The Yb protein defines a novel organelle and regulates male germline stem cell self-renewal in Drosophila melanogaster

Akos Szakmary,1 Mary Reedy,1 Hongying Qi,2,3 and Haifan Lin1,2,3

1Department of Cell Biology, Duke University Medical School, Durham, NC 27710
2Yale Stem Cell Center and 3Department of Cell Biology, Yale University School of Medicine, New Haven, CT 06520

Introduction

Stem cells are characterized by their ability to both self-renew and generate a large number of differentiated progeny. Tissue stem cells usually operate at a steady state, producing one daughter stem cell and one differentiated cell through each division. The division of stem cells is largely controlled by their inductive microenvironment, termed the stem cell niche, which is formed by stable supporting cells often called niche cells (for reviews see Lin, 2002; Morrison and Spradling, 2008). The concept of a stem cell niche was first proposed for the human hematopoietic system (Trentin, 1970). A major challenge in stem cell biology is to define the properties of niche cells and mechanisms through which they regulate the behavior of stem cells.

The organization of niche cells and their essential function in regulating stem cell self-renewal were first defined in the D. melanogaster ovary (Lin and Spradling, 1993, 1997; Cox et al., 1998; Xie and Spradling, 1998, 2000; King and Lin, 1999). The Drosophila ovary and testis are essentially tubular structures, with the apical end consisting of somatic niche cells that are in contact with germline stem cells (GSCs; Lin, 2004; for review see Lin, 2002). Each D. melanogaster ovary consists of 16–18 tubular structures called ovarioles. In each ovariole, 8–10 terminal filament (TF) cells form a stack at the very anterior end (Fig. 1, A–D). Adjacent to them are five to six cap cells that cover the anterior side of two to three GSCs. Oogenesis is initiated when a GSC divides asymmetrically to produce a daughter GSC and a differentiating daughter cell termed the cystoblast. The cystoblast then undergoes four rounds of synchronous divisions with incomplete cytokinesis to produce a germline cyst that contains 16 cells interconnected by cytoplasmic bridges called ring canals (for review see Deng and Lin, 2001). As a 16-cell cyst moves to the middle region of the gerarium, two to three somatic stem cells (SSCs) at the periphery of this region divide to produce a monolayer of follicle cells that encapsulate the cyst to form an egg chamber (Margolis and Spradling, 1995). The egg chamber then buds off of the gerarium and continues to grow and mature. In the D. melanogaster testis, the 10–12 anterior somatic niche cells form a dense sphere of similar size that appears to be RNA rich. There are one to two Yb bodies/cell, often located close to germline cells. The N-terminal region of Yb is required for hh expression in niche cells, whereas the C-terminal region is required for localization to Yb bodies. The entire Yb protein is necessary for piwi expression in niche cells. A double mutant of Yb and a novel locus show male germline loss, revealing a function for Yb in male germline stem cell maintenance.

Yb regulates the proliferation of both germline and somatic stem cells in the Drosophila melanogaster ovary by activating piwi and hh expression in niche cells. In this study, we show that Yb protein is localized as discrete cytoplasmic spots exclusively in the somatic cells of the ovary and testis. These spots, which are different from all known cytoplasmic structures in D. melanogaster, are evenly electron-dense spheres 1.5 µm in diameter (herein termed the Yb body). The Yb body is frequently associated with mitochondria and a less electron-dense sphere of similar size that appears to be RNA rich.

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D. melanogaster germ cells contain several specialized cytoplasmic structures, including the spectrosome, fusome, Balbiani body, sponge body, nuage, and U body. The spectrosome (Lin et al., 1994; Lin and Spradling, 1995; Deng and Lin, 1997) is a cytoplasmic organelle rich in membrane skeletal proteins and vesicular materials. In female GSCs, it anchors one pole of the GSC spindle to a niche cell such that the daughter GSC remains associated with the niche cell. The other daughter cell receives a smaller portion of the spectrosome and differentiates to the cystoblast. During the subsequent four cystoblast divisions, a group called the hub (Fig. 1 H). The hub is surrounded and contacted by six to nine GSCs, each of which is flanked by a pair of SSCs (Jones et al., 2004). A GSC and its two flanking SSCs undergo asymmetrical division in synchrony. After each division, the daughter GSC and SSCs remain in contact with the hub, whereas their sibling cells differentiate into a gonialblast and two somatic cyst cells, respectively. The gonialblast, sandwiched by the two somatic cyst cells, enters four rounds of mitotic divisions to form a 16-cell germline cyst. Subsequent meiotic divisions result in 64 interconnected spermatids.
divisions, the spectrosome grows into an intercellular structure called the fusome that connects the resulting 16 cells in a germ-line cyst through ring canals. In addition, the fusome is involved in specifying 1 of the 16 cells in the cyst as an oocyte (Deng and Lin, 1997).

*D. melanogaster* germline cysts also contain Balbiani bodies that are aggregates of mitochondria, ER, Golgi, and other cytoplasmic organelles (Cox and Spradling, 2003). Balbiani bodies have been described for many organisms, with a proposed role in the development of germ plasm. In addition to Balbiani bodies, nurse cells and oocytes contain sponge bodies that are associated with RNA and frequently with mitochondria (Wilsch-Bräuninger et al., 1997). Sponge bodies consist of ER-like cisternae, but they are not specialized Golgi complexes. Instead, they are linked to the assembly and transport of maternal products within the oocyte during *D. melanogaster* oogenesis. At present, it is not known whether Balbiani and sponge bodies are related organelles. Interestingly, the sponge body appears to have overlapping functions with the yeast P body, a cytoplasmic structure which is responsible for programmed mRNA degradation (Sheth and Parker, 2003). This is because Me31B, the *D. melanogaster* orthologue of a key component of the P body in yeast called Dhh1p (an activator of decapping; Coller et al., 2001), is localized to the sponge body (Nakamura et al., 2001). A recently identified U body is also localized in the cytoplasm of the nurse cells and oocyte (Liu and Gall, 2007). U bodies contain major U small nuclear RNPs (snRNPs) and associate with P bodies, suggesting that U bodies may be involved in the assembly and storage of snRNPs and regulation of the snRNP’s metabolism.

Another germline-specific organelle is the nuage, a perinuclear structure which is rich in RNA and proteins (Mahowald, 1971). The nuage is made up of electron-dense granules. It contains the putative DEAD box RNA helicase Vasa (Hay et al., 1988; Lasko and Ashburner, 1990) and the RNA-binding protein Aubergine (Harris and Macdonald, 2001) as well as Maelstrom (Findley et al., 2003) and Tudor (Bardsley et al., 1993). Because Vasa and Tudor are also essential components of polar granules, which are germ plasm–specific organelles in the *D. melanogaster* embryo, and because polar granules display nuagelike texture under an electron microscope, the nuage is considered a possible precursor of polar granules and has been suggested to function as an intermediate of cytoplasmic RNP particle assembly (Findley et al., 2003). In contrast to these specialized structures in the germline, no comparable cytoplasmic structure has been described in somatic cells of the ovary or testis.

*Yb* was the first gene identified to regulate two different types of stem cells in an organ (King and Lin, 1999; King et al., 2001). It encodes a novel protein that is specifically required in ovarian somatic niche cells for the self-renewal of both GSCs and SSCs, which produce germline and follicle cells, respectively. Loss of *Yb* function eliminates GSCs and reduces SSC division, whereas overexpression of *Yb* increases GSC number and causes SSC overproliferation. *Yb* appears to achieve this dual regulatory function via the *piwi*– and *hh*–mediated signaling pathways that emanate from the same niche cells to control GSC and SSC division, respectively (King et al., 2001). In addition to its stem cell function, *Yb* is required for follicle cell differentiation by interacting with the *Notch* group of neurogenic genes (Johnson et al., 1995). The *Notch* group genes function in both prefollicular and follicle cells in the germarium as well as in follicle cells in postgerminal egg chambers (Ruohola et al., 1991). Therefore, *Yb* is also expected to be expressed in all of these somatic cells for its function in follicle cell differentiation. Because *Yb*-null mutant males are fertile, *Yb* is not required in spermatogenesis (King and Lin, 1999). However, it remains to be determined whether *Yb* is involved in spermatogenesis.

The biochemical mechanism by which *Yb* achieves its regulation of GSC and SSC divisions is also unknown. *Yb* encodes a novel protein with localized regions bearing similarity to DEAD/DEAH box RNA helicases and a Tudor domain. DEAD/DEAH box RNA helicases are involved in many biological processes (Rocak and Linder, 2004), whereas the Tudor domain is a protein–protein interaction domain (Selenko et al., 2001; Sprangers et al., 2003). To explore the biochemical function of the *Yb* protein, in this study, we report the determination of its expression and subcellular localization pattern. This has led to the discovery of a novel cytoplasmic structure in the somatic cells of the ovary and testis. Furthermore, we describe discrete functional domains of *Yb* that are responsible for regulating GSC self-renewal, SSC division, and localization to the Yb body. Finally, we report the novel function of *Yb* in regulating male GSC self-renewal.

**Results**

**Yb** protein is localized to discrete spherical structures in ovarian somatic cells

To explore the molecular function of the *Yb* protein, we examined its expression and localization pattern. Previous RNA in situ hybridization experiments indicated that the *Yb* mRNA is most abundant in TF cells and cap cells of the ovary (King and Lin, 1999). To detect the *Yb* protein, we produced affinity-purified polyclonal rabbit antibodies against two different *Yb* peptides (see Materials and methods). Immunofluorescence localization of *Yb* using these antibodies reveals labeling of discrete spots in all somatic cells in the ovary (Fig. 1, A and E–G). These bright spots are present in high density in the cap cell region and are less dense in the follicle cell layer. However, they are absent from many TF cells. In addition to the spots, a low level of diffuse labeling is detectable in the cytoplasm of both somatic and germline cells.

We verified that the bright, discrete spots but not the diffuse staining represent the *Yb* protein by the following three additional and independent criteria. First, in the *Yb*Δ2 mutant ovary, in which the *Yb* protein is truncated upstream of the antibody epitope regions, no discrete *Yb* spot is detectable, yet the low level of diffuse staining still persists (Fig. 1 B). The much smaller spots seen in Fig. 1 B (which is shown at a three times higher magnification than Fig. 1 A) are spectrin-rich spheres that have been reported for *Yb* mutants (King and Lin, 1999). Second, when an N-terminal His6–Flag–tagged *Yb* transgene that fully rescues the defects of *Yb* mutants (see Materials and...
methods and Fig. 5) was stained using three different monoclonal anti-Flag antibodies and a polyclonal Flag antibody, the same staining pattern was observed (Fig. 1 C). The Flag-positive spots are again not detectable in the sibling flies that do not carry the His6-Flag-Yb transgene (Fig. 1 D). Third, we constructed a new N-terminal 5× Flag–tagged Yb transgene that also fully rescues the Yb mutant (see Materials and methods). In this transgene, each Flag is separated by 3 aa residues. When ovaries from flies carrying this transgene were costained by a polyclonal Flag antibody and an anti-Yb antibody, the two antibodies labeled the same discrete spots (Fig. 1 E). Once again, these spots were no longer detectable by anti-Flag antibody in the sibling flies that lack the transgene (unpublished data).

Inspection of the discrete Yb antibody–reactive spots at high magnification revealed that they have a distinctive shape and are present singly or in pairs in each cell. The high density of the spots in the cap cell region (Fig. 1, A, C, E, and G) is the result of the small size and squamous shape of these cells, whereas the one to two spots/cell distribution is clearly seen in the larger follicle cells that envelop individual egg chambers (Fig. 1, F and G). This distribution is clearly seen even in a low magnification image focused on the follicle cells at the surface of an egg chamber (Fig. 1 F). It is also evident in a sagittal section image of an ovariole (Fig.1 G), in which the Yb spots highlight the somatic cells in germaria and egg chambers.

Yb protein is also localized to discrete spheres in somatic cells of the D. melanogaster testes

Although Yb mutant males are normal and fertile, previous RNA hybridization and Northern blot analyses indicate that Yb is expressed in the testis (King and Lin, 1999). To examine the expression of the Yb protein in the testis, we conducted similar immunofluorescence microscopy to the wild-type D. melanogaster testis. Like in the ovary, the Yb protein is localized in discrete spots (Fig. 1, H and I). Moreover, these spots are highly concentrated in hub cells (the cap cell equivalents in the testis) and present in somatic cyst cells (the follicle cell equivalents in the testis), but they are not detectable in the germline. This testicular expression pattern of Yb, which is parallel to its ovarian expression pattern, indicates that the Yb protein might perform the same function in females and males, even though such function may be redundant in males.

The Yb-containing sphere does not colocalize with known cytoplasmic organelles in D. melanogaster
To examine whether the Yb-containing spheres present any known cytoplasmic organelles, we conducted immunocolocalization experiments of Yb and markers of the known cytoplasmic organelles in the D. melanogaster reproductive system. Double labeling of 5× Flag–tagged Yb transgenic fly ovaries with anti-Flag antibody and one of the antibodies against markers of the ER (Trap-ε; Nicchitta et al., 1991), the nuage (Vasa; Lasko and Ashburner, 1990), the P body (Tral; Wilhelm et al., 2005), the sponge body (Me31B; Nakamura et al., 2001), and the U body (dLsm11; Liu and Gall, 2007) reveals that the Yb-containing spherical structure does not colocalize with any of these structures (Fig. 2). The results, together with electron microscopic analysis (see next section), indicate that the Yb-containing sphere represents a novel cytoplasmic organelle.

The Yb-containing sphere represents a novel cytoplasmic organelle in gonadal somatic cells

To further investigate the subcellular localization and function of the Yb protein, we conducted immunoelectron microscopy. Ovaries of 5× Flag–tagged Yb transgenic flies were dissected and stained with the anti-Flag antibody and labeled by a secondary antibody conjugated with both 1.4-nm gold particles and a fluorescent label (see Materials and methods). The combination of these two labels allowed visualization of the same sample by both light and electron microscopy. Immunoelectron microscopy revealed that the Yb protein is localized in discrete electron-dense structures in all somatic cells in the gerarium (Figs. 3 and 4), including cap cells (Fig. 3, A–E), escort stem cells and escort cells in region I (Figs. 3 A and 4 A), follicle stem cells (also known as SSCs) in region IIa (Figs. 3 A and 4 C), invaginating prefOLLICULAR cells in region IIb (Figs. 3 A and 4 E and F), and follicle cells in region III (Figs. 3 A and 4 G). In addition, the Yb-enriched, electron-dense structures are also present in follicle cells of postgermarial egg chambers (unpublished data). We termed these novel structures Yb bodies.

In all of these somatic cell types, Yb bodies show a relatively uniform morphology and localization pattern. Yb bodies are evenly electron-dense round structures, with diameters around 1.5 µm (Figs. 3 and 4). They do not contain membranes, microtubules, microfilaments, or ribosomes, morphological features which are obvious under conventional transmission electron microscopy (Fig. 3, compare B with E; and Fig. 4, compare A with B and C with D). There are one or two Yb bodies per cell, usually in close proximity to mitochondria. In cap cells and SSCs, at least 90% of Yb bodies are adjacent to an unusual round structure of similar size with lower electron density (Figs. 3, B and C; and 4 C). In some cases, the gold labeling of Yb flanks this less electron-dense structure, suggesting that Yb protein may form a toroid encircling other components in cap cells and SSCs. Cap cells often contain one Yb body that is located at the apex region near the TF cells (Fig. 3, B and C) and a second Yb body located adjacent to GSCs (Fig. 3 D). In prefollicular and follicle cells, Yb bodies...
suspected that Yb bodies may have similar functions. To examine whether the Yb body contains RNA, we costained the whole-mount wild-type ovaries with SYTO RNAsensor fluorescent cell stain to visualize RNA and the anti-Yb antibody to identify the Yb body (see Materials and methods). Indeed, Yb bodies in both ovaries and testes are often closely apposed to a spherical structure highly enriched in RNA (Fig. 1, K and L). The size and the association of this RNA-rich structure correspond to the less electron-dense structure that is associated with the Yb body.

Figure 2. Yb-containing spheres do not co-localize with known organelles in the D. melanogaster ovary. (A–E) Immunofluorescent costaining of Yb (green) and a marker (red) of the ER (A and A’), the nuage (NU; B), the sponge body (SP; C and C’), the P body (P; D and D’), or the U body (U; E) in wild-type germaria and early stage egg chambers. All of the germaria and egg chambers were oriented with the apical side to the left and the basal side to the right. tf, TF cells; cc, cap cells. s1, s2, and s3 designate stage 1, 2, and 3 egg chambers, respectively. Bar, 10 µm.
in cap cells and SSCs. The close association of the Yb body with the RNA-rich structure suggests that these two structures might be involved in RNA-related processes.

To quantify the Yb protein distribution between the Yb body and the rest of the cytoplasm, we counted the number of gold particles in the Yb body versus the rest of the cytoplasm. In cap cells and other somatic cells, 93.1 ± 1.9% of gold particles are in the Yb body, whereas only 6.9 ± 1.9% of gold particles are in the rest of the cytoplasm. This quantification indicates that almost all Yb protein molecules are localized to the Yb body, which is consistent with the immunofluorescence staining analysis that the diffuse anti-Yb staining does not represent the Yb protein.
Yb is involved in male GSC maintenance

Although Yb protein localizes in a comparable pattern in ovaries and testes (Fig. 1), null Yb mutations do not result in sterility in males (Fig. 5, A and B). Thus, if Yb performs similar functions in males, another protein or pathway must exist to compensate for Yb function in the mutant males. To search for this compensatory gene, we discovered that Yb mutant flies became sterile when they were also homozygous for a P-element insertional line called P[myc-piwi]G38 (herein abbreviated G38), in which P[myc-piwi] designates the P-element construct containing a functional myc-tagged piwi transgene, whereas G38 designates this particular insertional line (Cox et al., 2000). Immunofluorescence microscopic analysis of the Yb\textsuperscript{G38} double mutant further revealed that its testes are often completely, and less often nearly completely, depleted of the germline (Fig. 5). In the case of nearly complete depletion, there are only a small number of arrested early spermatogenic cells. Similar, although somewhat less severe, defects are also seen for the Yb\textsuperscript{G38} double mutant (unpublished data). These defects clearly indicate that functional GSCs are not maintained during spermatogenesis.

The Yb\textsuperscript{G38} synthetic phenotype is likely caused by the disruption of a gene at the insertion site by P[myc-piwi] but not caused

\textbf{Figure 4.} Electron microscopy of the \textit{D. melanogaster} germarium revealing Yb bodies in regions II and III. (A) An immunelectron micrograph of the frequently observed association of a Yb body (arrow) with a less electron-dense round structure (asterisk), a denser round structure, and mitochondria in somatic inner sheath cells [S] in region Ila. (B) A similar structure in an escort cell [arrow] seen by transmission electron microscopy. An adjacent gray spherical body (asterisk) presumably corresponds to the RNA-enriched spot in Fig. 1 J, based on its size, location, and association with the Yb body. (C) SSCs at the region Ila-IIb border show gold labeling (arrow) of a Yb body on one side of a less electron-dense structure. (D) A similar structure in an SSC (arrow) seen by transmission electron microscopy. (E and F) Immunoelectron micrographs showing that Yb bodies (arrows) in prefollicular cells are usually associated with mitochondria [M]. (G) A Yb body (arrow) in follicle cells (FC) in region III associated with mitochondria. EMS, epithelial muscle sheath; G, germline cyst; GN, germline nucleus; PFC, prefollicle cell; SN, somatic nucleus. Bars, 2 µm.
by the expression of the myc-piwi transgene or any other effect of the inserted P construct. This is based on the following two lines of evidence. First, the same construct inserted at other genomic sites such as P[myc-piwi]G47 also expresses the functional myc-piwi at normal levels (Cox et al., 2000). However, mutants homozygous for P[myc-piwi] at these sites and Yb do not display the synthetic male sterile phenotype. This indicates that the synthetic effect is insertion site specific but not caused by the expression of myc-piwi or another effect of the construct. Second, wild-type flies homozygous for both G38 and P[myc-piwi]G47 insertions are normal and fertile (Fig. 4 B). Because both G38 and P[myc-piwi]G47 express functional myc-piwi, this indicates that even six copies of functional piwi genes (four transgenic and two endogenous) in flies do not affect male fertility (Cox et al., 2000; Megosh et al., 2006). Thus, the loss of fertility in these flies is caused by the disruption of a gene by the P element at the insertion site rather than by the increased dosage of piwi. Together, these observations reveal that Yb is involved in GSC maintenance, even though this role is not essential.

The N-terminal region of Yb is required for hh expression, whereas the entire Yb protein is needed for piwi expression and SSC maintenance

The Yb protein contains a 166-aa region (residues 396–562) homologous to the DEAD/DEAH box RNA helicases and a 127-aa Tudor-like domain (residues 817–944; Fig. 5). The DEAD/DEAH box RNA helicases family has many members and is involved in numerous processes. However, the only known D. melanogaster proteins that contain both DEAD/DEAH box RNA helicase and Tudor domains are Homeless (also known as Spindle-E) and two uncharacterized proteins with similarities to Yb (CG31755 and CG11133).

To explore the functions of the different regions of the Yb protein, we generated cDNA genomic DNA transgenes that express various deleted or point-mutated variants of Yb and used them for in vivo rescue experiments. The deletion/point mutation series were designed according to Yb sequence features. Fig. 6 shows a schematic summary of all of these constructs sequentially aligned from the N to C terminus. For comparison, the mutational sites of the four known Yb mutant alleles are also indicated on the wild-type transgene. None of these transgenes was able to rescue the fertility of the null Yb72 allele beyond slight improvement (Fig. 6), indicating that all regions of Yb are essential for fertility.

Despite this, we noticed that the phenotypes of the ovaries varied depending on whether they carried either an N-terminal deletion or other mutations. This suggested different effects of mutations on oogenesis. Specifically, N-terminal deletions show fused egg chambers, suggesting underproliferation of follicle cells that resembles the hh mutant phenotype (Fig. 7;...
from D. melanogaster), revealed a very strong conservation for the first N-terminal 150 aa, followed by a lower degree of conservation in short stretches of fewer than 10 aa residues, up to the RNA helicase domain (Fig. 6). The Yb proteins show very high homology in the last 470 residues at the C-terminal region. This pattern of homology corresponds well with our deletion/mutation analysis, suggesting that the N-terminal 150 aa residues may be important for hh expression in cap cells, whereas the C-terminal 470 residues may be essential for Yb protein localization via binding to protein partners.

Yb\(^1\), Yb\(^4\), and Yb\(^6\) mutant proteins fail to localize to the Yb body

The Yb\(^1\), Yb\(^4\), and Yb\(^6\) alleles are epithelial muscle sheath–induced point mutations that are genetic null (Young and Judd, 1978; Mohler and Carroll, 1984; King and Lin, 1999). Sequence analysis shows that Yb\(^1\) is a missense mutation in Arg972 that is conserved in all Tudor domains so far studied and therefore is likely to be essential for Tudor domain function. The crystal structure of the Tudor domain in the survival of motor neuron (SMN) protein indicates that this domain likely functions as a protein–protein-binding domain (Selenko et al., 2001, Sprangers et al., 2003).

Yb\(^4\) and Yb\(^6\) are missense mutations at residues 747 and 858, respectively, located between the DEAH box RNA helicase domain and the Tudor domain. Using the polyclonal antibody against native Yb protein, we revealed that these proteins fail to localize to the Yb body (Fig. 6).

To further identify functionally important regions of Yb, we compared its sequences among several Drosophila species. We identified Yb homologues in Drosophila simulans, Drosophila yakuba, Drosophila erecta, Drosophila ananassae, Drosophila virilis, and Drosophila mojavensis. (Drosophila pseudoobscura genome data [release 1.0] did not contain a homologue because of a sequencing gap.) The Yb sequences of more closely related species, D. simulans and D. yakuba (2.6 myrs and 10 myrs separation from D. melanogaster), revealed a very strong conservation for the first N-terminal 150 aa, followed by a lower degree of conservation in short stretches of fewer than 10 aa residues, up to the RNA helicase domain (Fig. 6). The Yb proteins show very high homology in the last 470 residues at the C-terminal region. This pattern of homology corresponds well with our deletion/mutation analysis, suggesting that the N-terminal 150 aa residues may be important for hh expression in cap cells, whereas the C-terminal 470 residues may be essential for Yb protein localization via binding to protein partners.

Forbes et al., 1996). Because Yb-controlled hh expression is mostly responsible for follicle stem cell division (King et al., 2001), this raises the possibility that the N-terminal region of Yb is more responsible for regulating hh expression. However, transgenes with more C-terminal deletions show pinched and split egg chambers, often with very few nuclei, suggesting that follicle proliferation is improved, but germ line cells are underproliferated (Fig. 8). Therefore, we tested these transgenes for their ability to restore hh expression in cap cells or piwi expression in somatic niche cells. N-terminal deletions often showed no rescue of hh expression in cap cells (Figs. 6 and 7, C–E). All other transgenes tested restored hh expression in cap cells (Fig. 8, B and C). However, none of the deletion transgenes rescued piwi expression in somatic niche cells (Figs. 6 and 8, D and E). Therefore, the N-terminal region is required for hh expression and follicle cell proliferation, whereas the entire Yb protein is necessary for piwi expression and GSC self-renewal during oogenesis.

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regions responsible for hh- and piwi-mediated niche signaling and for subcellular localization as well as revealed its role in the self-renewal of male GSCs. These results expand our understanding of cellular and molecular mechanisms involved in niche signaling, which regulates GSC division in D. melanogaster, with implications for niche signaling in other stem cell systems.

Yb body as a novel organelle

Subcellular localization experiments revealed that Yb localizes to a new discrete electron-dense structure in somatic cells of the ovary, testis, and embryonic gonad that we call the Yb body. Immunofluorescence and immunoelectron microscopy reveals that the Yb body is distinct from known cytoplasmic organelles such as endosomes, lysosomes, mitochondria, the ER, nuages, sponge bodies, U bodies, and vesicular elements of the Golgi complex. Moreover, unlike nuages, sponge bodies, and U bodies, Yb bodies are only in the ovarian somatic cells, especially dense in the cap cells, which is consistent with the in situ Yb mRNA localization pattern, but not in the germline. However, the Yb bodies are not abundant in TF cells, in which Yb mRNA is also strongly detectable. This could reflect a differential regulation of Yb mRNA translation in TF and cap cells.

Yb body association region is distinct from the aforementioned sequences shown to be essential for hh expression in cap cells.

Discussion

In this paper, we have reported a novel and distinctive subcellular structure that localizes the Yb protein and defined Yb functional regions responsible for hh- and piwi-mediated niche signaling and for subcellular localization as well as revealed its role in the self-renewal of male GSCs. These results expand our understanding of cellular and molecular mechanisms involved in niche signaling, which regulates GSC division in D. melanogaster, with implications for niche signaling in other stem cell systems.

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The Yb body shares certain morphological and structural similarities with nuages, sponge bodies, and P bodies (also known as GW bodies in mammalian cells; Eystathioy et al., 2002), all of
The P bodies in yeast and mammalian cells (Eystathioy et al., 2002; Sheth and Parker, 2003) are granular, electron-dense bodies devoid of membranous structures, which are similar to the ultrastructural features of the Yb body. Immunoelectron microscopy further revealed that P bodies contain electron-dense fibrils (Yang et al., 2004). The ultrastructural similarities between nuage, the sponge body, the P body, and the Yb body all hint that the Yb body is involved in RNA-related processes.

The possibility that the Yb body functions in RNA-related processes is intriguing. Compartmentalization of mRNA and trans-acting proteins in controlling translation and stability of mRNAs is likely to be important in the D. melanogaster ovarian somatic cells (cap cells, previtellogenic cells, and postgermarial follicle cells), which must maintain a specific polarity in their contact and communication with neighboring germ cells. Indeed, the Yb body is frequently located near the germline-contacting side of the cytoplasm. It is possible that the Yb body represents a somatic counterpart to the germline-specific organelles such as nuage, sponge bodies, or Balbiani bodies. The Yb body may be involved in regulating the expression of the signaling machinery at the mRNA level. It will be interesting to identify additional protein and, possibly, RNA components of the Yb bodies and to determine

Figure 8. C-terminal region of Yb is required for PIWI but not HH expression in niche cells. (A) DAPI image of Yb72/Yb72 ovarioles carrying a transgene with a C-terminal deletion (pAS10). These ovarioles show improved follicle cell (FC) proliferation, implicating improved hh signaling, which is essential for follicle stem cell division (Farbes et al., 1996). (B and C) Consistent with this, the cap cell (CC) expression of hh is fully restored in Yb72/Yb72 mutants by introducing a C-terminally deleted Yb (pAS9 in B and pAS10 in C). This suggests that the C-terminal region of Yb is not required for HH expression. (D and E) A Yb72/Yb72 gerarium, carrying a pAS10 transgene and an myc-piwi gene, stained for DAPI (D) and myc-PIWI (E), respectively, is shown. The expression of piwi in TF and cap cells is not restored. This is also the case when any other mutant transgene in Fig. 6 is introduced into the Yb72/Yb72 mutants (see Fig. 6 for summary). Thus, the full-length Yb protein is required for piwi expression. GC, germline cell; NC, nurse cell. Bars, 20 µm.
and are involved in numerous processes. It has been proposed recently that most of these helicases release proteins from RNA rather than unwinding RNA (Fairman et al., 2004). However, the helicase homologous domain of the Yb protein contains no obvious DEAD box motif (King and Lin, 1999); thus, Yb is unlikely to be an RNA helicase but might still be involved in other RNA-related function. The Tudor domain is currently of unknown function but is present in several RNA-binding proteins (Ponting, 1997). However, based on the structure of the Tudor domain in the SMN protein, it is more likely to mediate protein–protein interaction (Selenko et al., 2001). The Tudor domain of the SMN protein binds to methylated RG tails of SMN proteins with particularly high affinity (Sprangers et al., 2003). Finally, it is rare to see proteins with a combination of a DEAD/DEAH box RNA helicase domain and a Tudor domain. In *D. melanogaster*, two novel and uncharacterized genes, CG31755 and CG11133, encode proteins with this domain combination and share moderate sequence similarity with Yb over most of their lengths. Our search has also revealed CG11133-like genes in *Pan troglodytes* and *Homo sapiens* but no Yb homologue outside of the *Drosophila* genus. In *D. melanogaster*, although the Homeless protein displays little sequence homology to the Yb protein, they share the same domain combination. A short part of the Tudor domain is conserved between Yb and Homeless, but the conserved sequence does include the Arg residue critical whether the composition of the Yb body varies in different types of cells and tissues.

The role of Yb in regulating male GSC self-renewal

The failure of the male GSC maintenance in the *Yb;G38* double mutant but not their corresponding single mutants indicates that Yb and a second gene affected by the *G38* mutation positively regulate the self-renewal of these GSCs in a compensatory manner. This finding provides an opportunity to explore Yb-mediated niche signaling mechanism in regulating male GSC self-renewal by cloning and systematically characterizing the gene disrupted by the *G38* mutation.

Functional domains of the Yb protein

Sequence homology searches revealed two conserved protein domains in Yb, a putative DEAD/DEAH box RNA helicase domain and a Tudor domain. These two domains are essential for Yb function because the pAS6 mutation in the Walker B motif of the helicase domain and the *Yb*1 mutation that mutates an extremely well conserved Arg residue in the Tudor domain to a Gln residue both result in loss of GSC maintenance and failure of the protein localization to the Yb body. The precise function of these domains cannot be predicted from sequence alone. DEAH box RNA helicases domains are found in many proteins and are involved in numerous processes. It has been proposed recently that most of these helicases release proteins from RNA rather than unwinding RNA (Fairman et al., 2004). However, the helicase homologous domain of the Yb protein contains no obvious DEAD box motif (King and Lin, 1999); thus, Yb is unlikely to be an RNA helicase but might still be involved in other RNA-related function. The Tudor domain is currently of unknown function but is present in several RNA-binding proteins (Ponting, 1997). However, based on the structure of the Tudor domain in the SMN protein, it is more likely to mediate protein–protein interaction (Selenko et al., 2001). The Tudor domain of the SMN protein binds to methylated RG tails of SMN proteins with particularly high affinity (Sprangers et al., 2003). Finally, it is rare to see proteins with a combination of a DEAD/DEAH box RNA helicase domain and a Tudor domain. In *D. melanogaster*, two novel and uncharacterized genes, CG31755 and CG11133, encode proteins with this domain combination and share moderate sequence similarity with Yb over most of their lengths. Our search has also revealed CG11133-like genes in *Pan troglodytes* and *Homo sapiens* but no Yb homologue outside of the *Drosophila* genus. In *D. melanogaster*, although the Homeless protein displays little sequence homology to the Yb protein, they share the same domain combination. A short part of the Tudor domain is conserved between Yb and Homeless, but the conserved sequence does include the Arg residue critical...
for function. Thus, even though Homeless is known to be involved in RNA localization and possibly translational regulation in the oocyte, considerable effort is needed to elucidate the biochemical activities of Yb.

Although sequence homology analysis has not revealed obvious biochemical activities for Yb, our deletional analysis has defined distinct regions of Yb that are responsible for its three different molecular functions: (1) activation of HH expression in somatic niche cells, (2) activation of PIWI expression in somatic niche cells, and (3) Yb body localization and/or organization. This analysis suggests that Yb is a functionally tripartite protein with the N-terminal half required for hh expression in cap cells, the C-terminal half required for Yb localization to Yb bodies, and the entire protein needed for piwi expression in somatic niche cells. The Yb body localization function requires the Tudor domain but not the helicase homology domain, yet the PIWI expression function requires both domains, including a conserved ATP/GTP-binding site in the helicase domain. Future studies, including the identification of protein interactors specific to different domains of Yb, should help decipher the biochemical functions of these domains as well as reveal other components of the Yb body.

Materials and methods

**Drosophila strains and culture**

All strains were grown at 25°C on yeast-containing molasses/agar medium. The following fly strains were used: The P{myc-piwi}/G38 insertion on the second chromosome, with an expression pattern representative of most P{myc-piwi} transgenic lines, was used to visualize Piwi protein localization [Cox et al., 2000]. w^118 served as the wild-type strain in all experiments unless indicated otherwise. Yb^3, Yb^6, and Yb^60 are previously described Yb point mutations (King and Lin, 1999), whereas Yb^27 is a truncation allele that causes a premature termination of the 1,042-aa Yb protein at residue 134 (King et al., 2001); all of these alleles are considered to be null (King and Lin, 1999; King et al., 2001). The hh-loz2 chromosome used is from the Bloomington BL-5530 sy50 sy[1] E[Fy(+)17] y[+] strain. loz2 staining was performed as previously described [Lin et al., 1994].

**Construction of transgenic flies**

To produce a His-Flag epitope fused to the Yb sequence, the following primer pair was purchased from Integrated DNA Technologies: the 5′-CTTGTCGTCGTCGTCCTTATAGTCAATAATGGT-3′ and the 3′-Yb Flag primer (5′-CTTGTCGTCGTCGTCCTTATAGTCCATGCTGCG-3′) and the 3′-Yb Flag primer (5′-CTTGTCGTCGTCGTCCTTATAGTCCATGCTGCG-3′). These primers were phosphorylated, mixed, and filled in by PCR reaction. PCR product of the correct length was gel purified, digested with Nhel–Mhel, and inserted into the pJ35 ΔXhol plasmid. pJ35 ΔXhol contains only the N-terminal sequence of Yb and allows the use of the first Mhel restriction site right after the ATG codon. The resulting His-Flag-tagged 5′ Yb fragment was digested with Nhel–BsiWI and used to replace the corresponding sequence in pCa4Stu (genomic Yb rescue construct; King and Lin, 1999), resulting in a P-element transformation vector containing a genomic Yb rescue fragment with a His-Flag sequence after the ATG start codon. This construct also completely rescued the phenotype of the Yb^27 mutant.

Transgenes in P-element transformation vectors were introduced into w^118 flies via germline transformation (Rubin and Spradling, 1982). The transgenes were injected at the Duke University Model System Genomics Facility (http://www.biology.duke.edu/model-system).

**Staining in RNA in whole-mount ovaries**

Ovaries were dissected and fixed for 10 min or 2 h in prechilled methanol at −20°C (no difference was observed between these two Fixing times) followed by several washes in 1× PBS. Labeling was performed with SYTO RNAselect green fluorescent cell stain (Invitrogen). All of the following procedures were performed in the dark. Ovaries were incubated for 20 min at RT in a 500-nM solution of the dye in 1× PBS followed by several washes in PBS. Counterstaining with Yb antibody was subsequently performed as described in the next section (beginning with the blocking step).

**Antibody production and affinity purification**

MAP–8–conjugated peptides of [H-KENPTPSSEDYKAEA] 8× MAP (Yb pep3) and [H-DITLSDSRTYKDSPA] 8× MAP (Yb pep4) were purchased from Invitrogen. These peptides correspond to amino acid residues 187–203 and 322–338 of Yb, respectively. The peptides were each injected into one rabbit by Cocalico Biologicals, Inc. according to standard protocol. Serum was obtained from rabbits immunized with peptides Yb pep3 [serum DU159] and Yb pep4 [serum DU160]. IgG was isolated from both sera by purification on a protein A-Sepharose column (GE Healthcare). Both IgG fractions were further purified through affinity matrix and prepared by cross-linking peptides Yb pep3 and Yb pep4 to Reacti-Gel 6× (Thermo Fisher Scientific). For the cross-linking, 23 mg of peptide Yb pep3 and 10 mg of peptide Yb pep4 were separately incubated with 1 ml of activated resin in 0.1 M sodium carbonate buffer, pH 10, for 24 h at 4°C. The reaction was stopped by incubation with 1 M methanolamine, pH 10.0, for 24 h. After extensive washing, peptide columns were loaded with the protein A–purified IgG and incubated overnight at 4°C. The columns were washed extensively and eluted with 0.05 M of Gly buffer, pH 2.5, containing 0.15 M NaCl. Fractions were immediately neutralized with 0.1 V 1 M Tris-HCl, pH 8.0, pooled, and dialyzed against a buffer containing 0.025 M Hepes-KOH, pH 7.5, and 0.15 M KCl. The purified antibodies were aliquoted, quick-frozen in liquid nitrogen, and stored at −80°C.

**Immunohistochemistry**

Ovaries and testes were dissected, fixed, and stained as previously described [Lin et al., 1994]. For immunofluorescence staining, the following antisera were used: polyclonal rabbit anti-Vasa (1:2,000; Hay et al., 1990), monoclonal mouse anti-181 (which recognizes spectrosomes and fusomes; 1:1; Zaccai and Lipshitz, 1996), polyclonal rabbit anti-a-spectrin (1:200; Byers et al., 1987), monoclonal mouse anti-MYC epitope 1PE10.2 (1:50; Even et al., 1989), monoclonal mouse anti-Flag M2 (1:1,000; Sigma-Aldrich), polyclonal rabbit anti-Flag F7425 (1:500; Sigma-Aldrich), monoclonal
mouse anti-Ma31B (1:400; provided by A. Nakamura, RIKEN Center of Developmental Biology, Kobe, Japan; Nakamura et al., 2001), polyclonal rabbit anti-Flag (1:750; provided by J. Wilhelm, University of California, San Diego, La Jolla, CA; Wilhelm et al., 2005), polyclonal rabbit dlsm11 (1:3,000; provided by J.G. Gall, Carnegie Institute of Washington, Baltimore, MD; Liu and Gall, 2007), and polyclonal rabbit Traps (1:200; provided by C. Nicchitta, Duke University, Durham, NC; Nicchitta et al., 1991). Affinity-purified rabbit anti-Yb pep3 antibody was used at 1:250 dilution. All of the fluorescence-conjugated secondary antibodies were purchased from Jackson ImmunoResearch Laboratories and used at 1:200 dilution except for Alexa Fluor 488 (Invitrogen), which was used at 1:500 dilution. Immunofluorescently labeled samples were also counterstained with DAPI as described previously (Lin and Spradling, 1993). Photomicrographs were taken using either a microscope (Axioskop; Carl Zeiss, Inc.) with the image captured by a cooled charged-coupled device camera (Star-1; Photometrics) or a confocal microscope (LSM510; Carl Zeiss, Inc.) using its built-in camera. Both microscopes used a 10× NA 0.3 Plan-Neofluar or a 40× NA 1.3 Plan-Neofluor oil differential interference contrast objective for low, medium, and high magnification imaging, respectively. All images were processed using Photoshop (Adobe).

Transmission electron microscopy

The conventional transmission microscopy of wild-type germaria was conducted according to Lin et al. (1994) or Reedy and Beall (1993). Images were taken using an electronic microscope (EM420; Philips) in the Department of Cell Biology at Duke University.

Immunoelectron microscopy

Ovaries were dissected from 5× Flag–Yb transgenic flies, fixed, and stained with mouse monoclonal anti-Flag M2 antibodies for immunocytochemistry as previously described (Lin et al., 1994). Anti–mouse Nanogold and Alexa Fluor 488 double-conjugated secondary antibodies were used for secondary labeling (1:3). Ovaries were washed, lightly cross-linked with 0.1% glutaraldehyde and aldehyde quenched with 50 mM Gly in PBS, according to the protocol from Nanoprobes, Inc. Several ovaries were directly inspected for their valuable suggestions on the manuscript, Dr. Jochen Genschel for help with Yb antibody purification, Dr. Zhong Wang for help in BLAST searching sequence assembly, and Dr. Heather Weggoh for help in Western blot analysis of the Yb mutant proteins. We also thank Drs. A. Nakamura, J. Wilhelm, J.G. Gall, and C. Nicchitta for providing Me31B, Tral, dlsm11, and Traps antibodies, respectively.

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