Phosphatidylinositol 4,5-bisphosphate Directs Spermatid Cell Polarity and Exocyst Localization in *Drosophila*

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During spermiogenesis, *Drosophila melanogaster* spermatids coordinate their elongation in interconnected cysts that become highly polarized, with nuclei localizing to one end and sperm tail growth occurring at the other. Remarkably little is known about the signals that drive spermatid polarity and elongation. Here we identify phosphoinositides as critical regulators of these processes. Reduction of plasma membrane phosphatidylinositol 4,5-bisphosphate (PIP₂) by low-level expression of the PIP₂ phosphatase SigD or mutation of the PIP₂ biosynthetic enzyme Skittles (Sktl) results in dramatic defects in spermatid cysts, which become bipolar and fail to fully elongate. Defects in polarity are evident from the earliest stages of elongation, indicating that phosphoinositides are required for establishment of polarity. Sktl and PIP₂ localize to the growing end of the cysts together with the exocyst complex. Strikingly, the exocyst becomes completely delocalized when PIP₂ levels are reduced, and overexpression of Sktl restores exocyst localization and spermatid cyst polarity. Moreover, the exocyst is required for polarity, as partial loss of function of the exocyst subunit Sec8 results in bipolar cysts. Our data are consistent with a mechanism in which localized synthesis of PIP₂ recruits the exocyst to polarity. Moreover, the exocyst is required for polarity, as partial loss of function of the exocyst subunit Sec8 results in bipolar cysts. Our data are consistent with a mechanism in which localized synthesis of PIP₂ recruits the exocyst to promote targeted membrane delivery and polarization of the elongating cysts.

INTRODUCTION

Cell polarity is a prerequisite for differentiation, proliferation and morphogenesis in all organisms. The activity and localization of the cytoskeleton, polarity complexes, signaling networks, and membrane trafficking associated with polarization are regulated by the lipid composition of organelles and membrane domains, in particular by phosphorylated derivatives of phosphatidylinositol (PI), also called phosphoinositides, which function as spatially restricted molecular signals (Odorizzi et al., 2000; Behnia and Munro, 2005; Di Paolo and De Camilli, 2006). Phosphoinositides are membrane-tethered lipid molecules synthesized from PI by the sequential action of lipid kinases (Figure 1A). Local concentrations of particular phosphoinositides are controlled by the subcellular localization or activation of lipid kinases, lipases, and phosphatases (Roth, 2004). For example, plasma membrane PI 4,5-bisphosphate (PIP₂) is synthesized from PI 4-phosphate (PIP₄) by PI4P 5-kinases (PI4Ks) and can in turn be phosphorylated by PI 3-kinases (PI3Ks) to yield PI 3,4,5-trisphosphate (PIP₃). PIP₂ can also be hydrolyzed by phospholipase C (PLC) to produce the second messengers inositol trisphosphate (IP₃) and diacylglycerol (DAG) or dephosphorylated by phosphoinositide phosphatases to regenerate PI4P. Importantly, in addition to serving as a precursor for other signaling molecules, PIP₂ itself is thought to act as a potent membrane signal (Di Paolo and De Camilli, 2006).

PIP₂ regulates diverse targets, including actin regulators and membrane-trafficking proteins (Takenawa and Itoh, 2001; Yin and Janmey, 2003; De Matteis and Gоди, 2004; Behnia and Munro, 2005; Niggli, 2005; Balla, 2006; Oude Weernink et al., 2007). High levels of PIP₂ promote filamentous (F) actin assembly by directly binding and regulating proteins that control actin polymerization (Yin and Janmey, 2003). PIP₂ links F-actin to the plasma membrane via membrane-cytoskeleton cross-linkers such as spectrin and moesin (Niggli et al., 1995; Hirao et al., 1996). Subunits of the exocyst—a conserved octameric protein complex that directs localized membrane addition—contain basic regions that interact with PIP₂ (He et al., 2007; Liu et al., 2007; Zhang et al., 2008; Yamashita et al., 2010). The exocyst directs membrane trafficking events required for cell polarization and growth during development in plants and animals and was recently shown to be required for ciliary membrane formation in Madin-Darby canine kidney (MDCK) cells (Hsu et al., 2004; Somers and Chia, 2005; Hala et al., 2008; Zuo et al., 2009). However, the role of PIP₂ binding in exocyst function during development of a complex multicellular organism remains unknown.

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To study the role of phosphoinositides in cell polarity, we are using *Drosophila* sperm development as a model system. Spermatogenesis in *Drosophila* initiates by division of a germline stem cell to form a gonial cell, which then divides four times to produce a group of 16 primary spermatocytes. The spermatocytes grow in size and undergo meiosis to generate a group of 64 haploid spermatids. Because germ cell cytokinesis is incomplete, groups of cells remain connected through stable intercellular bridges called ring canals and develop within a syncytial cytoplasm or cyst. Male germline ring canals contain the actin-associated protein anillin and associate with the fusome, a spectrin-rich membrane organelle that interconnects the developing germ cells and localizes to the growing end of the sperm tails in a honeycomb structure (Hime *et al.*, 1996).

*Drosophila* spermatids undergo dramatic changes in shape as they develop from small round cells—12 μm in diameter to elongated cells nearly 1.8 mm in length (Figure 1B; Tates, 1971; Tokuyasu, 1975). In early round spermatids, the basal body forms a stable attachment to the nuclear envelope (Figure 1Bii). The microtubule-based flagellar axoneme grows out from the basal body, and two mitochondrial derivatives elongate along the length of the axoneme (Figure 1B, i–iii). The distal (growing) end of the elongating axoneme is covered by a flagellar membrane—topologically equivalent to a ciliary membrane—that remains contiguous with the plasma membrane throughout the axoneme assembly (Figure 1Biv; Fritz-Niggli and Suda, 1972; Tokuyasu, 1975). During elongation, each cyst becomes highly polarized, with nuclei localizing to one end and the fusome and ring canals localizing to the growing end (Hime *et al.*, 1996; Ghosh-Roy *et al.*, 2004). Spermatid elongation and individualization of mature sperm require a substantial increase in cell surface area. Hence, membrane addition is a critical aspect of sperm maturation.

Here, we show that polarized organization of elongating spermatids within cysts depends on normal levels of Pip2. When Pip2 levels are reduced, nuclei localize to both ends of the cysts and sperm tails grow toward the middle. The exocyst, which normally colocalizes with Pip2, at the growing end, is uniformly distributed when Pip2 levels are reduced, suggesting a defect in targeted membrane delivery. We demonstrate that both the Pip5K Skt1 and the exocyst are required for normal polarization and elongation of spermatid cysts. Our results suggest that local synthesis of PIP2 is predicted to produce a truncated protein of 581 rather than 672 amino acids (Blankenship *et al.*, 2007). Evidence that funnel cakes (fun) encodes *Drosophila* Sec8 will be presented elsewhere (T. J. Blankenship, M. Giansanti, C. Robinett, M. Gatti, and M. T. Fuller, unpublished data). Both mutants were examined in trans to deficiencies that uncover the corresponding genes (Df(3R)Esp13 in the case of *fun* and Df(3R)Exel6145 in the case of *skt1*). β-Tubulin-GFP (*Inoue et al.*, 2006) was used as a germ cell marker in all experiments (Tate, 2008; Blankenship *et al.*, 2010). α-tubulin (6-11-B) antibodies, which were generously provided by James Baker and Maurice Kernan (SUNY Stony Brook, NY), skt1−/− is a hypomorphic allele of *skt1* that affects polarity during *Drosophila* oogenesis (Gervais *et al.*, 2008). Clones of germ cells mutant for *skt1* were generated in flies containing *hspFLP/Y, FRT42B, skt1−/−FRT42B, and Ubi-GFP* lines using the FLP-FRT (fragment length polymorphism and recognition target system, respectively; Golic and Lindquist, 1989). hs-FLP and FRT42B, Ubi-GFP* lines were obtained from the Bloomington *Drosophila* Stock Center (Bloomington, IN). Complete genotypes of fly stocks are available upon request.

**Fluorescence Microscopy**

For live preparations of *Drosophila* male germ cells, testes from newly eclosed males were dissected in testis isolation buffer (Casal *et al.*, 1990) containing 8.3 μg/ml Hoechst 33342 (Sigma-Aldrich, St. Louis, MO) to stain DNA and squashed under a coverslip, as previously described (Wei *et al.*, 2008). Spermatid cyst length was determined from phase-contrast micrographs of fixed preparations obtained from 1-d-old adult males. Germ cells were stained for F-actin and DNA (see below) to mark the ends of the cysts and to determine the stage of cyst elongation. Full-length wild-type cysts were identified by the presence of actin-containing investment cones. Skt1−/− cysts were measured at the latest identified stages, based on the shape and distribution of the nuclei. Because cysts were often bent, segmental measurements were obtained using Axiovision software (Carl Zeiss, Germany) and then summed to determine the total length. Average length and SD were calculated using Microsoft Excel (Microsoft, Redmond, WA).

Sample preparation for fluorescent staining was performed as described (Wei *et al.*, 2008). Primary antibodies used were used at the following concentrations: 1:250 mouse IgM anti-Pip2 (Echelon Biosciences, Logan, UT); 1:500 guinea pig anti-Sec7 and 1:1000 guinea pig anti-Sec8 (gifts from Slobodan Bersonja and Ulrich Tepass [University of Toronto, Toronto, Canada]; Bersonja *et al.*, 2004); 1:1000 mouse anti-acetylated-α-tubulin (6-11B, Sigma-Aldrich); 1:500 mouse anti-phospho-moesin (p-moesin; gift from Sebastien Carreno [Institute for Research in Immunology and Cancer, Montreal, Canada]; Carreno *et al.*, 2008); 1:100 rabbit anti-anillin (from Christine Field [Harvard Medical School, Boston, MA]) or from our lab; 1:1000 rabbit anti-Sec8 (gifts from Slobodan Bersonja and Ulrich Tepass [University of Toronto, Toronto, Canada]; Bersonja *et al.*, 2004); 1:500 mouse anti-GFP (Molecular Probes, Eugene, OR); F-Antin was stained with 1 U/ml rhodamine phalloidin or Alexa 488 phalloidin, as recommended by the manufacturer (Molecular Probes). Secondary antibodies were conjugated to Alexa fluorochromes (Molecular Probes) and were used at 1:1000. DAPI (5 μg/ml; Molecular Probes) was used to stain DNA.

Preparations were examined on an upright Zeiss Axioplan 2 epifluorescence microscope equipped with an Axiocom black and white camera using Axiovision software (Carl Zeiss). Confocal images were obtained on an inverted optical microscope with a Zeiss LSM510 confocal microscope (Zeiss). Images were obtained using Axiovision software (Carl Zeiss, Oberkochen, Germany) or on an inverted Zeiss LSM510 laser-scanning confocal microscope using LSM Image software (The Hospital for Sick Children Imaging Facility). Unless indicated, groups of images were adjusted in an identical manner for brightness, contrast and pseudocolor with Adobe Photoshop (Adobe Systems, San Jose, CA).

**Immunoblotting**

Samples for immunoblotting were prepared using 20 pairs of testes per lane for each genotype. Testes from newly eclosed males were dissected in testis isolation buffer and boiled for 5 min in Laemmli sample buffer (Sambrook *et al.*, 1989). Proteins were separated in a 10% SDS-polyacrylamide gel and transferred to a nitrocellulose membrane (Akersham, CE Healthcare, Little Chalfont, Buckinghamshire, United Kingdom) using a Trans-Blot Semi-Dry transfer apparatus (Bio-Rad, Mississauga, ON, Canada). Blots were probed sequentially with mouse secondary antibodies (Amersham) were diluted 1:10,000 and visualized using chemiluminescence (ECL Plus kit, Amersham). The distribution of the nuclei. Because cysts were often bent, segmental measurements were obtained using Axiovision software (Carl Zeiss, Germany) and then summed to determine the total length. Average length and SD were calculated using Microsoft Excel (Microsoft, Redmond, WA).

**Electron Microscopy**

Testes samples for transmission electron microscopy (TEM) were prepared as described (Bazinet and Rollins, 2003; Wei *et al.*, 2008). Samples for immunoelectron microscopy were prepared using a protocol modified from Peters *et al.* (2006). Briefly, *Drosophila* testes were dissected in phosphate buffer, pH 7.4, placed immediately in 1% permagnate, and fixed overnight at 4°C. The samples were rinses twice with 1.5 M glycine in phosphate-buffered saline (PBS) for 5 min and then taken through serial dilutions of 1, 5, and 12% gelatin (Electron Microscopy Sciences, Hatfield, PA), for 10 min each. Testes were

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**MATERIALS AND METHODS**

**Fly Stocks**

Flies were raised and maintained on standard cornmeal molasses agar at 25°C. Transgenic flies were generated by injection of w1118 balancer FM7i, which carries a

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transferred to a warm Chang mold for flat embedding (Electron Microscopy Sciences) and covered with warm 12% gelatin. Molds were placed at 4°C for 30 min. Regions of interest were excised from the gelatin blocks, transferred to vials with PBS (without Ca\(^{2+}\) and Mg\(^{2+}\)) for 10 min, and then placed in a 2.3 M sucrose solution in PBS (without Ca\(^{2+}\) and Mg\(^{2+}\)) overnight at 4°C. Individual testes were cut from the gelatin blocks, transferred to color-marked pins, and then immersed in liquid nitrogen. Pins were stored in liquid nitrogen before sectioning with a Leica Ultracut UCT microtome with an FCS cryochamber (Leica Microsystems, Wetzlar, Germany). Cryosections (80 nm thick) were blocked for 30 min with 5% cold water fish gelatin, incubated with 1:100 rabbit anti-GFP for 30 min, rinsed five times for 2 min with PBS, fixed 10 min with 2.5% glutaraldehyde, rinsed two times for 2 min with PBS followed by five times for 2 min with distilled water, and finally stained for 10 min with 1:100 rabbit anti-GFP for 30 min, rinsed five times for 2 min with PBS, and with 2.5% glutaraldehyde, rinsed two times for 2 min with PBS followed by five times for 2 min with distilled water, and finally stained for 10 min with 2% methylcellulose and 0.4% uranylacetate.

Sections were viewed with a JEOL JEM 1200EX TEM, a JEOL JTE 141011 (JEOL, Peabody, MA; The Hospital for Sick Children Electron Microscopy Facility), or a Tecnai TEM (Tecnai, Hillsboro, OR; Advanced Bioimaging Centre at The Hospital for Sick Children and Mt. Sinai Hospital). Images were obtained using AMTv542 (Advanced Microscopy Techniques, Danvers, MA) and Gatan Digital Micrograph (Gatan, Pleasanton, CA) acquisition software and were manipulated only for brightness and contrast using Adobe Photoshop.

**In Vitro Culture of Elongating Spermatids**

Isolated cysts of *Drosophila* spermatids were cultured as described (Noguchi and Miller, 2003). Briefly, testes from newly eclosed wild-type or SigD-low adult males were dissected and transferred into MM3 culture medium containing 10% fetal calf serum (Sigma-Aldrich) and standard penicillin/streptomycin cocktail. Time-lapse differential interference contrast (DIC) videos were recorded from onion stage to early elongation stages. Elongation was recorded using a DeltaVision system (Applied Precision, Issaquah, WA) on a IX70 (Olympus, Tokyo, Japan) microscope equipped with 40x UPlanApo objective (Olympus). QuickTime videos (10 fps) were created by ImageJ (http://rsb.info.nih.gov/ij/) and compressed by MPEG-4.

**RESULTS**

**Reduction of Plasma Membrane PIP\(_2\) Causes Male Sterility**

During experiments to test the roles of phosphoinositides during *Drosophila* spermat development, we generated lines of transgenic flies expressing the *Salmonella* PIP\(_2\) phosphatase SigD (also called SopB) under control of the spermatocyte-specific \(\beta\)-tubulin promoter (see Materials and Methods). Two of the SigD-expressing lines (denoted R1 and R7) had superficially normal testes that contained elongated spermatid cysts (Figure 1C). R1 and R7, unlike the other SigD-expressing lines with severe phenotypes, carried transgenes on the X chromosome and expressed lower levels of SigD (Figure 1D; R7, not shown). We refer to these lines as SigD-low to distinguish them from the previously described lines, which we call SigD-high. Because R1 and R7 behaved identically, for the purposes of this study we focused on R1.

Expression of low levels of SigD caused a significant reduction in plasma membrane PIP\(_2\), as visualized by localization of fluorescent markers that specifically bind PIP\(_2\). In wild-type spermatocytes, early spermatids and elongating spermatids (Figure 2, A' and D'), PLC\(_6\)-PH-GFP, a green fluorescent marker that specifically binds PIP\(_2\), was mainly associated with the plasma membrane, whereas in SigD-low, PLC\(_6\)-PH-GFP was cytoplasmic, with low or undetectable levels on the plasma membrane (Figure 2, B' and E').

There was no obvious difference in the distribution of RFP-PH-FAPP, a red fluorescent PI4P-binding protein, in SigD-low versus wild-type cells, suggesting that levels of PI4P were not significantly affected (not shown). However, we cannot rule out a subtle increase in PI4P below the level of detection with the fluorescent marker. Overexpression of the PIP5K Sktll (Sktll)—which synthesizes PIP\(_2\) from PI4P—suppressed loss of plasma membrane PIP\(_2\) in the context of SigD-low; PLC\(_6\)-PH-RFP was restored to the plasma membrane in spermatocytes and early spermatids (Figure 2C') and partially restored in elongated spermatids (Figure 2F').

**PIP\(_2\) Is Required for Spermatid Cyst Polarity and Elongation**

Analysis of the morphology of developing germ cells in SigD-low males revealed defects in spermatid cyst elongation and polarity. Wild-type cysts of elongating spermatids were 1.75 ± 0.19 mm long (n = 17) and unipolar, with nuclei clustered at one end of the cyst (Figure 3, A, E, I, M, and O).
In contrast, early elongating cysts from SigD-low males showed scattered nuclei (Figure 3N) and later stage SigD-low cysts were much shorter than wild-type cysts of comparable stage (0.91 ± 0.09 mm long; n = 24) and appeared bipolar, with nuclei distributed between the two ends (Figure 3, B and F) and sperm tails growing toward the middle of the cyst (Figure 3, B, F, J, and P). Bipolarity of SigD-low cysts was partially rescued by coexpression of Sktl (Figure 3, B and F) and sperm tails growing toward the middle (Figure 3, B, F). Coexpression of Sktl with SigD-low partially restores plasma membrane PIP$_2$ in early bipolar cysts (Figure 3, A and C). PIP$_2$ is found at the plasma membrane of wild-type early round spermatids (A') and elongating spermatids (D'). Low levels of SigD reduce PIP$_2$ at the plasma membrane (B') and cause accumulation of PLC$_6$-PH-GFP in the cytoplasm (B' and E'). Coexpression of Sktl with SigD-low partially restores plasma membrane PIP$_2$ in early (arrowhead, C') and elongating (E') spermatids. Scale bars, 10 μm.

**PIP$_2$ Reduction Disrupts Polarity during Early Stages of Elongation**

Proper levels of PIP$_2$ are required at the earliest stages of elongation to ensure correct polarized orientation of individual elongating spermatids, as revealed by real-time imaging of wild-type and SigD-low spermatids cultured on polylysine-coated glass dishes (Figure 4A). In these experiments, groups of interconnected spermatids remained attached by ring canals at the elongating end, which adhered to the dish. Thus, the nuclear ends of the individual spermatids appeared to grow outward, away from each cluster of cells. In wild-type spermatids, polarity was established early (Figure 4A, top panels and diagrams; see also Supplemental Video 1). Each nucleus-basal body pair became associated with one end of the elongating mitochondrial derivative, the nucleus appeared to form a tight association with the plasma membrane, and all sperm tails elongated in the same direction (Figure 4A, t = 05:26, red arrows). In SigD-low cysts, the relationship of the nucleus with the mitochondrial derivative was more fluid (Figure 4A, bottom panels and diagrams; see also Supplemental Video 2). The nucleus-basal body pair appeared to move freely along the length of the early elongating spermatid, the nucleus failed to associate tightly with the plasma membrane, and different spermatids within a group elongated in different directions (Figure 4A, S = 07:02, red arrows).

**Basal Body Orientation Requires PIP$_2$**

Normal levels of PIP$_2$ are required for basal body docking and proper orientation of the nucleus and basal body with respect to the direction of growth of the spermatid cyst. Analysis of the basal body marker Unc-GFP and the centrosomal protein Cnn revealed that in wild-type spermatids Unc-GFP and Cnn were associated with a single basal body per nucleus (Figure 4, B and C). All nucleus-basal body pairs were oriented in the same direction, with the nuclei adjacent to the plasma membrane and basal bodies oriented away from the membrane (Figure 4, B and C, and inset in B). In SigD-low spermatids, Unc-GFP and Cnn were generally present at the expected position of the basal body (Figure 4, E and F), and most basal bodies were near a nucleus. However, nucleus-basal body pairs were oriented in different directions, with some of the basal bodies found next to the plasma membrane (Figure 4, E and F, and inset in E). Immunoelectron microscopy showed that Unc-GFP, which localized to the basal body in wild type (Figure 4D, inset), was properly localized in SigD-low, but its levels were reduced on PIP$_2$ levels. As previously reported, spectrin showed a polarized localization to the fusome in early stages of elongation (not shown) and, in later stages, to a honeycomb-like structure along the membranes at the growing end of wild-type cysts (Figure 3, I and L). In SigD-low cysts, the honeycomb structure failed to form and a fusome-like spectrin structure was observed in the middle of the bipolar cysts (Figure 3J). The polarized distribution of spectrin was largely restored by coexpression of Sktl (Figure 3K). In wild-type cysts, the highest concentration of actin filaments was found at the growing end (Figure 3, M and O), near the ring canals, which contain anillin (Figure 3M). P-moesin also localized to ring canals (Figure 3O). SigD-low cysts had fewer actin filaments, and these were fairly evenly distributed throughout the cysts (Figure 3, N and P). Anillin and P-moesin remained associated with ring canals, but these were scattered at early stages of elongation (Figure 3N; P-moesin, not shown) and relocalized to the middle of the bipolar cysts at later stages (Figure 3P; anillin, not shown).
In a few cases, basal bodies of apparently normal architecture were found in the absence of any obvious nuclear envelope (Figure 4G), similar to what we had observed for SigD-high.

**Sktl Localizes to the Growing End of Spermatid Cysts**

Sktl becomes concentrated at the growing end of spermatid cysts, as determined by examining distribution of YFP-Sktl in wild-type spermatids. In early round spermatids, Sktl was evenly distributed along the plasma membrane (Figure 5A) and was concentrated at ring canals, where it colocalized with anillin (not shown). In early elongating spermatids, Sktl was found along the plasma membrane and the growing end (Figure 5C, arrow) on a linear structure of the expected length and morphology to be the flagellar membrane (Figure 5I, white arrowhead). By mid-elongation, Sktl was enriched in punctate and elongated structures (Figure 5, E and J, white arrowhead), many of which localized near ring canals at the growing end (Figure 5, I’ and J’; arrowheads, and insets in I’ and J’). At later stages, Sktl became concentrated on membranes near the growing end (Figure 5G). In contrast, YFP-Sktl failed to strongly concentrate at the growing end in SigD-low spermatids (Figure 5, D, F, and H, arrows).

**PIP2 Colocalizes with and Recruits the Exocyst to the Growing End of Spermatid Cysts**

The exocyst, like PIP2 and Sktl, was concentrated along membranes at the growing end of spermatid cysts, as revealed by immunofluorescence using antibodies that specifically recognize the exocyst subunits Sec8 and Sec6. Although the exocyst was present in cytoplasmic puncta that showed no obvious asymmetry in early round and elongating spermatids (not shown), at later stages of elongation the exocyst exhibited a highly polarized distribution. Sec8 and Sec6 were found in a honeycomb-like structure at the growing end of the spermatid cysts (Figure 6, B, C, and F, and insets), with PIP2 appearing to surround the exocyst patches (Figure 6, E and C). In SigD-low spermatids, the localization of Sec8 and Sec6 was dramatically affected, with the two proteins showing a diffuse distribution throughout the entire cyst (Figure 6, D and G, and inset in G). Localization of
Sec8 and PIP₂ to the growing end was largely restored by coexpression of Sktl with SigD-low (Figure 6, E and H), suggesting that PIP₂ recruits the exocyst to membranes near the sites of flagellar axoneme assembly and membrane addition.

**The Exocyst Is Required for Spermatid Cyst Polarity during Elongation**

Loss-of-function mutations in genes encoding exocyst subunits cause defects in spermatid cyst elongation and polarity reminiscent of the defects caused by forced expression of SigD. The **onr** gene encodes the *Drosophila* homolog of Exo84, and the **fun** gene encodes *Drosophila* Sec8. Testes from **onr**/Df flies revealed severe defects in spermatid elongation, with round cysts (Figure 7A) that completely lacked polarity (Figure 7, C and E), similar to the phenotype of SigD-high spermatids. Testes from **fun**/Df flies showed milder defects in spermatid elongation (Figure 7B) and had bipolar cysts (Figure 7, D and F), similar to the phenotype of SigD-low (Figure 3B). Consistent with **fun** being a partial loss-of-function allele, mutant testes showed reduced Sec8 staining, with no accumulation at the growing end of the cysts (Figure 7G). Sec6 was delocalized, but its levels were not visibly reduced in **fun**/Df testes (Figure 7H). The cytoskeletal proteins actin, anillin and spectrin did not redistribute to the middle of the bipolar cysts, but accumulated at the growing end of the cysts in **fun**/Df testes (Figure 7, I and J), in contrast to the mislocalization to the middle of the cysts observed in SigD-low testes. This difference raised the possibility that the defects in spermatid cyst polarity observed in exocyst mutants might be due to a primary defect in localized membrane trafficking rather than to secondary effects on cytoskeletal proteins. Ultrastructural analysis by TEM confirmed that both SigD-low and **fun** spermatid cysts are indeed bipolar, with axonemes growing from opposite directions. Mature wild-type axonemes contain nine doublet and two central-pair microtubules and are decorated with accessory microtubules and flagellar dyneins that confer rotational polarity to the axonemes. In wild-type cysts, all axonemes were oriented in the same direction (Figure 7K, curved red arrows), whereas axonemes in SigD-low or **fun** cysts were antiparallel (Figure 7, L and M, curved red arrows).

**PIP₂ and the Exocyst Are Required for Plasma Membrane Addition during Spermatid Elongation**

Ultrastructural analysis by TEM revealed defects in the plasma membrane in both SigD-low and **fun** spermatids, suggesting that reduction of PIP₂ by SigD-low and reduced function of Sec8 in **fun** mutants cause defects in membrane deposition during elongation. In wild type, each spermatid was surrounded by a plasma membrane that separated it from adjacent spermatids (Figure 7K). In contrast, the plasma membranes separating individual spermatids were largely absent in SigD-low and **fun**/Df spermatid cysts (Figure 7, L and M), indicating a defect in membrane addition during spermatid elongation.

**DISCUSSION**

We provide evidence for a critical role for PIP₂ as a regulator of spermatid cyst polarization during spermio-
Figure 5. Sktl localizes to the growing end of spermatid cysts. (A–H) Epifluorescence micrographs showing distribution of YFP-Sktl in wild type (A, C, E, and G) and SigD-low (B, D, F, and H). Arrows indicate the growing ends of spermatid cysts. In wild-type early round spermatids (A), Sktl is evenly distributed along the plasma membrane (also shown in I). In wild-type early elongating spermatids (C), Sktl localizes along the plasma membrane and at the growing end (arrow and inset). In wild-type spermatids at midstages of elongation (E), Sktl is enriched in few focal spots along the length of the spermatid membrane and is concentrated at the growing end (inset; also shown in J). In wild-type late elongated cysts (G), Sktl is concentrated near the growing end. In early SigD-low cysts (B), Sktl localization at the plasma membrane appears normal. In contrast, in elongating SigD-low cysts (D, F, and H), Sktl fails to strongly concentrate at the growing end. (I–J) Epifluorescence micrographs showing localization of YFP-Sktl (grayscale and green), anillin in the ring canals (Anil, red) and DNA (blue) in wild-type spermatids. Dur- ing early stages of elongation (I′–I′), Sktl is found all along the plasma membrane and on the flagellar membrane (I′ and I′; insets in I′ and I′; white open arrowheads), which is located near the ring canals (I′ and I′; insets in I′ and I′; white open arrowheads). Note that the ring canals in this cell are viewed end-on and appear as linear rather than circular structures. In later stages (J–J′), anillin remains in ring canals at the growing end (J′ and J′; black open arrowheads; insets in J′ and J′), whereas Sktl is enriched in regions adjacent to ring canals (J′ and J′; white arrowheads; insets in J and J′). Scale bars, 10 μm.

The factors that regulate localization of PIP5Ks are not well understood. In C. elegans, concentration of the Sktl homolog PPK-1 at the posterior pole of early embryos is dependent on casein kinase 1, which negatively regulates its distribution. In mammalian tissue culture cells, plasma membrane association of PIP5Kα is regulated by Rho and Rac (Chatah and Abrams, 2001). In neurons, mammalian PIP5K1y661 is recruited and activated by binding to the focal adhesion protein talin and the clathrin adaptor AP-2 (Di Paolo et al., 2002; Nakano-Kobayashi et al., 2007). Our finding that Sktl localization is abnormal in SigD-low cysts suggests the existence of a positive feedback loop, whereby locally high concentrations of PIP2 retain the PIP5K, which in turn stimulates local PIP2 synthesis. Such a feedback loop could operate by maintaining locally activated Rho family G proteins, which—together with their activators—bind PIP2 (Heo et al., 2006; Yeung et al., 2008; Yoshida et al., 2009) and which in turn could recruit PIP5K (Yang et al., 2004). Alternatively, the feedback mechanism could be more direct, because PIP5Ks were recently shown to localize to the plasma membrane via positively charged amino acids that bind phosphoinositides (Arioka et al., 2004; Fairn et al., 2009). In either case, mislocalization of YFP-Sktl in SigD-low cysts likely reflects a failure to retain sufficiently high levels of PIP2 at the plasma membrane in the presence of SigD. The mechanism by which Sktl concentrates at the growing ends of spermatid cysts remains an area for future study.

We show for the first time that PIP2 levels affect polarized exocyst distribution in a developmental context. In a survey of proteins that were candidates to be regulated by PIP2 during spermatid cyst polarization, we discovered that actin cytoskeletal proteins (anillin, P-moesin, and spectrin) associate with the growing ends of the flagellar axonemes in the middle of the bipolar SigD-low cysts. These cytoskeletal proteins do not seem to be involved in establishing or maintaining spermatid polarity in Dro sophila. For example, anillin is dispensable for spermatid cyst polarity (Goldbach et al., 2010), and PIP2 binding by β-spectrin is not required for viability or male fertility (Das et al., 2008). In contrast, the exocyst was completely delocalized upon PIP2 reduction, and exocyst mutants show defects in spermatid cyst polarity.

Based on previous studies from yeast and mammalian cells, regulation of the exocyst by PIP2 is likely to be direct.
Indeed, binding of the exocyst subunits Sec3 and Exo70 to PIP2 appears critical for exocyst function (He et al., 2007; Liu et al., 2007; Zhang et al., 2008). Consistent with this idea, we found that the exocyst subunit Sec8 localizes immediately adjacent to PIP2 at the growing end of spermatid cysts and that localization of Sec8 and Sec6 is strongly influenced by PIP2 levels: reduction of PIP2 caused complete delocalization of the exocyst, and rescue of PIP2 by Sktl expression restored colocalization. Furthermore, the exocyst is required for polarity of Drosophila spermatid cysts. A hypomorphic mutation in fun, which encodes Sec8, caused defects in cyst polarity, whereas loss of function of onr, which encodes Exo84, caused formation of apolar cysts. These defects in polarity, common to exocyst mutants and SigD transgenic flies, were not due to failure of cytokinesis, because cytokinesis was normal in SigD-low flies. Instead, our data suggest that establishment of spermatid cyst polarity relies on localized recruitment of the exocyst, which drives targeted membrane delivery to the growing end. Alternatively, or in addition, PIP2 and the exocyst may help establish the axis of polarization by linking the spermatid nuclear envelope to the plasma membrane, as suggested from analysis of sktl2.3 mutant clones during oogenesis (Gervais et al., 2008) or by regulating formation of the flagellar membrane that surrounds the growing end of the axoneme (Fritz-Niggli and Suda, 1972; Tokuyasu, 1975). Indeed, given the structural similarity between flagella and cilia and the requirement for the exocyst in ciliogenesis, PIP2 could play a conserved role in regulating this critical process.

Figure 6. PIP2 is required for exocyst localization at the growing end of spermatid cysts. (A–E) Confocal micrographs showing colocalization of PIP2 and Sec8 by immunofluorescence. In wild type, PIP2 (A) and Sec8 (B) are enriched at the growing end, with PIP2 surrounding Sec8 (C). In SigD-low cysts, Sec8 fails to concentrate at the growing end (D). Coexpression of Sktl with SigD-low partially restores association of Sec8 with PIP2, at the growing end (E). (F–H) Confocal micrographs showing the distribution of Sec8 (grayscale) and Sec6 (green) in elongating spermatid cysts from wild type (F), SigD-low (G), and SigD-low coexpressing Sktl (H). Arrows indicate the growing ends. In wild type, Sec8 and Sec6 localize in a honeycomb pattern at the growing end (F). Sec8 and Sec6 are delocalized in SigD-low cysts (G). Sec8 localization is partially restored by coexpression of Sktl with SigD-low (H). Scale bars, 10 μm.

Figure 7. The exocyst and PIP2 are required for spermatid elongation, polarity, and membrane addition. (A and B) Phase-contrast micrographs of whole testes, and (C and D) epifluorescence micrographs of spermatid cysts expressing β2-tubulin GFP (green) and stained for DNA (magenta). onr mutant cysts fail to elongate and are apolar (A and C). fun mutant cysts elongate and are bipolar (B and D). (E and F) Diagrams illustrating the distribution of nuclei in onr (E) and fun (F) mutant cysts. (G–J) Epifluorescence micrographs showing distribution of Sec8 (red, G), Sec6 (green, H), F-actin (red, I), α-spectrin (red, J), and anillin (green, J) relative to DNA (blue, magenta, or cyan) in fun mutant cysts. Arrows, the growing end (I and J). (K–M) Transmission electron micrographs showing ultrastructure of axonemes, mitochondrial derivatives and plasma membranes in early spermatids. Curved red arrows indicate rotational polarity of axonemes. White arrowheads, mitochondrial derivatives filling with electron-dense paracrystalline material. In wild-type cysts (K), axonemes show the same rotational polarity, and each axoneme is associated with a major (m1) and a minor (m2) mitochondrial derivative. In SigD-low (L) and fun (M) cysts, axonemes are antiparallel. Mitochondrial derivatives in SigD-low (L) and fun (M) contain multiple sites of filling with paracrystalline material. Scale bars (A and B), 50 μm; (C, D, and G–J), 20 μm; (K–M), 200 nm.
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