Coupling of Hedgehog and Hippo pathways promotes stem cell maintenance by stimulating proliferation

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It is essential to define the mechanisms by which external signals regulate adult stem cell numbers, stem cell maintenance, and stem cell proliferation to guide regenerative stem cell therapies and to understand better how cancers originate in stem cells. In this paper, we show that Hedgehog (Hh) signaling in Drosophila melanogaster ovarian follicle stem cells (FSCs) induces the activity of Yorkie (Yki), the transcriptional coactivator of the Hippo pathway, by inducing yki transcription. Moreover, both Hh signaling and Yki positively regulate the rate of FSC proliferation, both are essential for FSC maintenance, and both promote increased FSC longevity and FSC duplication when in excess. We also found that responses to activated Yki depend on Cyclin E induction while responses to excess Hh signaling depend on Yki induction, and excess Yki can compensate for defective Hh signaling. These causal connections provide the most rigorous evidence to date that a niche signal can promote stem cell maintenance principally by stimulating stem cell proliferation.

Introduction

Self-renewal of adult stem cells is generally supported by a specialized niche microenvironment, which can serve to maintain stem cells in an appropriate location, in an undifferentiated state or in an appropriate proliferative state. For example, a short-range bone morphogenetic protein (BMP) signal produced in directly adjacent cap cells (Fig. 1 A) promotes germline stem cell (GSC) maintenance in the Drosophila melanogaster ovary by repressing a key differentiation factor (Chen et al., 2011b). BMP similarly represses differentiation in the male GSC lineage, whereas a Janus kinase–signal transducer and activator of transcription (JAK-STAT) pathway ligand produced in directly adjacent hub cells represses differentiation of somatic stem cells in the Drosophila testis (Losick et al., 2011; Matunis et al., 2012). These and other findings firmly established repression of differentiation as a, likely widespread, mechanism for niche signals to maintain stem cells (Losick et al., 2011). Other potential mechanisms for niche signals to support stem cells have not been fully explored and validated.

Although stem cells must proliferate to serve their regenerative function, relative quiescence was once thought to be universally important for long-term maintenance of stem cells. Several types of stem cells, including mammalian hematopoietic stem cells (HSCs) and muscle stem cells, indeed divide infrequently, whereas normal cycles of hair growth depend on maintaining quiescence of hair follicle stem cells (FSCs) in between brief periods of activation (Pietras et al., 2011; Chakkalakal et al., 2012; Hsu and Fuchs, 2012). However, it is now clear that some adult stem cells, including mammalian epidermal and gut stem cells proliferate constitutively and extensively (Simons and Clevers, 2011; Hsu and Fuchs, 2012). In all of these cases, there are likely specific niche signals, some of which have been identified, that stimulate or inhibit stem cell proliferation to produce an appropriate supply of tissue-replenishing daughters. However, an important question that has been hard to resolve is whether regulation of the rate of stem cell proliferation by niche factors is also a major mechanism for regulating the maintenance of the stem cells themselves.

Drosophila FSCs provide a particularly informative stem cell paradigm for understanding niche function because FSC maintenance depends on multiple known extracellular signals.
FSCs are constitutively active in well-fed flies, supporting the production of up to two egg chambers per day from each germarium (Margolis and Spradling, 1995), and are therefore a good model for highly proliferative mammalian stem cells. We had previously suggested that FSC proliferation promotes FSC maintenance based on artificial genetic conditions: several homozygous mutations expected to decrease cell proliferation, including partial loss of function cyclin E (cycE) mutations, caused cell-autonomous loss of FSCs, whereas excess CycE compensated for the loss of several otherwise essential FSC factors (Wang and Kalderon, 2009; Wang et al., 2012). We therefore sought to understand how proliferation is regulated in FSCs and whether regulation of the rate of FSC proliferation by niche signals affects stem cell maintenance or stem cell numbers.

Here, we show that Hh signaling induces the Hippo (Hpo) pathway coactivator Yorkie (Yki) in FSCs and that Yki, in turn, regulates FSC maintenance and expansion by stimulating FSC proliferation via induction of CycE and by preventing apoptosis. These causal connections provide clear evidence that a niche signal can regulate stem cell maintenance by controlling the rate of stem cell proliferation.

and FSCs compete for niche occupation (Nystul and Spradling, 2007, 2010; Vied et al., 2012). FSCs are maintained midway along the anterior–posterior (AP) axis of the germarium, at the region 2a/2b border; here, rounded stage 2a germline cysts shorten along the AP axis to form lens-shaped stage 2b cysts that span the width of the germarium for the first time (Fig. 1 A; Margolis and Spradling, 1995). FSCs self-renew and produce follicle cell (FC) daughters, which proliferate as they form a monolayer epithelium around passing germline cysts and also become specialized stalk cells that separate egg chambers as they bud from the posterior of the gerarium (Fig. 1 A; Nystul and Spradling, 2007, 2010; Wu et al., 2008b). FSC maintenance depends on Wnt and BMP niche signals, but the Hedgehog (Hh) and JAK-STAT pathways are the most potent regulators of FSC maintenance and expansion (Song and Xie, 2003; Kirilly et al., 2005; Vied et al., 2012). Hh and the JAK-STAT pathway ligand, Unpaired, derive from opposite ends of the gerarium, producing reciprocal AP gradients that overlap in region 2a/b to induce sufficiently high activity in each pathway to support FSCs (Vied et al., 2012). How those signals act within FSCs to promote their maintenance or duplication has not yet been determined.

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Results

Yki is required for FSC maintenance

The Hpo pathway has recently emerged as a major regulator of organ size and of the growth, cycling, and survival of individual cells in Drosophila and mammals, motivating exploration of its potential roles governing stem cell proliferation (Pan, 2010; Zhao et al., 2011; Ramos and Camargo, 2012). To investigate Hpo pathway function in FSCs, we induced marked homozygous mutant FSCs at a fixed time by using a heat shock–inducible flp recombinase and FRT recombination targets located at the base of the appropriate chromosome arm. We then scored the percentage of ovarioles that retained marked (GFP negative) FSC and FC descendants (an FSC clone; Fig. 1, A and D) over time. The earliest time interval used in this well-established assay is generally 7 d (Vied and Kalderon, 2009; Wang and Kalderon, 2009; Vied et al., 2012; Wang et al., 2012) to ensure that “all-marked” cells originated from an FSC. Derivatives of FRT-mediated recombination in FCs (transient FC clones) normally pass through the ovariole in 5 or less (Margolis and Spradling, 1995). FSC clones lacking activity of Yki, the key regulated transcriptional coactivator of the Hpo pathway, were lost very rapidly relative to controls when clones were induced in adults or in larvae (shortly before FSC establishment; Fig. 1, B–E; Vied et al., 2012).

We examined earlier time points by focusing only on the germarium, which should not retain transient FC clones beyond 3 d. At 4 d after clone induction, yki mutant FSCs were present in 13% of germaria compared with 71% for controls. Moreover, only 5% of germaria contained marked wild-type FCs but no FSC, whereas 30% of germaria contained yki mutant FCs but no FSC, directly revealing accelerated yki mutant FSC loss (Fig. S1 A).

In contrast, yki mutant GSCs, which also reside on the germarium and can be assayed in parallel by exactly analogous methodology, were maintained normally over 18 d (Fig. 1, B and E). A selective requirement in FSCs but not GSCs has been observed previously for several other regulators of FSC behavior (Wang et al., 2012).

Most FSC daughters divide about eight times over a 3–4-d period before differentiating as mature FCs and dying 2–3 d later (Margolis and Spradling, 1995). The proliferation of these FCs is not impaired by partial loss-of-function cycE mutations or by several other mutations in FSC-selective factors that drastically reduce FSC persistence (Wang and Kalderon, 2009; Wang et al., 2012). However, yki mutant clones (marked by loss of GFP) induced in FCs and captured five or fewer days after induction were much smaller than the “twin spot” wild-type clones (marked by two copies of the ubiquitin [ubi]-GFP gene), which are generated simultaneously (Fig. 2, A–C). Yki commonly regulates both cell cycling and apoptosis. Apoptosis was detected in yki mutant FCs by activated Capase-3 staining in >15% of yki mutant FSC clones 4 d after induction compared with <2% of control FSC clones (Fig. S1, B–D). We also found that expression of a DIAP1 transgene, which inhibits apoptosis, in (GFP positive) yki mutant FC clones made by the mosaic analysis with a repressible cell marker (MARCAM) method (Lee and Luo, 2001), restored their size to normal; in contrast, expressing excess CycE did not alter yki mutant FC clone size (Fig. 2, D–H). Thus, net expansion of yki mutant FC clones is limited principally by cell death.

Coexpression of DIAP1 in MARCM clones did not, however, prevent the rapid loss of yki mutant FSCs (Fig. 1, C and F). Excess CycE was similarly ineffective. However, excess CycE and DIAP1 together substantially rescued yki FSC maintenance (Fig. 1, C and G). Hence, yki has an important antiapoptotic function throughout the FSC lineage, but it also has a selective function in FSCs connected with stimulating cell cycling that is not critical for FCs or indeed for GSCs.

FSCs with increased Yki activity compete better

In Drosophila and mammals, upstream components of the Hpo pathway regulate the specific activity of Yki proteins. A major mechanism involves regulation of the nuclear entry and activity of Yki orthologues by phosphorylation at conserved sites, including S168 in Drosophila Yki (Pan, 2010; Staley and Irvine, 2012). The protein kinase directly responsible, Warts (Wts), is itself phosphorylated and activated by the Hpo protein kinase in a complex with the scaffolding protein Salvador (Sav). Hence, loss of hpo, sav, or wts generally activates Yki protein. We found that hpo, wts, and sav mutant FSC clones produced striking overrepresentation phenotypes, indicating enhanced FSC activity (Fig. 3, A and C; and Fig. S2 B). Each germarium contains more than one FSC. Consequently, marked wild-type FSC clones are almost always accompanied by unmarked FSC derivatives in mosaic ovarioles (Fig. 1, A and D). However, a significant proportion of hpo, wts, and sav mutant FSC clones occupied the entire ovariole (all-marked clones; Fig. 3, A and C; and Fig. S2 B), as seen previously for genetic changes (patched [ptc] mutation) that increase Hh (Fig. 3 D), JAK-STAT, or phosphatidylinositol 3'-kinase pathway activities (Vied et al., 2012; Wang et al., 2012). Production of an all-marked clone from a single FSC implies that the marked FSC duplicated and out-competed unmarked FSCs in the same germarium (NYstul and Spradling, 2007; Vied et al., 2012). The frequency of all-marked clones continued to increase over time (36% for hpo with 28% mosaic and 39% for ptc with 34% mosaic at 26 d after clone induction in adults), as might be expected for FSCs with a stable competitive advantage. hpo, wts, and sav mutant FSC clones were also maintained better than wild-type FSC clones (Fig. 3 A). The frequency of all-marked ovarioles for hpo, wts, and sav was significantly higher for FSC clones induced in larvae rather than in adults (Fig. 3 A), as observed previously for excess Hh, phosphatidylinositol 3'-kinase, or JAK-STAT pathway activity, consistent with the idea that competition among potential stem cells is particularly intense during FSC establishment (Vied et al., 2012; Wang et al., 2012).

The all-marked hpo, wts, and sav FSC clones also produced an excessive number of FCs, which accumulated between egg chambers or formed multilayered epithelia (Fig. 3 C and Fig. S2 B). Those phenotypes are reminiscent of ptc mutant FSC clones (Fig. 3 D), which have elevated Hh pathway activity, and differ from the phenotypes produced by FSCs with increased Yorkie relays Hedgehog stem cell niche signal • Huang and Kalderon
phosphatidyl inositol 3'-kinase or JAK-STAT pathway activity (Vied et al., 2012; Wang et al., 2012).

FSC clones lacking both Hpo and Yki activity were lost just as rapidly as yki mutant FSCs (Fig. 3 A), consistent with the expectation that hpo mutant phenotypes are induced entirely by excessive activation of Yki. Indeed, phenotypes analogous to those of hpo, wts, and sav were produced by expression of Yki with an activating S168A alteration in GFP-positive MARCM FSC clones (Fig. 4 D and Fig. S2, F and G). Thus, FSCs with elevated Yki activity outcompete wild-type FSCs for retention in the gerarium and become overrepresented.

Merlin (Mer), Expanded (Ex), and Kibra are key upstream regulators of the Hpo pathway in FSCs

Kibra can associate with the cortical FERM (4.1 protein, Ezrin, Radixin, and Moesin) domain proteins Ex and Mer to connect external stimuli to activation of core Hpo pathway components (Boggiano and Fehon, 2012; Staley and Irvine, 2012). Both ex and mer mutant FSC clones survived better than wild-type clones and produced a significant frequency of all-marked ovarioles (Fig. 4 D and Fig. S2, F and G). Thus, FSCs with elevated Yki activity outcompete wild-type FSCs for retention in the gerarium and become overrepresented.

Hpo-Yki pathway regulates FSCs by altering the rate of FSC proliferation

We then wished to test whether the Hpo pathway regulates FSC maintenance and FSC expansion by influencing cell proliferation, as suggested by our earlier finding that excess CycE

![Figure 2. Yki is important for net FC amplification principally by preventing apoptosis.](image-url)
We then tested whether the regulation of FSC maintenance and competition by the Hpo pathway depends on CycE, which is commonly induced by Yki to stimulate cell cycling (Pan, 2010). Here, we used ex mutations to alter Hpo pathway activity because ex and cycE are on the same chromosome arm, facilitating testing of double mutants. These tests used both a null allele of cycE (cycE_{AR95}) and an allele (cycE_{WX}) that retains enough activity to support FC proliferation (Wang et al., 2012). Although ex mutant FSCs induced in larvae normally survive better than control FSC clones and produce an elevated frequency of all-marked clones (Fig. 4, D and E), cycE_{ex} double mutant FSCs were instead lost prematurely (Fig. 4 D). Similarly, expression of a constitutively active form of Yki (UAS-Yki^{S168A}) alone produced a strong all-marked FSC clone phenotype but failed even to rescue FSC maintenance in cycE mutant FSCs (Fig. 4 D). FSC maintenance can be rescued for both cycE alleles by constitutive expression of UAS-Yki^{S168A} (driven by constitutive GAL4 transgenes tubulin [ub]-GAL4 and actin [act]-GAL4; Wang and Kalderon, 2009), whereas expression of UAS-cycE in otherwise wild-type FSC clones has no effect.
depend on the induction of cycE and proliferation by Yki because yki is required for both maintenance and normal cell cycling of FSCs (Fig. 1 B, Fig. S1, and Fig. 4 A), whereas excess CycE (together with DIAP1) can rescue yki mutant FSC maintenance (Fig. 1 C).

The Hh pathway regulates FSC proliferation and intersects with the Hpo–Yki pathway

(Fig. 4 D), suggesting that UAS-cycE is expressed at roughly normal physiological levels in FSCs. The expression of UAS-cycE in ex cycE mutant FSCs restored normal FSC maintenance but did not produce a high frequency of all-marked FSC clones or the increased longevity characteristic of ex mutant FSCs (Fig. 4, D and F). Thus, the induction of FSC duplication by activated Yki to form all-marked FSC clones depends on the presence of an active Yki-inducible cycE gene. Moreover, EdU incorporation was no longer enhanced by loss of Ex in cycE mutant FSCs expressing UAS-cycE (Fig. S5 A), showing that regulation of CycE expression is also critical for activated Yki to stimulate FSC proliferation. Normal FSC maintenance also appears to depend on the induction of cycE and proliferation by Yki because yki is required for both maintenance and normal cell cycling of FSCs (Fig. 1 B, Fig. S1, and Fig. 4 A), whereas excess CycE (together with DIAP1) can rescue yki mutant FSC maintenance (Fig. 1 C).

The Hh pathway regulates FSC proliferation and intersects with the Hpo–Yki pathway

The similarity between ptc and hpo mutant FSC phenotypes (Fig. 3, C and D; and Fig. S2, H–J) prompted us to test whether Hh signaling also regulates FSC proliferation and how the Hh
and Hpo signaling pathways intersect in FSCs. The EdU labeling index was increased almost twofold for both ptc mutant FSCs and germarial FCs, whereas region 1 ECs remained quiescent (Fig. 4 A and Table S1). FSC proliferation has previously been shown to be increased by excess Hh signaling (Hartman et al., 2010). Conversely, the EdU labeling index was very low in smoothened (smo) mutant FSCs and germarial FCs, which cannot respond to Hh (Fig. 4 A). Thus, FSC proliferation is highly dependent on the level of Hh pathway activity. Furthermore, double mutant ptc yki FSC clones showed strongly reduced EdU labeling in FSCs (Fig. 4 A). Double mutant ptc yki FSC clones were also lost just as rapidly as yki FSC clones (see Fig. 6 A), showing that the effects of excess Hh on FSC duplication, maintenance, and proliferation all depend absolutely on Yki activity.

Hh signaling induces Yki activity specifically in the FSC lineage

To test whether Hh signaling might directly stimulate Yki activity, we examined two Yki target gene reporters, DIAP1-lacZ and ex-lacZ (Karpowicz et al., 2010). Both reporters were readily detected in somatic cells in the ovary but not in the germline and showed increased expression cell autonomously in hpo mutant clones, confirming that they do report Yki activity (Fig. 5, A, B, F, G, and K; and Fig. S3, A, B, D, E, and G). Interestingly, FSCs expressed significantly lower levels of both DIAP1-lacZ and ex-lacZ than more posterior FC daughters or more anterior ECs (Fig. 5, A, F, and K; and Fig. S3, A, D, and G), implying that upstream Hpo pathway components (Kibra, Ex, Mer, Sav, Wts, and Hpo) are particularly active (limiting Yki protein activity) in FSCs. Loss of yki activity reduced DIAP1-lacZ expression in FCS to a level similar to that of FSCs (Fig. S4, A, C, and I), supporting the inference that Yki activity is normally higher in FCs than FSCs.

Strikingly, loss of ptc activity markedly increased DIAP1-lacZ and ex-lacZ expression in both FSCs and FCs (Fig. 5, A, C, F, H, and K; and Fig. S3, A, C, D, F, and G), indicating that excess Hh pathway activity significantly stimulates Yki activity. The cell-autonomous activation of Yki by Hh signaling seen in the FSC lineage was not observed in wing discs, where ex-lacZ was instead induced in a ring around each anterior ptc mutant clone (Fig. S3, H–M). The nonautonomous induction of ex-lacZ is consistent with the recent observation that Decapentaplegic (Dpp) signaling activates Yki in wing discs (Kagey et al., 2012). Dpp is induced cell autonomously by Hh pathway activity in wing discs, but the ectopic Dpp acts principally on surrounding cells because Hh pathway activity also cell-autonomously inhibits expression of the Dpp receptor Thickveins (Funakoshi et al., 2001).

A previous study suggested that the transcriptional coactivator Mastermind (Mam), which normally contributes to Notch signaling, has an atypical role in FSCs, in which it collaborates closely with the Hh pathway (Vied and Kalderon, 2009). We therefore tested whether DIAP1-lacZ expression was induced by excess Hh pathway activity in the absence of Mam function. We found that DIAP1-lacZ was not increased in ptc mam double mutant FSCs or FCs (Fig. 5, E, J, and K). We also found that mam mutant FSCs have reduced EdU incorporation (Fig. S5 B) and that the maintenance of mam mutant FSCs was rescued by expression of activated Yki (Fig. 6, A and F). Thus, Mam appears to be critical for excess Hh signaling to activate Yki in the FSC lineage, and mam mutant FSC loss may be caused by insufficient Yki activity.

Hh induces yki transcription in the FSC lineage

Hh signaling normally influences cells by transcriptionally inducing cell type–specific target genes. We therefore tested whether Yki activation by Hh signaling in FSCs might be mediated by transcriptional induction of yki itself. Because reporters of Yki activity are activated by excess Hh pathway activity in both FSCs and FCs, we reasoned that the underlying mechanism was probably common to all cells of the FSC lineage and could be studied in samples containing both FSCs and FCs. We therefore isolated RNA from samples that included the gerarium and all egg chambers of an ovariole up to stage 9. Analogous samples were collected from wild-type ovarioles and from ovarioles that were populated mostly by derivatives of either ptc or hpo mutant FSCs. We then measured the relative levels of key RNAs by quantitative RT-PCR, using rp49 mRNA as an internal control. All three common Yki target genes tested (DIAP1, cycE, and ex) were induced in extracts from hpo mutant ovarioles and, to almost the same extent, in extracts from ptc mutant ovarioles (Fig. 5 L). Those observations confirmed our previous findings with ex-lacZ and DIAP1-lacZ reporter genes and also showed directly that both Hh and Hpo–Yki pathways induce cycE in the FSC lineage. Crucially, yki RNA was also induced in samples from ptc mutant FSC lineages (Fig. 5 L), showing that Hh signaling does indeed induce yki expression in the FSC lineage. In contrast, yki RNA was not induced in hpo mutant FSC lineages (Fig. 5 L), as expected because Hpo regulates Yki activity posttranscriptionally.

Hh regulates FSCs by inducing Yki transcriptionally and thereby stimulating FSC proliferation

If Hh regulates FSCs by inducing yki transcription, excess Yki should mimic the effects of excess Hh signaling. Conversely, in the absence of Hh signaling, Yki protein levels may be too low to respond to inactivation of upstream Hpo pathway regulators. We found that excess wild-type Yki was indeed sufficient to produce increased FSC longevity and a significant frequency of all-marked clones (Fig. 6, A and D) as well as increased EdU incorporation (Fig. S5 B). The percentage of all-marked FSC clones was similar for FSCs with increased Hh pathway activity (ptc FSCs), increased Yki expression (from UAS-yki), or both alterations together (ptc + UAS-yki), consistent with the idea that Hh acts by increasing yki expression (Fig. 6 A). Likewise, the phenotype of ptc hpo double mutant FSCs was similar to the phenotype of ptc or hpo single mutant FSCs (Fig. S5 C). These results are consistent with the idea that maximal Hh signaling, inactivation of upstream Hpo pathway regulators of Yki, and excess Yki expression provide three routes to activate Yki, each of which produces a saturatingly high input from Yki into the proliferation rate and competitive behavior of FSCs.
Conversely, inactivation of the upstream Hpo pathway regulator Ex, which normally leads to posttranscriptional activation of Yki, enhanced FSC maintenance, and frequent all-marked FSC clones did not induce all-marked FSC clones in the absence of smo activity and did not even counter the rapid loss of smo7.6.6 (Fig. 6 B) or smo7 mutant FSCs (Fig. S5 F). Similarly, smo hpo and smo wts double mutant FSCs did not produce all-marked clones but instead exhibited accelerated FSC loss, similar to smo single mutant FSCs (Fig. S5 D). Thus, the requirement for Hh pathway input is downstream of, or parallel to, the input of conventional Hpo pathway regulators, consistent with our demonstration that Hh acts by inducing yki RNA, whereas upstream Hpo pathway components regulate the activity of Yki protein. It also appears that regulation of Yki activity by the Hpo pathway in FSCs is only effective when normal Hh signaling provides sufficient, but not excessive, yki transcription.

Figure 5. Hh signaling increases Yki activity at the transcriptional level in the FSC lineage. (A–J) FSC clones of the indicated genotypes, marked by loss of GFP in germaria stained for Fas3 and β-galactosidase (red) encoded by a diap1-lacZ reporter. diap1-lacZ expression was lower in wild-type (WT) FSCs (arrowhead; A) than FCs (white lines) and was increased in both FSCs (arrowheads) and FCs (arrows in F–J) for hpo and ptc genotypes (B, C, G, and H) but not in ptc yki; tub-yki (D and I) or ptc mam (E and J) genotypes. Red bars, 10 µm. (K) Quantification of diap1-lacZ expression levels in FSCs and FCs of the indicated genotypes. Error bars show SDs from three measurements of the same samples, as described in the Materials and methods (n > 18 for FSCs and n > 300 for FCs). Dotted lines indicate values for wild-type FSCs and FCs. (L) Relative mRNA levels of diap1, cycE, ex, yki, and ptc (indicated on top) in ovarioles containing control (wild type, blue bars), hpo (red bars), or ptc (green bars) FSC clones induced 12 d earlier in larvae. rp49 mRNA was used to normalize total mRNA concentration for each genotype. Error bars show SDs; n = 3. Values significantly different from wild type at P < 0.05 by Student’s t test in K and L are indicated by asterisks.
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Figure 6. Regulation of FSCs by Hh is mediated principally by induction of Yki. (A–C) Percentage of ovarioles containing positively marked FSC clones 12 d after larval heat shock induction; dark red indicates all-marked clones. Each set of genotypes has the appropriate control (wild type [WT]) to the left. Horizontal black lines clarify correspondence between bars and genotypes. Black bars indicate critical comparisons. (B and C) All tests used a hypomorphic smo allele (smo[7.6.6], right, angled stripes) and some (C) also used a strong, effectively null allele (smo[2], left, solid shading). Error bars show SDs; n = 3, with ≥100 ovarioles scored in each measurement. Significant differences from wild type for all-marked FSC clones using Fisher’s exact two-tailed test with P < 0.05 are indicated by asterisks. (D–G) Ovarioles with clones induced 12 d earlier and marked by the presence of GFP, shown together with Fas3 and mam FSC clones. All-marked FSC clones were induced by excess Yki (D) but not by ptc when yki is replaced by a tub-yki transgene (E). mam FSC clones were rescued by activated Yki (F), and smo FSC clones were rescued by excess CycE together with DIAP1 (G). Red bars, 10 µm.

To test definitively whether yki induction by Hh is critical for FSC responses to Hh, we eliminated the transcriptional induction of yki by Hh signaling. To achieve this, we used a tub-yki transgene, in which the yki coding sequence is expressed from a heterologous, constitutive low-level promoter, to replace normal yki gene function. This transgene has previously been shown to rescue viability of yki mutant animals and must therefore be expressed at roughly physiological levels (Huang et al., 2005). Indeed, tub-yki almost fully rescued yki mutant FSC clone maintenance (Fig. 6 A) and EdU incorporation (Fig. S5 B) and restored normal DIAP1-lacZ expression to yki mutant FSCs (Fig. S4, C, D, and I). Also, tub-yki did not detectably increase DIAP1-lacZ expression in wild-type FSCs (Fig. S4, A, B, and I), and unlike UAS-yki or UAS-yki[270] expression, tub-yki did not cause any increase in proliferation of wild-type FSCs (Fig. S5 B). Hence, tub-yki must produce almost the same levels of Yki as in normal FSCs. When the constitutive tub-yki transgene was used to complement yki in FSCs, excess Hh signaling did not elevate DIAP1-lacZ reporter activity in ptc yki FSC clones in animals with one copy of tub-yki in all cells, ptc yki; tub-yki (Fig. 5, D, I, and K). We therefore conclude that elevation of Yki activity by excess Hh pathway activity in otherwise normal FSCs is achieved by inducing yki expression.

Crucially, ptc yki; tub-yki FSCs, which have elevated Hh pathway activity but do not have elevated Yki activity, did not have an increased all-marked frequency (Fig. 4 A) and did not produce all-marked clones (Fig. 6, A and E). Thus, all responses to excess Hh pathway activity seen in ptc mutant FSC clones (Yki activation, increased proliferation, and increased FSC competitiveness) were lost when normal regulation of Yki was substituted by constitutive Yki expression at a level appropriate for normal FSC function.

We also tested whether upstream Hpo pathway components can regulate Yki activity when normal transcriptional regulation of yki is prevented. We found that the phenotype of hpo yki; tub-yki FSCs was very similar to that of hpo mutant FSCs, producing many all-marked FSC clones (Fig. S5 C). Thus, upstream Hpo pathway components do not regulate Yki activity by modulating yki transcription, in accordance with precedent in other tissues and with our observation that yki RNA levels are unaltered in hpo mutant FSC clones (Fig. S5 L). Thus, substitution of yki with tub-yki clearly outlines the contrast between...
the effects of the Hh pathway on yki RNA levels and Hpo pathway components on Yki protein activity and defines Yki as the point of convergence of these two pathways in FSCs.

To investigate the importance of yki induction for FSC responses to normal levels of Hh signaling, we first tested whether excess Yki activity could restore deficits of smo mutant FSCs. We found that expression of activated Yki restored EdU incorporation in smo mutant FSCs and FCS to almost normal levels (Fig. 4 A) and substantially rescued smo^1^6^6 (Fig. 6 B) and smo^2^ mutant FSC maintenance (Fig. S5 F). Thus, both the deficits in FSC proliferation and FSC maintenance as a result of loss of Hh pathway activity are substantially complemented by provision of excess Yki activity. We also found that smo mutant FSC maintenance was substantially rescued by the combined expression of excess CycE and DIAP1, two key targets of Yki in FSCs, but not by either alone (Fig. 6, C and G), just as observed for rescue of yki mutant FSCs (Fig. 1 C). Finally, we saw that DIAP1-lacZ activity in FCS was reduced in smo mutant FCSs and restored by tub-yki (Fig. S4, G–I). DIAP1-lacZ expression in FSCs is too low to determine whether it can be reduced further by yki or smo mutations. These results support the idea that induction of Yki is an essential response to Hh signaling in normal FSCs.

However, some observations indicate that Yki induction is not the only way that Hh normally influences FSCs. Most convincing, excess wild-type or activated Yki induced many all-marked FSC clones, even in yki mutant FSCs but not in smo mutant FSCs (Fig. S5 E). Second, low levels of Yki, supplied by the tub-yki transgene, did not rescue smo FSC maintenance completely. In fact, rescue was slightly less effective than for yki mutant FSC maintenance (Fig. 6, A and B), even though Yki activity is compromised more by yki than by smo mutations (Fig. S4 I). Thus, activation of Yki and stimulation of cell cycling appear to be critical components of the response to normal Hh pathway activity in FSCs, whereas excessive induction of Yki and hyperproliferation are essential for the duplication and increased competitiveness of FSCs responding to excess Hh pathway activity.

Discussion

Regulation of proliferation is clearly critical for all types of adult stem cells to ensure adequate, but not excessive, production of daughter cells for tissue maintenance. Our studies with FSCs have revealed a specific mechanism for regulating stem cell proliferation that centers on the Hpo–Yki pathway, a major conserved pathway that was initially recognized for its roles in regulating tissue size. Most importantly, the inductive connections among Hh, Yki, and CycE, coupled to their tightly correlated effects on FSC proliferation and maintenance, provide compelling evidence that the longevity and competitive strength of an FSC are regulated by controlling the rate of FSC proliferation.

Our results clearly show that an FSC lacking Yki or Hh pathway activity has reduced proliferation and is rapidly lost, whereas an FSC with excessive Yki or Hh pathway activity has an increased proliferation rate, extended longevity, and frequently duplicates and outcompetes wild-type FSCs in the same germarium. It is also clear that excess Hh pathway activity increases Yki activity in FSCs and that the increased proliferation, longevity, and competitive strength of such FSCs is completely suppressed when the endogenous yki gene is inactivated in animals that have a tub-yki transgene. We argue that the latter observation shows that all responses to excess Hh pathway activity depend on inducing Yki activity.

We measured a three- to fourfold increase in FSC Yki activity using a DIAP1-lacZ reporter and an induction of less than twofold for both yki and diap1 RNA in quantitative RT-PCR measurements using ovarioles with ptc mutant FSC clones. The latter experiment almost certainly underestimates induction in FSCs because the samples included some wild-type FSC lineages and many ptc mutant FCSs, which show less induction of Yki activity than FSCs (Fig. 5 K). We therefore surmise that a three- to fourfold increase in Yki activity gives an FSC a large competitive advantage.

FSCs expressing UAS-Yki and UAS-Yki^S168A presumably also reached this threshold of excessive Yki activity because they induced many all-marked FSC clones and increased EdU incorporation in FSCs. In contrast, the tub-yki transgene likely produces Yki levels similar to those in wild-type FSCs because DIAP1-lacZ activity and EdU incorporation in yki mutant FSCs expressing tub-yki were not significantly higher or lower than in wild-type FSCs (Fig. S4 and Fig. S5). Hence, complementation of yki loss of function with tub-yki provided a suitable, normal level of yki expression to test the effects of elevating Hh pathway activity. In those tests, there was no measurable response at all to loss of ptc, even though ptc mutations in a wild-type background strongly increase the expression of reporters of Yki activity (DIAP1-lacZ and ex-lacZ) and produce very strong phenotypes of increased FSC proliferation, FSC longevity, and FSC duplication. Hence, we have no reservations in concluding that excess Hh pathway activity makes an FSC more competitive by increasing yki transcription and hence Yki activity.

The next critical question is whether increased Yki activity makes an FSC more competitive by increasing its proliferation rate. We showed that the competitive advantage of FSCs with increased Yki activity (in ex mutant FSC clones) was lost when induction of CycE activity was lost (cycE^465^5) or greatly reduced (cycE^S168A) by complementing cycE mutations with a UAS-cycE transgene. The same UAS-cycE transgene almost fully complemented the FSC maintenance defects of both cycE alleles without inducing all-marked FSC clones in these or otherwise normal FSCs, as shown previously (Wang and Kalderon, 2009) and repeated here. We therefore conclude that CycE activity provided by the UAS-cycE transgene is likely slightly lower than in wild-type FSCs. Hence, CycE activity induced by excess Yki activity acting on a normal cycE gene must be much higher than the activity provided by UAS-cycE. The failure of Yki activation to induce all-marked FSCs, or indeed to alter the longevity of cycE; UAS-cycE FSCs at all, can therefore be attributed, logically and reasonably, to a failure to induce cycE transcriptionally. We conclude that increased CycE activity is critical for increased Yki activity, and therefore, by inference, for excess Hh pathway activity, to make an FSC more competitive. Whether increased DIAP1 activity is also critical was not tested.
The substantial rescue of smo FSC maintenance and smo FSC EdU incorporation by a tub-yki transgene that approximates normal levels of yki expression in FSCs (Fig. 4 A and Fig. 6 B) suggests that yki induction by Hh signaling is also important for the proliferation and maintenance of normal FSCs. The logical extrapolation that cycE induction (via Yki) is also a critical consequence of normal Hh signaling is further supported by the finding that smo mutant FSC clones are substantially rescued by coexpression of DIAP1 and a CycE transgene at a level that produces approximately normal levels of CycE activity (Fig. 6 C). As discussed earlier, Yki does not appear to be the only significant target for normal Hh signaling in FSCs because excessive Yki activity supplied by a transgene (UAS-yki or UAS-yki

Importantly, FSCs have provided an opportunity to understand how an essential proliferative action of the Hpo pathway is integrated into the normal physiological regulation of a stem cell population, revealing two key relationships. First, for FSCs, we find that regulated stem cell proliferation and stem cell maintenance are causally and positively linked. That relationship was originally considered unlikely for any stem cell based on considerations of the potential for replicative DNA damage and evidence of HSC loss in response to enforced proliferation. However, even for HSCs, the direct impact of regulated proliferation has been difficult to assess definitively because a few genetic manipulations that increase HSC division rates (such as loss of Cdkn2c/p18 or c-Myb) do not impair HSC maintenance, whereas the few direct manipulations of niche factor signaling responses that affect both stem cell proliferation and maintenance do not prove a causal connection between these two responses (Orford and Scadden, 2008; Pietras et al., 2011; Yamazaki et al., 2011; Rossi et al., 2012). In contrast, it is likely that maintenance of highly proliferative stem cells is, at worst, indifferent to high rates of stem cell division. Indeed, Wnt signaling can promote proliferation of mammalian intestinal stem cells, and loss of Wnt signaling pathway components or a key transcriptional target, c-Myc, implicated in growth control was shown to result in stem cell loss, suggesting that proliferation likely stimulates intestinal stem cell maintenance (Clevers, 2013). However, even in this well-studied paradigm, it remains possible that responses to Wnt or c-Myc other than changes in stem cell proliferation rate play an important role in regulating stem cell maintenance (Hsu and Fuchs, 2012; Clevers, 2013). A key strength of our experiments was the demonstration that it is specifically the proliferative response to Hh signaling, which acts directly in a cell-autonomous manner to regulate the maintenance and competitive behavior of a stem cell. It will be very important to establish rigorously whether regulated stem cell proliferation and maintenance are indeed causally linked in some mammalian stem cells, as in FSCs, because this has important consequences for cancer biology. As illustrated by ptc or hpo mutations in FSCs, activation of either regulatory pathway in a single stem cell would, in one step, induce expansion and increased longevity of the activated stem cell and confer inherited hyperproliferation on daughter cells.

Second, we discovered how stem cells can be regulated by tissue-specific coupling of two widely used and conserved signaling pathways, Hh and Hpo. Normal FSC behavior was previously shown to require a precise level of Hh pathway activity (Vied et al., 2012), which we now find is translated into Yki and CycE activities sufficient to stimulate appropriate, but not excessive, FSC proliferation. This is achieved by linking Hh and Hpo pathways through transcriptional induction of yki, involving a noncanonical role of the Notch coactivator Mam. This mode of coupling is inherently versatile because the transcriptional targets of Hh and other signaling pathways vary extensively according to cell type; indeed, the robust coupling we observed in the FSC lineage was not evident in wing disc cells. An analogous tissue-specific linkage in mammals would provide a mechanism for selectively regulating a specific stem cell or progenitor cell type, with the accompanying susceptibility to...
cancer-initiating mutations. In fact, it has already been found that Sonic Hh stimulates both transcription of the Yki ortholog YAP1 and proliferation in human cerebellar granule neural precursors and that elevated YAP1 activity contributes to the cancerous phenotype of some murine medulloblastomas induced by excess Hh pathway activity (Fernandez-L et al., 2009). It remains to be established whether YAP1 is involved in the regulation of other Hh-responsive stem cells or Hh-initiated tumors in mammals.

Materials and methods

**Drosophila stocks and clonal analysis**

For clonal analyses, we used FRT19A (control), 32-47 (control), 342-47 (control), ex342-47 (control), FRT40A, ed270 FRT40A, ed72 FRT40A, fo7 FRT40A, d315 FRT40A, FRT42D (control), FRT42D yk38, FRT42D hpo2-47, FRT42D hpo2-472, FRT42D ptcB2, 82-04 NM (control), FRT40A dRasfl36, FRT82B crb1422, FRT82B crb2916, FRT82B sav, FRT82B wts2, and FRT82B kibra82, ex4 FRT82B kibra82. These flies were crossed to hs-flp; ubi-mRFp. nls FRT19A, hs-flp, ubi-gFP; nls FRT40A, hs-flp; FRT42D ubi-gFP; nls, or hs-flp; FRT42D ubi-GFP; nls flies for negatively marked clones. For positively marked clones, we either used alleles on an FRT40A chromosome (NM, smo, smo2-6.6, eyET4a, eyET4b, eyET4c, eyET4d, ex342-47 smo, or ex47 smo2-6.6) crossed to hs-flp, UAS-gFP, tub-GAL4; tub-GAL80 FRT40A/Cyo; act5-CD2-Ds-Gal4 or alleles on an FRT42D chromosome (sha, yki3, hpo, ptc1, ptc2, yki1, man1, or ptc2 man1) crossed to hs-flp, UAS-gFP; tub-GAL4; FRT42D tub-GAL80/Cyo; act5-CD2-Ds-Gal4, UAS-cycE, UAS-DIAP1, UAS-DIAP1 + UAS-cycE, UAS-yki1868, UAS-yki1869, tub-yki1, UAS-yki1, UAS-yki RNAi, UAS-yki RNAi, and UAS-yki2 RNAi were added to the clones of these crosses when needed. Third chromosome Hpo pathway reporters DIAPI-1acx and exacx were added to stocks for negative clone marking. To induce smo hpo double mutant clones, we used flies of the genotype yh hs-flp; smo1 FRT42D hpo2-47/smo1 FRT42D P(sm0) ubi-gFP; nls and yw hs-flp; FRT42D hpo2-47 (or hpo1)/sm0 FRT42D P(sm0) ubi-gFP; nls flies to generate hpo mutant and control clones in a smo+ background. To induce smo wts double mutant clones, we used flies of the genotype yh hs-flp; smo1 FRT40A/ubi-gFP; FRT40A; TR32B wts2/ubi-mRFp FRT82B H2Av-RFP with controls lacking the smo mutation. Larvae or adult flies of the appropriate genotype were heat shocked for 1 h at 37°C and then dissected 12 h after heat shock. The germarium and early egg chambers (up to stage 9) were collected using the hyperplastic ptc and hpo phenotypes to identify ovarioles harboring those mutant FSC clones. RNA was isolated using the RNaseasy Mini kit (QIAGEN) according to the manufacturer’s protocol. Quantitative RT-PCR was performed in the StepOne Plus Real-Time PCR System with Power SYBR Green RNA-to-Ct Kit (Applied Biosystems). Primers for amplifying 100–200 bp of each PCR product are listed in Table S2. RT-PCR reactions were performed for 30 min at 48°C followed by 10 min at 95°C and then followed by 40 cycles of two-step PCR for 15 s at 95°C and 1 min at 60°C. Each sample was performed in triplicate. The mRNA levels of target genes (DIAP1, cycE, ey, yki, and ptc) were normalized to rp49 mRNA, and the relative quantification in gene expression between mutants and control was determined using the 2−ΔΔCt method (Livak and Schmittgen, 2001).

**Online supplemental material**

Fig. S1 shows that yki mutant FSCs are lost very rapidly and that yki mutant FCS have increased apoptosis. Fig. S2 shows that core components of the Hpo pathway regulate FSCs, whereas several known upstream regulators do not. Fig. S3 shows that excess Hh activity induces Yki activity, measured with an exacx reporter, cell autonomously in FSCs but not in wing disc cells. Fig. S4 shows that a tub-yki transgene rescues Yki activity in yki, ptc yki, and smo mutant FCS but does not confer excessive Yki activity on wild-type cells. Fig. S5 shows that EdU incorporation is no longer enhanced by excess Yki activity in FSCs when cycE is not inducible, that Yki transgenes complement proliferation defects and FSC maintenance defects in yki, smo, and mammalian FSCs to different degrees, that the hpo mutant phenotype is not altered by additional loss of ptc or substitution of yki with a constitutive yki transgene, and that smo mutant FSC maintenance is restored by a hyperactive Yki transgene but not by activation of Yki in response to loss of the upstream regulators Hpo, Wts, or Ex. Table S1 shows fractions of GFP-labeled cells of the indicated genotypes and cell types with EdU labeling. Table S2 shows primers used for quantitative real-time PCR. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201309141/DC1.

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