8.79	±0.35	19	
92	S3H	upd TS > +, Fs <sup>null -/-</sup>	0 days	9.71	±0.52	17	
93	S3H	upd TS > +, Fs <sup>null -/-</sup>	2 weeks	6.92	±0.48	38	
94	S3H	upd TS > +, Fs <sup>null -/-</sup>	4 weeks	6.20	±0.59	11	
95	S3H	upd TS > Fs, Fs <sup>null -/-</sup>	0 days	9.95	±0.60	19	
96	S3H	upd TS > Fs, Fs <sup>null -/-</sup>	2 weeks	9.71	±0.39	28	3e <sup>-5</sup> (vs #93)
97	S3H	upd TS > Fs, Fs <sup>null -/-</sup>	4 weeks	9.36	±0.75	19	0.0028 (vs #94)
98	S3I	fng > +, Fs <sup>null</sup> / +	4 weeks	9.14	±1.09	14	,
99	S3I	fng > Fs, Fs <sup>null</sup> / +	4 weeks	8.07	±0.89	14	
100	S3I	fng > +, Fs <sup>null -/-</sup>	4 weeks	5.00	±0.84	11	
101	S3I	fng > Fs, Fs <sup>null -/-</sup>	4 weeks	8.29	±0.53	24	0.0034 (vs #100)
102	S3J	Fs <sup>GFSTF.1</sup> / Fs <sup>MI11350</sup>	0 days	11.64	±0.48	36	,
103	S3J	Fs <sup>GFSTF.1</sup> / Fs <sup>MI11350</sup>	4 weeks	11.14	±0.30	37	0.3751 (vs #102)
104	S3J	Fs <sup>TG4.1</sup> / Fs <sup>MI</sup> -GAL4	0 days	11.39	±0.37	18	,
105	S3J	Fs <sup>TG4.1</sup> / Fs <sup>MI</sup> -GAL4	4 weeks	5.64	±0.67	39	6e <sup>-10</sup> (vs #104)
106	S3J	Fs <sup>GFSTF.1</sup> / Fs <sup>TG4.1</sup>	0 days	10.40	±0.46	25	,
107	S3J	Fs <sup>GFSTF.1</sup> / Fs <sup>TG4.1</sup>	4 weeks	7.88	±0.76	32	0.0067 (vs #106)
108	S3J	Fs <sup>GFSTF.1</sup> / Fs <sup>MI</sup> -GAL4	0 days	10.32	±0.58	19	,
109	S3J	Fs <sup>GFSTF.1</sup> / Fs <sup>MI</sup> -GAL4	4 weeks	9.55	±0.37	51	0.2744 (vs #108)
110	S3J	Fs <sup>MI</sup> -GAL4 / Fs <sup>MI11350</sup>	0 days	13.21	±0.40	39	,
111	S3J	Fs <sup>MI</sup> -GAL4 / Fs <sup>MI11350</sup>	4 weeks	11.48	±0.79	25	0.0576 (vs #110)
112	S3J	Fs <sup>TG4.1</sup> / Fs <sup>MI11350</sup>	0 days	11.47	±0.63	15	,
113	S3J	Fs <sup>TG4.1</sup> / Fs <sup>MI11350</sup>	4 weeks	3.41	±0.43	66	2e <sup>-11</sup> (vs #112)

Table S2: Genotypes, Related to Figures 1-5

Figure	Full genotype	Abbreviation used in Figure
1A,I	yw / Y; tj-GAL4 / +;	tj > +
1B,1	yw / Y; tj-GAL4 / UAS-Dp-RNAi (GD 12722);	tj > Dp-RNAi
1C,E,G	yw, hs-Flp, tub>GAL4, UAS-nlsGFP / Y; FRT <sup>42D</sup> tub>GAL80 / FRT <sup>42D</sup> ;	Control clones
1D,F,H	yw, hs-Flp, tub>GAL4, UAS-nlsGFP / Y; FRT <sup>42D</sup> tub>GAL80 /FRT <sup>42D</sup> Dp <sup>a3</sup> ;	<i>Dp</i> <sup>a3</sup> clones
11	yw / Y; tj-GAL4 / UAS-Dp-RNAi (GD 12722); UAS- GFP / +	tj > Dp-RNAi (GD), GFP
11	yw / Y; tj-GAL4 / +; UAS-Dp-RNAi (TRiP JF02519) / +	tj> Dp-RNAi (TRiP)
11	yw / Y; tj-GAL4 / UAS-Dp-RNAi (GD 12722); UAS- Dp / +	tj > Dp-RNAi (GD), Dp
11	yw / Y; tj-GAL4 / +: UAS-E2f1-RNAi / +	tj > E2f1-RNAi
11	yw / Y; tj-GAL4 / +: UAS-Rbf <sup>280</sup> / +	$tj > Rbf^{280}$
1J	yw / Y; tj-GAL4 / +: hh-GAL80 / +	tj, hh-GAL80 > +
1J	yw / Y; tj-GAL4 / UAS-Dp-RNAi (GD 12722); hh- GAL80 / UAS-LacZ	tj, hh-GAL80 > Dp-RNAi, LacZ
1J	yw / Y; tj-GAL4 / UAS-Dp-RNAi (GD 12722); hh- GAL80 / UAS-Dp	tj, hh-GAL80 > Dp-RNAi, Dp
2B,D	w / Y; tj-GAL4 / UAS-LacZ; UAS-LacZ / UAS-Dicer- 2	tj > LacZ, LacZ
2C,D	w / Y; tj-GAL4 / UAS-LacZ; UAS-Fs-RNAi / UAS- Dicer-2	tj > Fs-RNAi, LacZ
2D	w / Y; tj-GAL4 / UAS-Fs; UAS-Fs-RNAi / UAS-Dicer- 2	tj > Fs-RNAi, Fs
2D	w /Y; tj-GAL4 / UAS-Fs; UAS-LacZ / UAS-Dicer-2	tj > LacZ, Fs
2E,F	w /Y; Fs <sup>null</sup> / Fs <sup>null</sup>	Fs <sup>null</sup> -/-
2F	Oregon-R	wt
2G,I	w / Y; Fs-GAL4 / UAS-LacZ; UAS-GFP / +	Fs > GFP, LacZ
2H,I	w / Y; Fs-GAL4 / UAS-Dp-RNAi; UAS-GFP / +	Fs > GFP, Dp-RNAi
2J	yw / Y; tj-GAL4 / +; hh-GAL80 / +	tj, hh-GAL80 > +
2J	yw / Y; tj-GAL4 / UAS-Dp-RNAi (GD 12722); hh- GAL80 / UAS-LacZ	tj, hh-GAL80 > Dp-RNAi, LacZ
2J	yw/Y; tj-GAL4/UAS-Dp-RNAi (GD 12722); hh- GAL80 / UAS-Fs	tj, hh-GAL80 > Dp-RNAi, Fs
3A	w / Y; ; babo <sup>fTRG00444.sfGFP-TVPTBF</sup>	babo-GFP
3B	upd-GAL4, tub-GAL80™ / Y	upd <sup>TS</sup> > +
3B,C	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y;; UAS-babo <sup>QD</sup> / +	upd <sup>TS</sup> > babo <sup>QD</sup>
3D,F	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; ubiP63E-FRT-STOP- FRT-GFP, UAS-Flp / +; +	upd <sup>TS</sup> > lineage
3E,F	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; ubiP63E-FRT-STOP- FRT-GFP, UAS-Flp / +; UAS-babo <sup>QD</sup> / +	upd <sup>TS</sup> > babo <sup>QD</sup> , lineage
4B,D	C587-GAL4 / Y; QUAS-Flp / +; act>y[+]>LHV2- 86Fb, 13XlexAop2-myr::GFP / hh-QF	c587 > +, hh > lineage

4C,D	C587-GAL4 / Y; QUAS-Flp /UAS-Dp-RNAi (GD 12722); act>y[+]>LHV2-86Fb, 13XlexAop2- myr::GFP / hh-QF	c587 > Dp-RNAi, hh > lineage
4E	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; UAS-NLAP / + ; UAS- babo <sup>QD</sup> / +	NLAP (control)
4E	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; UAS-esg <sup>NLAP</sup> / + ; UAS-babo <sup>QD</sup> / +	esg <sup>NLAP</sup>
5A,B,C	w / Y; Fs-GAL4 / + ; UAS-GFP / +	Fs > GFP
5D	w / Y; tj-GAL4 / UAS-LacZ; UAS-Dicer-2 / +	tj-GAL4 > LacZ
5D	w / Y; tj-GAL4 / UAS-Fs; UAS-Dicer-2 / +	tj-GAL4 > Fs
5E	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; UAS-LacZ / +	upd <sup>TS</sup> > LacZ
5E	upd-GAL4, tub-GAL80 <sup>TS</sup> /Y; UAS-smox-RNAi/+	upd <sup>TS</sup> > smox-RNAi
5E	upd-GAL4, tub-GAL80 <sup>TS</sup> /Y; UAS-babo-RNAi/+	upd <sup>TS</sup> > babo-RNAi
5E	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; UAS-daw-RNAi <sup>HMS01110</sup> / +	upd <sup>TS</sup> > daw-RNAi- HMS01110
5E	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; UAS-daw-RNAi <sup>HMJ03135</sup> / +	upd <sup>™</sup> S > daw-RNAi- HMJ03135
5F	yw / Y; daw <sup>MI05383</sup>	daw-GFP
5G	w / Y; daw <sup>NP4661</sup> -GAL4 / + ; UAS-GFP.nls	daw <sup>NP4661</sup> > GFP
5H	Oregon-R	wt
S1A,B	yw, hs-Flp, tub-GAL4, UAS-nlsGFP / Y; FRT <sup>42D</sup> tub- GAL80 / FRT <sup>42D</sup> Dp <sup>a4</sup>	Dp <sup>a4</sup> clones
S1B	yw, hs-Flp, tub-GAL4, UAS-nlsGFP / Y; FRT <sup>42D</sup> tub- GAL80 / FRT <sup>42D</sup> ;	FRT <sup>42D</sup> (control)
S1B,C,E	yw, hs-Flp, tub-GAL4, UAS-nlsGFP / Y; FRT <sup>42D</sup> tub- GAL80 / FRT <sup>42D</sup> Dp <sup>a3</sup> ;	Dp <sup>a3</sup> clones
S1D	yw, hs-Flp, tub-GAL4, UAS-nlsGFP / Y; UAS-Dp-RNAi (GD 12722) / +; FRT <sup>82B</sup> tub-GAL80 / FRT <sup>82B</sup>	Dp-RNAi clones
S1F	yw, hs-Flp, tub-GAL4, UAS-nlsGFP / Y; ; FRT <sup>82B</sup> tub-GAL80 / FRT <sup>82B</sup> E2f1 <sup>729</sup>	E2f1 clones
S1G	yw, hs-Flp, tub-GAL4, UAS-nlsGFP / Y; ; FRT <sup>82B</sup> tub-GAL80 / FRT <sup>82B</sup>	FRT <sup>82B</sup> (control)
S2A	yw / Y; tj-GAL4 / +; hh <sup>P30</sup> / +	tj > +; hh-LacZ
S2B	upd <sup>PD1</sup> / Y; tj-GAL4 / +;	tj > +; upd-LacZ
S2C	yw / Y; tj-GAL4 / UAS-Dp-RNAi (GD 12722); hh <sup>p30</sup> / +	tj > Dp-RNAi; hh-LacZ
S2D	upd <sup>PD1</sup> / Y; tj-GAL4 / UAS-Dp-RNAi (GD 12722)	tj > Dp-RNAi; upd-LacZ
S2E	yw / Y; tj-GAL4 / +; tub-GAL80 <sup>TS</sup> / +	$tj^{TS} > +$
S2E	yw / Y; tj-GAL4 / UAS-Dp-RNAi (GD 12722); tub- GAL80 <sup>TS</sup> / +	$tj^{TS} > Dp$ -RNAi
S2E	yw / Y; tj-GAL4 / +; tub-GAL80 <sup>TS</sup> / UAS-E2f1-RNAi	$tj^{TS} > E2f1-RNAi$
S2F	yw, hs-Flp / Y; eyaA3-GAL4, M5-4-LacZ / +;	eyaA3 > +
S2F	yw, hs-Flp / Y; eyaA3-GAL4, M5-4-LacZ / UAS-Dp- RNAi (GD 12722);	eyaA3 > Dp-RNAi
S2F	C587-GAL4 / Y; ; Chd64-GFP / +	C587 > +
S2F	C587-GAL4 / Y; UAS-Dp-RNAi (GD 12722) / +; Chd64-GFP / +	C587 > Dp-RNAi
S2F	yw, hs-Flp / Y; tub-GAL80 <sup>TS</sup> / +; fng-GAL4 / +	$fng^{TS} > +$
S2F	yw, hs-Flp / Y; tub-GAL80 $^{TS}$ / UAS-Dp-RNAi (GD 12722); fng-GAL4 / +	$fng^{TS} > Dp-RNAi$

S2F	yw, hs-Flp / Y; tub-GAL80 <sup>™</sup> / +; fng-GAL4 / UAS- E2f1 RNAi	fng <sup>™</sup> > E2f1-RNAi
S3B,C,E	w / Y; tj-GAL4 / + ; UAS-Fs-RNAi / UAS-Dicer-2	tj > Fs-RNAi
S3E	w / Y; tj-GAL4 / + ; UAS-Dicer-2 / +	<i>tj</i> > +
S3E	Oregon-R	wt
S3E,G	w / Y; Fs <sup>null</sup> / Fs <sup>null</sup>	Fs <sup>null -/-</sup>
S3F	w / Y; nos-GAL4-VP16 / + ; +	nos > +
S3F	w / Y; nos-GAL4-VP16 / + ; UAS-Fs-RNAi / +	nos > Fs-RNAi
S3H	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; Fs <sup>null</sup> / + ; +	upd <sup>TS</sup> > +, Fs <sup>null</sup> / +
S3H	upd-GAL4, tub-GAL80 <sup>TS</sup> /Y; Fs <sup>null</sup> /+; UAS-Fs/+	upd <sup>TS</sup> > Fs, Fs <sup>null</sup> / +
S3H	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; Fs <sup>null</sup> / Fs <sup>null</sup> ; +	upd <sup>TS</sup> > +, Fs <sup>null -/-</sup>
S3H	upd-GAL4, tub-GAL80 <sup>TS</sup> / Y; Fs <sup>null</sup> / Fs <sup>null</sup> ; UAS-Fs /	upd <sup>TS</sup> > Fs, Fs <sup>null -/-</sup>
S3I	w / Y; Fs <sup>null</sup> / + ; fng-GAL4 / +	fng > +, Fs <sup>null</sup> / +
S3I	w / Y; Fs <sup>null</sup> / + ; fng-GAL4 / UAS-Fs	fng > Fs, Fs <sup>null</sup> / +
S3I	w/ Y; Fs <sup>null</sup> / Fs <sup>null</sup> ; fng-GAL4 / +	fng > +, Fs <sup>null -/-</sup>
S3I	w / Y; Fs <sup>null</sup> / Fs <sup>null</sup> ; fng-GAL4 / UAS-Fs	fng > Fs, Fs <sup>null -/-</sup>
S3J	w / Y; Fs <sup>GFSTF.1</sup> / Fs <sup>MI11350</sup>	Fs <sup>GFSTF.1</sup> / Fs <sup>MI11350</sup>
S3J	w / Y; Fs <sup>TG4.1</sup> / Fs <sup>MI</sup> -GAL4	Fs <sup>TG4.1</sup> / Fs <sup>MI</sup> -GAL4
S3J	$W/Y$ ; $Fs^{GFSTF.1}/Fs^{TG4.1}$	Fs <sup>GFSTF.1</sup> / Fs <sup>TG4.1</sup>
S3J	w / Y; Fs <sup>GFSTF.1</sup> / Fs <sup>MI</sup> -GAL4	Fs <sup>GFSTF.1</sup> / Fs <sup>MI</sup> -GAL4
S3J	w / Y; Fs <sup>MI</sup> -GAL4 / Fs <sup>MI11350</sup>	Fs <sup>MI</sup> -GAL4 / Fs <sup>MI11350</sup>
S3J	w / Y; Fs <sup>TG4.1</sup> / Fs <sup>MI11350</sup>	Fs <sup>TG4.1</sup> / Fs <sup>MI11350</sup>
S4A,C	yw / Y; tj-GAL4/Fs <sup>MI04308-GFSTF.1</sup> (Fs-GFP);	<i>tj</i> > +
S4B,C	yw / Y; tj-GAL4/ Fs <sup>MI04308-GFSTF.1 (Fs-GFP)</sup> ; UAS-E2f1- RNAi / +	tj > E2f1-RNAi
S5B,C	C587-GAL4 / Y; QUAS-Flp / +; ubiP63E-FRT-	+ (in B)
	STOP-FRT-GFP / hh-QF	C587 > +, hh > lineage
		(GTRACE) (in C)
S5B,D	C587-GAL4 / Y; QUAS-Flp / UAS-Dp-RNAi (GD	<i>Dp-RNAi</i> (in b)
	12722); ubiP63E-FRT-STOP-FRT-GFP / hh-QF	C587 > Dp-RNAi, hh >
		lineage (GTRACE) (in D)
S5E	C587-GAL4 / Y; QUAS-Flp / +; act>y[+]>LHV2-	C587 > +, hh > lineage
	86Fb, 13xlexAop2-myr::GFP / hh-QF	(FlexAmp)
S5F,G	C587-GAL4 / Y; QUAS-Flp / UAS-Dp-RNAi (GD	C587 > Dp-RNAi, hh >
	12722); act>y[+]>LHV2-86Fb, 13xlexAop2-	lineage (FlexAmp)
	myr::GFP / hh-QF	
S6A	w / Y; daw <sup>NP6274</sup> -GAL4, UAS-LacZ / + ; UAS-GFPnIs	daw <sup>NP6274</sup> > GFP
S6B	w / Y; ; babo <sup>fTRG00444.sfGFP-TVPTBF</sup>	babo-GFP
S6C	w / Y; ; Actβ-GAL4 / UAS-GFPnls	Actβ-GAL4 > GFP
S6D	W / Y; ; Actβ <sup>fTRG00506.sfGFP-TVPTBF</sup>	Actβ-GFP
S6E	w / Y; myo-GAL4 / UAS-GFPnls	myo-GAL4 > GFP
S6F	w / Y; ; myofTRG00161.sfGFP-TVPTBF	myo-GFP
S6G	w / Y; ; put <sup>fTRG00910.sfGFP-TVPTBF</sup>	put-GFP

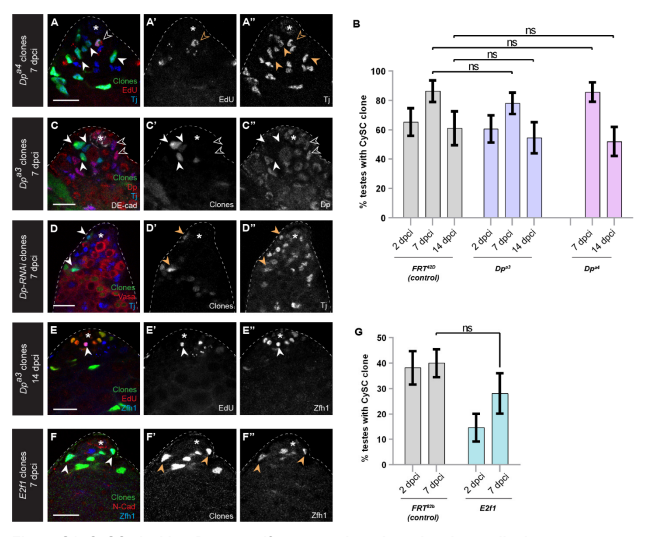


Figure S1: CySCs lacking *Dp* can self-renew and produce daughter cells that differentiate, Related to Fig. 1

- (A) CySC clones (green) mutant for  $Dp^{a4}$  can remain in the niche for at least 7 days after clone induction (dpci). Arrowheads mark the CySC clones and the open arrowhead a CySC clone positive for EdU (red, A'). Tj (blue, A") labels CySCs and early cyst cells.
- (**B**) Graph indicating the clone recovery rates at 2, 7 and 14 dpci for control clones (labeled  $FRT^{42D}$ , gray bars, n=26, 22, 18, respectively),  $Dp^{a3}$  mutant clones (blue bars, n=28, 32, 22, respectively), and  $Dp^{a4}$  mutant clones (purple bars, n=28, 25 respectively). ns = not significant, assessed by Fisher's exact test. Error bars represent SEM.
- (**C**) A *Dp*<sup>a3</sup> mutant clone (green, C') at 7 dpci, labeled with an antibody against Dp protein (red, C"). Mutant CySCs (closed arrowheads) lack detectable Dp. Neighboring wild type CySCs, marked with open arrowheads, express Dp protein (C").

- (**D**) CySC clones expressing the same RNAi construct targeting Dp as in Fig. 1 (arrowheads) can be recovered at 7 dpci. GFP (green) marks the clones. Tj (blue) marks CySCs and early cyst cell. Vasa (red) labels the germline.
- (E) A *Dp*<sup>a3</sup> mutant clone (green) at 14 dpci labeled with EdU. The clone contains several Zfh1-positive (blue, D"), EdU-positive (red, D') CySCs (arrowhead shows one such CySC).
- (**F**) CySC clones (green) mutant for *E2f1* can be recovered after 7 days. (The arrowheads show two such CySCs). Zfh1 (blue) marks CySCs and early cyst cells. N-Cad (red) marks the hub.
- (**G**) Graph indicating the clone recovery rates at 2 and 7 dpci for control clones (labeled  $FRT^{82B}$ , gray bars, n=35 and 80, respectively) and E2f1 mutant clones (blue bars, n=41 and 32, respectively). ns = not significant, assessed by Fisher's exact test. Error bars represent SEM. The hub is shown with an asterisk in A, C, D, E, F. Scale bar = 20  $\mu$ M.

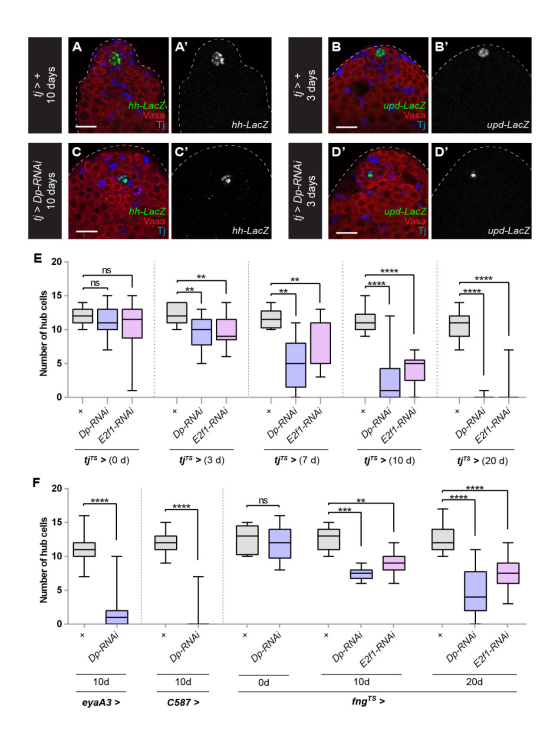


Figure S2: Hub cells are lost when Dp/E2f1 complex activity is depleted from CySCs, Related to Fig. 1.

(**A-D**) Hub cells in control tj > + testes (A,B) express hh-LacZ and upd-LacZ, but the number of cells expressing these markers is greatly reduced in tj > Dp-RNAi testes (C,D) after 10 and 3 days at 29°C, respectively.

- (**E**) Graph of the number of hub cells in  $tj^{TS}$  > + (gray bars, n=17, 21, 8, 35, 17 at 0, 3, 7, 10, 20 days, respectively),  $tj^{TS}$  > Dp-RNAi (purple bars, n=28, 14, 6, 39, 30 at 0, 3, 7, 10, 20 days, respectively);  $tj^{TS}$  > E2f1-RNAi (pink bars, n=18, 17, 11, 29, 22 at 0, 3, 7, 10, 20 days, respectively). Flies were reared at 18°C until eclosion and then reared at 29°C for the indicated time periods.
- (**F**) Graph of the number of hub cells in control testes (gray bars, n=32 for eyaA3 > +, n=35 for C587 > +, n=12, 7, and 37 for  $fng^{TS} > +$  at 0, 10 and 20 days, respectively), or in testes with knockdown of Dp (purple bars, n=43 for eyaA3 > Dp-RNAi, n=52 for C857 > Dp-RNAi, n=14, 9, 32 for  $fng^{TS} > Dp$ -RNAi at 0, 10, and 20 days, respectively) or E2f1-RNAi (pink bars, n=16 and 12 for  $fng^{TS} > E2f1$ -RNAi at 10 and 20 days, respectively) in eya >, C587 >, or  $fng^{TS} >$  at the indicated time periods.

Error bars represent the data range. \*\*\*\* P < 0.0001; \*\*\* P < 0.001; \*\* P < 0.01; ns = not significant, as assessed by Student's t-test. See Supplementary Table 1 for exact P values. Scale bar = 10  $\mu$ M.

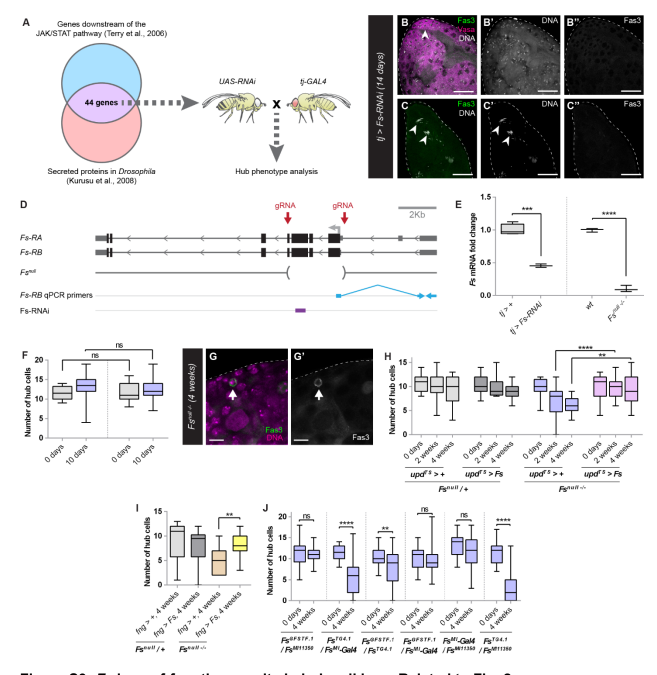


Figure S3: Fs loss-of-function results in hub cell loss, Related to Fig. 2.

- (A) Schematic of the screen performed to identify secreted factors from CySCs influencing hub fate. Briefly, we cross-referenced genes enriched in testis stem cells with the list of *Drosophila* secreted proteins. We obtained flies carrying RNAi lines targeting the resulting 44 genes and crossed them to a *tj-GAL4* line, and scored for phenotypes on hub cells.
- (**B**, **C**) Representative confocal images of testes from tj > Fs-RNAi at 2 weeks of adulthood. Note the absence of Fas3-positive hub cells in both images. Arrowhead in (B) indicates a

- differentiating 16-cell spermatogonia. Arrowhead in (C) indicates canoe stage spermatids that normally are located basally. Due to the absence of any mitotic cells, these differentiated cells are observed more apically. Vasa (magenta) labels germ cells in (B); Fas3 (green) marks hub cells and Topro (white) marks DNA.
- (**D**) Model of the *Fs* genomic locus and the two predicted *Fs* isoforms (*Fs-RA* and *Fs-RB*). The first line below the genomic locus shows the approximate extent of the deletion spanning coding exons 1-4 in the *Fs<sup>null</sup>* allele created with gDNAs in the positions shown with red arrows. The location of the primers used for qPCR and the sequence targeted by the RNAi is shown below. Gray boxes represent non-coding exons and black boxes the coding exons.
- (**E**) Left panel: expression of Fs mRNA in control tj > + (gray bar) and tj > Fs-RNAi (blue bar) as measured by qRT-PCR (n=4). Right panel: expression of Fs mRNA in wild (wt) type testes (gray bar) and  $Fs^{null}$  mutant testes (blue bar) as measured by qRT-PCR (n=4).
- (**F**) Number of hub cells in control *nos* > + (gray bar, n=10 and 12 at 0 and 10 days, respectively) *and nos* > *Fs-RNAi* (blue bar, n=12 and 17 at 0 and 10 days, respectively) at 0 and 10 days of adulthood.
- (**G**) Representative confocal image of a *Fs*<sup>null</sup> testis at 4 weeks of adulthood. Fas3 (green) staining reveals only one hub cell (arrow) remaining in these testes. Topro (magenta) marks DNA.
- (**H**) Graph of number of hub cells at different time points of adulthood in testes carrying the hub cell driver upd-GAL4 in an heterozygous  $Fs^{null}/+$  (light gray bars) or homozygous  $Fs^{null}/Fs^{null}$  (blue bars) background; or in testes in which upd-GAL4 drives UAS-Fs in an heterozygous  $Fs^{null}/+$  (dark gray bars) or homozygous  $Fs^{null}/Fs^{null}$  (purple bars) background. See Supplementary Table 1 for n values.
- (I) Graph of number of hub cells at different time points of adulthood in testes carrying the somatic cell driver fng-GAL4 in a heterozygous  $Fs^{null}$ /+ (light gray bar, n=14) or homozygous  $Fs^{null}$ / $Fs^{null}$  (dark gray bar, n=11) background; or in testes in which fng-GAL4 drives UAS-Fs in a heterozygous  $Fs^{null}$ /+ (brown bar, n=14) or homozygous  $Fs^{null}$ / $Fs^{null}$  (yellow bar, n=24) background.
- (J) Graph of number of hub cells at 0 and 4 weeks of adulthood in heteroalleic combinations of the indicated Fs hypomorphic mutant alleles. See Supplementary Table 1 for n values. Error bars represent the data range. \*\*\*\* P < 0.0001; \*\*\* P < 0.001; \*\* P < 0.01; ns = not significant, assessed by Student's t-test. Scale bar = 50  $\mu$ M.

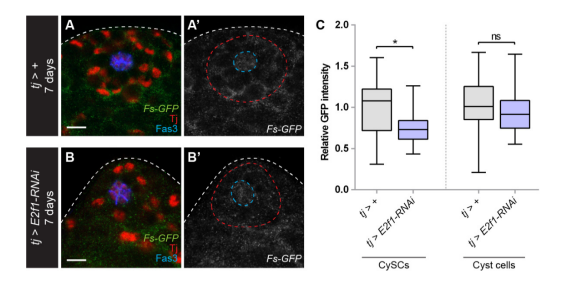


Figure S4: Depletion of *E2f1* from the cyst lineage causes a significant decrease in Fs protein in the testis stem cell microenvironment, Related to Fig. 2.

(**A, B**) Representative confocal images of the expression pattern of a Fs protein trap (Fs<sup>GFP</sup> in green) in control tj > + testis (A) and a tj > E2f1-RNAi testis in which E2f1 was depleted from the cyst cell lineage for 7 days (B). Tj (red) marks CySCs and early cyst cells and Fas3 (blue) marks hub cells. Dashed blue line indicates cluster of hub cells. Red dashed line indicates CySCs within 1 cell diameter of the hub.

(**C**) Graph of relative GFP intensity in tj > + testes (gray bars, n=15) or tj > E2f1-RNAi testes (blue bars, n=23) of CySCs (i.e., the area between the dashed blue and red lines in A,B) and in cyst cells (i.e., the area basal to the red dashed line in A,B).

Error bars represent the data range. \* P < 0.05; ns = not significant, assessed by Student's t-test.

Scale bar =  $10 \mu M$ .

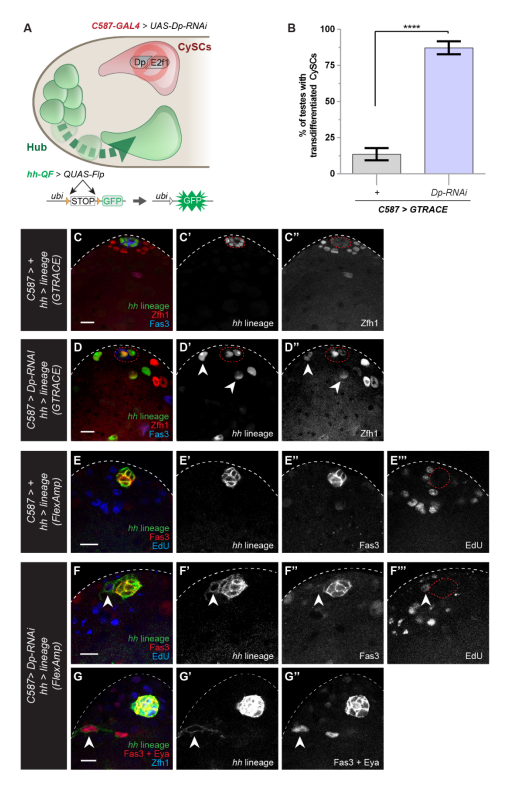


Figure S5: Depletion of *Dp* from the cyst lineage causes transdifferentiation of hub cells into CySCs, Related to Fig 4.

(A) Model indicating the alternative experimental design (referred to as "hh > lineage (GTRACE)") to deplete Dp from CySCs while concomitantly lineage tracing hub cells using

independent binary expression systems, complementary to the design shown in Figure 4A (referred to as "hh > lineage (FlexAmp)"). Experimental data are shown in (B). To trace the lineage of hub cells, we used *hh-QF*, which is expressed in only hub cells, to induce *QUAS-FLP*. In turn, FLP recombines *FRT* sites in the *ubi>STOP>GFP* GTRACE transgene. This leads to the labeling of hub cells and their lineage with nuclear GFP. At the same time, *C587-GAL4* drives expression of a *Dp-RNAi* transgene, which depletes Dp from CySCs but not from hub cells.

- (**B**) Graph showing the percentage of testes with hub lineage-positive CySCs in *C587* > + (gray bar, n=67) and *C587* > *Dp-RNAi* (blue bar, n=48) testes, using the experimental design indicated in (A).
- **(C, D)** Example images of *C587* > + (C) and *C587* > *Dp-RNAi* (D) testes, using the experimental design indicated in (A). Only in the latter genotype can GFP (green)-positive cells from the hub lineage (D, arrowheads) be found outside the hub. Zfh1 (red) marks CySCs; Fas3 (blue) marks hub cells. A red dashed line outlines the hub.
- (**E-G**) There are GFP (green)-positive cells lacking the hub cell marker Fas3 (red) (F-F", arrowhead) located outside the cluster of hub cells in a C587 > Dp-RNAi; hh > lineage (FLEXAMP) testis but not in a control C587 > +; hh > lineage (FLEXAMP) testis (E), following the experimental design indicated in **Fig. 4A**. These GFP-positive cells also incorporate EdU, showing that they undergo DNA replication (F", arrowhead) and differentiate into cyst cells, which express Eya and have thin, long membranes wrapping differentiating germ cell cysts (G-G", arrowhead). A red dashed line outlines the hub. The hh lineage is green in (E-G). EdU is blue and Fas3 is red in (E,G). Fas and Eya are red and Zfh1 is blue in (G). Error bars represent SEM. \*\*\*\* P < 0.0001, assessed by Fisher's exact test. Scale bar = 10 μM.

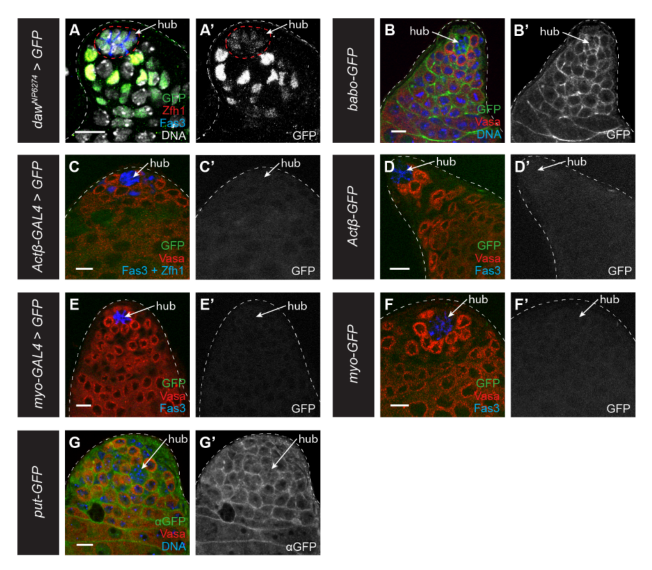


Figure S6: Expression pattern of Activin ligands and receptors in the *Drosophila* testis, Related to Fig. 5.

- **(A)** A *daw-GAL4* enhancer trap driving *UAS-GFP* (GFP). The *GAL4* insertions drives GFP expression in Fas3-positive hub cells and Zfh1-positive CySCs. GFP is green; Fas3 is blue; Zfh1 is red; DAPI (white) marks DNA.
- **(B)** Babo, as assessed by a fosmid *babo-GFP* transgene, is present at the surface of hub cells, CySCs and cyst cells and early germ cells. GFP is green; Vasa is red and Topro, which marks DNA, is blue.
- **(C)** *Actβ-GAL4* driving expression of *UAS-GFP*. GFP is not detected in any cells of the testis stem cell niche. GFP is green; Vasa is red; Fas3 and Zfh1 are both blue.

- (**D**) Act $\beta$ , as assessed by a fosmid *Act\beta-GFP* transgene, is not detected in any cells in the testis stem cell niche. GFP is green; Vasa is red; and Fas3 is blue.
- **(E)** *myo-GAL4* driving expression of *UAS-GFP*. GFP is not detected in any cells of the testis stem cell niche. GFP is green; Vasa is red; Fas3 is blue.
- **(F)** Myo, as assessed by a fosmid *myo-GFP* transgene, is not detected in any cells in the testis stem cell niche. GFP is green; Vasa is red; and Fas3 is blue.
- **(G)** The type II receptor Put is not detected in hub cells in the testis stem cell niche, as assessed by a fosmid *put-GFP* transgene. GFP is green; Vasa is red; and Fas3 is blue. Scale bar =  $10 \, \mu M$ .