

Asterless is required for centriole length control and sperm development

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Centrioles are the foundation of two organelles, centrosomes and cilia. Centriole numbers and functions are tightly controlled, and mutations in centriole proteins are linked to a variety of diseases, including microcephaly. Loss of the centriole protein Asterless (Asl), the *Drosophila melanogaster* orthologue of Cep152, prevents centriole duplication, which has limited the study of its nonduplication functions. Here, we identify populations of cells with Asl-free centrioles in developing *Drosophila* tissues, allowing us to assess its duplication-independent function. We show a role for Asl in controlling centriole length in germline and somatic tissue, functioning via the centriole protein Cep97. We also find that Asl is not essential for pericentriolar material recruitment or centrosome function in organizing mitotic spindles. Lastly, we show that Asl is required for proper basal body function and spermatid axoneme formation. Insights into the role of Asl/Cep152 beyond centriole duplication could help shed light on how Cep152 mutations lead to the development of microcephaly.

Introduction

The centrosome is the major microtubule organizing center (MTOC) of many cells, serving as a site of microtubule (MT) nucleation and minus end organization. These nonmembrane bound organelles are critical for a variety of cellular processes including cell migration, immune cell function, neuronal pathfinding, and axon selection, among others (Bettencourt-Dias et al., 2011; Bornens, 2012; Angus and Griffiths, 2013; Sakakibara et al., 2013; Elric and Etienne-Manneville, 2014). In many cells, centrosomes serve as spindle poles and help construct and organize a bipolar mitotic spindle (Vitre and Cleveland, 2012; Helmke et al., 2013). Centrosome number and activity are tightly regulated to guarantee proper MTOC function and avoid detrimental effects at the cellular, tissue, and organismal level, which have been linked to human diseases, including microcephaly and cancer (Noatynska et al., 2012; Korzeniewski et al., 2013; Nigg et al., 2014).

Centrosome number is controlled by limiting its duplication to once per cell cycle. Building a centrosome involves assembling its two major components, centrioles and pericentriolar material (PCM). Centrioles are barrel-like structures composed of nine triplet MTs and many highly conserved proteins that are recruited and arranged in a stepwise assembly process that ensures proper centriole size and function (Pelletier et al., 2006). For example, Sas-6 and Ana2/STIL are recruited early

in the process to build the cartwheel structure that sets the diameter and radial symmetry of the centriole (Kitagawa et al., 2011; van Breugel et al., 2011; Dzhindzhev et al., 2014; Ohta et al., 2014). Cep97, CP110, and Sas-4/CPAP are then recruited to ensure proper centriole length (Spektor et al., 2007; Kohlmaier et al., 2009; Schmidt et al., 2009; Tang et al., 2009; Franz et al., 2013). When serving as MTOCs, the mother centriole recruits and organizes the PCM, from which MTs are nucleated. Some of the major PCM components include Pericentrin-like protein (PLP)/Pericentrin, Centrosomin (Cnn)/Cdk5Rap2/Cep215, and Spd2/Cep192, which then recruit gamma tubulin (γ -tub; Pelletier et al., 2004; Zimmerman et al., 2004; Fong et al., 2008; Giansanti et al., 2008; Zhu et al., 2008).

A major challenge to understanding the role of many centrosome proteins is that their loss leads to a loss of centrosome duplication and the subsequent dilution of centrosomes from the cell population (Goshima et al., 2007; Dobbelaere et al., 2008; Gönczy, 2012; Balestra et al., 2013). This precludes analysis of the potential roles of multifunctional centrosome proteins in PCM assembly, MTOC function, and ciliogenesis, among other processes. It is likely that proteins critical for centrosome duplication have unappreciated functions in other aspects of centrosome biology. One such protein is Asterless (Asl), the *Drosophila melanogaster* orthologue of vertebrate Cep152. In the absence of Asl or Cep152, centriole duplication halts because of its role in recruiting and stabilizing the master centriole duplication kinase Plk4 (Cizmecioglu et al., 2010;

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Abbreviations used in this paper: Ac-tub, acetylated tubulin; Asl, Asterless; BB, basal body; Cnn, Centrosomin; fGSC, female germline stem cell; γ -tub, γ -tubulin; IP, immunoprecipitation; mGSC, male germ line stem cell; MT, microtubule; MTOC, microtubule organizing center; PA, pharate adult; PCM, pericentriolar material; PLP, Pericentrin-like protein; SC, spermatocyte; SIM, structured illumination microscopy; ST, spermatid; Unc, Uncoordinated; wt, wild type; Y2H, yeast two hybrid.

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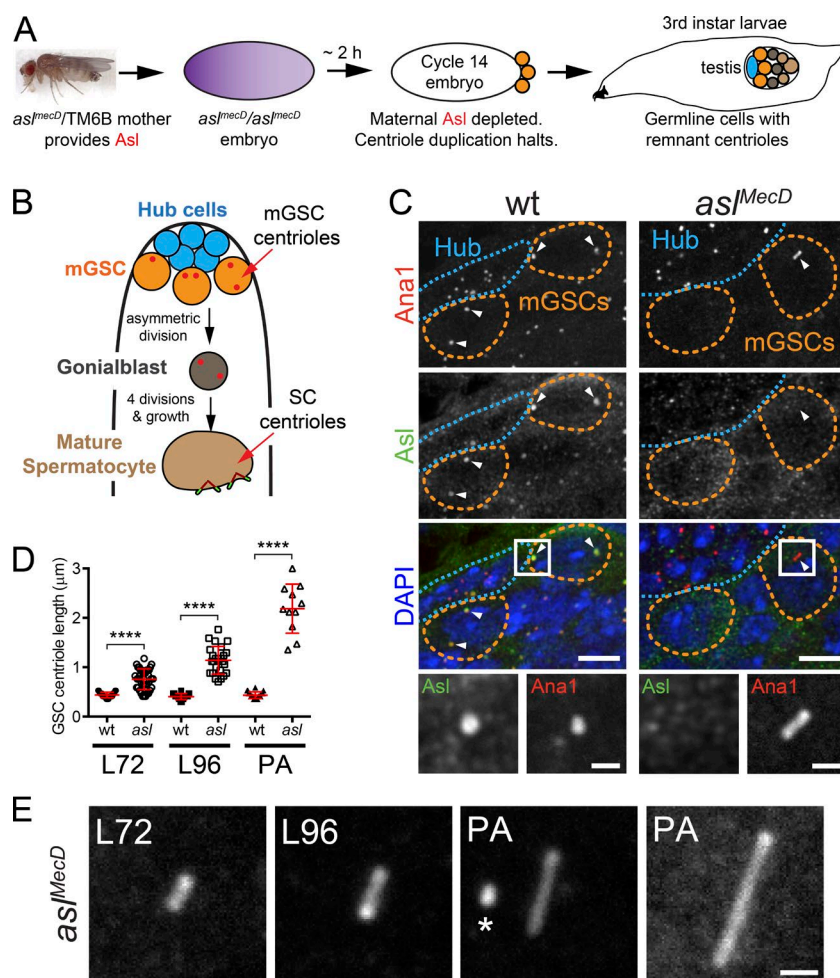


Figure 1. Asl is required for proper centriole length control. (A) *asl* embryos receive maternal Asl protein and mRNA (purple) used for the first 2 h of development. When this protein runs out, centriole duplication halts. Centrioles present at this point are reliably inherited by the developing testes (remnant centrioles). (B) Cell types in the *Drosophila* testis (centrioles, red). (C) Remnant centrioles (Ana1::tdTomato, red) in larval *asl* mGSCs do not have Asl (green) and are longer than in wt. Arrowheads, mGSC centrioles. Boxed region enlarged. DAPI, blue. (D) Centriole length (Ana1::tdTomato length) in mGSCs in wt (*asl^{mecD}/TM6B*) and *asl* mutants in larvae 72 h after egg laying (L72), larvae 96 h after egg laying (L96), or PAs. Mean \pm SD, red. Comparison: ordinary one-way analysis of variance followed by Tukey's multiple comparisons test. ****, $P < 0.0001$. (E) Representative centrioles in mGSCs at indicated stages. The longest centriole observed is on the right. Asterisk, centriole in adjacent hub cell. Bars: (C) 5 μ m; (insets) 1 μ m; (E) 1 μ m.

Dzhindzhev et al., 2010; Hatch et al., 2010; Kim et al., 2013; Sonnen et al., 2013; Klebba et al., 2015). Asl may also “license” centrioles for their first duplication in *Drosophila* embryos (Novak et al., 2014). Although it is clear that Asl is critical for centriole duplication, little is known about other roles it may play in centrosome biology.

Several lines of evidence suggest Asl has other critical roles. Asl localizes along the length of the outer centriole surface (Varmark et al., 2007; Blachon et al., 2008). This differs from Plk4, which is found as a ring or spot on the centriole's proximal end (Kim et al., 2013), suggesting Asl could do more than recruit Plk4, like structurally organizing the centriole. Also, Asl and Cep152 adopt a radially extended conformation with their C termini adjacent to the centriole wall and N termini extending into the PCM (Mennella et al., 2012; Sonnen et al., 2012). This suggests Asl/Cep152 could link the centriole wall with PCM, possibly directly scaffolding PCM proteins. There is evidence both supporting (Bonaccorsi et al., 1998; Varmark et al., 2007; Dzhindzhev et al., 2010; Conduit et al., 2014) and opposing (Blachon et al., 2008) the hypothesis that Asl functions in PCM organization. Finally, the removal of Asl from basal bodies during late spermatogenesis is important for zygotic development after fertilization (Khire et al., 2015).

Given this evidence and the critical nature of Asl/Cep152 for development and human health (Kalay et al., 2011; Poulton et al., 2014), we sought to identify functions for Asl beyond centriole duplication. We examined an Asl-free centriole population

in the testes of *asl* mutant flies, focusing primarily on the male germline stem cells (mGSCs). We identified two novel roles for Asl. Most striking, we show that Asl plays a role in controlling centriole length, primarily via Cep97. We find that Asl-free centrioles recruit PCM and can organize mitotic spindles, indicating that Asl is not essential for PCM function. Finally, we show that *asl* mutant centrioles have defects in assembling into sperm, indicating a novel role for Asl in basal body function.

Results

Asl-free centrioles in *asl* mutant mGSCs

To elucidate roles for Asl beyond centriole duplication, we examined a population of cells containing centrioles, but lacking Asl, found in *Drosophila* testes. We refer to these centrioles as “remnants,” because they were originally built during early embryogenesis using maternally provided Asl (Fig. 1 A; Blachon et al., 2008). Using this strategy, we generated *asl^{mecD}* zygotic mutants and performed a detailed examination of these Asl-free, remnant centrioles. We confirmed that *asl^{mecD}* testes showed complete loss of full-length Asl protein (Fig. S1, A and B; Blachon et al., 2008).

The testis contains several cell types including mGSCs, the hub (stem cell niche of somatic origin), somatic cyst stem cells, and the progeny of these stem cells (Fig. 1 B). In wild-type (wt) larval testes, all of these cell types contain Asl-positive

centrioles (Figs. 1 C and S1 C). In testes from *asl^{mecD}* homozygous mutant flies, remnant centrioles were located in four distinct cell populations (Fig. S1 D), two at the proximal tip (zones 1 and 2), one at the distal end (zone 4), and one in the central region (zone 3; Fig. S1, D vs. the even distribution in controls in C). The majority of remnant centrioles in the proximal tip are in the hub (Fig. 1 C, blue outline; and Fig. S1 D, zone 1). These cells have been quiescent since early development. The presence of these Asl-free centrioles reinforces that Asl is dispensable for centriole maintenance. The centrioles in zone 2 were in the mGSCs (Fig. 1 C, orange outline; and Fig. S1 D, zone 2), which are actively proceeding through the cell cycle. We found that 42% of mGSCs in *asl^{mecD}* third-instar larvae (larvae 96 h after egg laying) contained Asl-free remnant centrioles (Figs. 1 C and S1 E). This was expected, as *asl^{mecD}* mutants produce no full-length protein and a small amount of a ~60 kD, C-terminally truncated protein lacking the centriole localization region (Fig. S1, A and B; Blachon et al., 2008; Klebba et al., 2015). The mGSCs of *asl^{mecD}* are actively proceeding through the cell cycle and reproducibly contain Asl-free centrioles. They are therefore ideal for studying the nonduplication roles of Asl.

Asl is required for proper centriole length control

Examining *asl^{mecD}* mGSC remnant centrioles revealed a striking result: by the late larval stage, all centrioles were longer than wt (Fig. 1 C). These centrioles get longer over time, averaging >2 μ m long by the pharate adult (PA) stage. Some are almost 3 μ m long (Fig. 1, D and E), 30-fold longer than the 100 nm length observed in wt mGSCs (Gottardo et al., 2015). This length increase is not simply a consequence of the time since Asl protein was exhausted during embryogenesis, because centrioles do not grow long in the hub. The major difference between these cells is that unlike hub cells, mGSCs have undergone multiple cell cycles since early embryogenesis. A model consistent with these observations is that centrioles retained in the *asl* mutant mGSCs get longer with each cell cycle. Interestingly, we found that the frequency with which centrioles are found in mGSCs decreases as larval and pupal development proceed (Fig. S1 F). This indicates that centrioles are being lost from the mGSCs over time, presumably as a consequence of cell division. We also observed remnant centrioles lost from the mGSCs in another centriole duplication mutant, *sas-4^{S2214}* (Fig. S1 G), indicating that this is a general phenomenon, not something specific to loss of Asl.

To ensure that the long centriole phenotype was a consequence of Asl loss, we examined other *asl* allelic combinations. Both *asl^{mecD}/Dff(3R)ED5177* and *asl²/Dff(3R)ED5177*, an independently derived, severe loss-of-function allele (Blachon et al., 2008) that produces no full-length protein (Fig. S1 B), also have abnormally long centrioles in mGSCs (Fig. S2, A–B'). These long centrioles are not a result of centriole duplication failure, because remnant centrioles in mGSCs of another centriole duplication mutant, *sas-4^{S2214}*, were indistinguishable in length from wt (Fig. S2, C and D). These results strongly support that Asl controls centriole length. For all subsequent experiments, we used *asl^{mecD}* homozygotes, which we refer to as *asl*.

To understand the role of Asl in controlling mGSC centriole length, we examined the structure and composition of *asl* long centrioles using structured illumination microscopy (SIM). Centriole diameter and length in wt and *asl* hub cells were indistinguishable (Fig. 2 A), confirming that hub centrioles do

not grow longer in *asl*. The radial distribution of Ana1 and PLP was normal on *asl* mGSC centrioles, confirming they are indeed centrioles and not unusual protein aggregates (Fig. 2 B). This is again consistent with the increased centriole length of Asl-free centrioles requiring passage through multiple cell cycles, a common circumstance in frequently dividing mGSC, but not in the quiescent hub.

One formal possibility is that the centrioles in *asl* mGSCs are long as a result of a premature execution of the normal developmental program of spermatogenesis in which centrioles elongate during basal body (BB) formation. To investigate this, we compared the “giant” centrioles in wt spermatocytes (SCs) to the long centrioles in *asl* mGSCs. Two defining characteristics of SC centrioles are the recruitment of the Uncoordinated (Unc) protein and the formation of small cilia at their distal tips marked by acetylated tubulin (Ac-tub; Riparbelli et al., 2012). The long centrioles in *asl* mGSCs did not contain Unc or Ac-tub (Fig. 2, C and D, blue boxes), indicating that *asl* long centrioles are a consequence of loss of Asl protein function and not an ectopic execution of the SC centriole elongation program. The absence of Ac-tub also suggests that centriole elongation in *asl* mutants arises by a mechanism distinct from the one in *cp110* mutants, where centrioles form extensive Ac-tub projections at their distal ends (Franz et al., 2013). Thus, Asl normally functions to prevent precocious centriole elongation and its loss does not accelerate BB development.

Centrioles in *asl* mGSCs are proximalized

To understand the nature of the extension of centriole length and to identify possible causes, we examined localization of key centriole proteins. Sas-6, a component of the internal centriole cartwheel structure, and Sas-4, localize to diffraction-limited spots in wt mGSCs and to the proximal ends of wt SC giant centrioles (Fig. 3, A and B, left and middle). In *asl* mGSC long centrioles, Sas-6 and Sas-4 localize uniformly along the entire length (Fig. 3, A and B, right columns, blue boxes, and A' and B'). The elongation of Sas-6 and Sas-4 indicates that Asl controls both overall centriole length and the distribution of proximal centriole characteristics.

We next investigated the radial organization of Sas-6, Cep135, Sas-4, and PLP on *asl* long centrioles in mGSCs and wt SC giant centrioles (Fig. 3 C). Within the limits of this technique, the radial distribution of proteins on *asl* centrioles in mGSCs is normal. This provides additional evidence that these structures are centrioles, not protein aggregates or tubular extensions similar to the effects of Sas-6 and Ana2 overexpression (Gopalakrishnan et al., 2010; Stevens et al., 2010).

To determine if *asl* mGSC long centrioles retained distal characteristics, we examined Cep97 and CP110, proteins normally localized to the distal end of centrioles that function to control length (Spektor et al., 2007; Schmidt et al., 2009; Delgehyr et al., 2012; Franz et al., 2013). The antibodies we raised against these proteins were not suitable for immunofluorescence, so we generated animals expressing GFP-tagged versions of these proteins. As expected, Cep97 and CP110 localize as one or two dots in wt mGSCs and SCs (Fig. 3, D and E, left and middle columns). In *asl* mutant mGSCs, both Cep97 and CP110 are recruited to one end of long centrioles at almost normal levels (Fig. 3, D' and E'), with the caveat that both are overexpressed. Thus, *asl* centrioles, although long and mostly of proximal character, form a distal end.

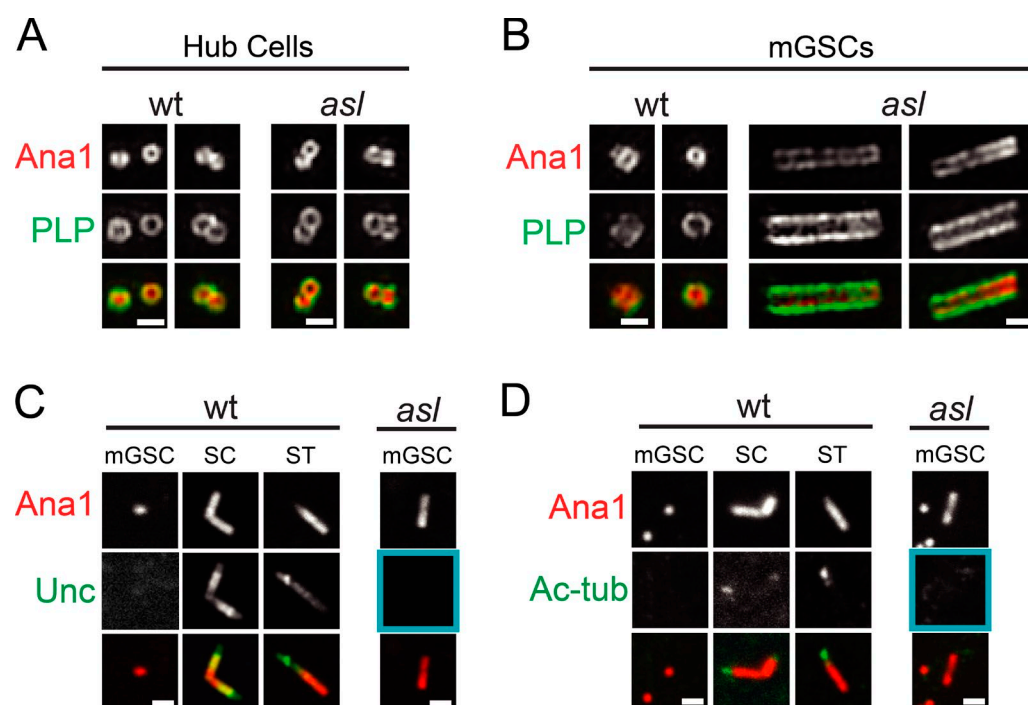


Figure 2. Loss of Asl does not trigger premature centriole to basal body transition in mGSCs. (A and B) SIM of hub (A) and mGSC (B) centrioles in wt (*asl^{mecD}/TM6B*; left) and *asl* (right) larval testes. Ana1::tdTomato, red; PLP, green. Two examples of each. (C and D) Confocal images of wt and *asl* centrioles. (C) Unc (green) was not on *asl* (blue box) or wt mGSC centrioles (Ana1, red) but was on wt SC and ST centrioles. (D) Ac-tub was on centrioles in 75% of SC and 100% of ST. Ac-tub was not on centrioles in wt or *asl* (blue box) mGSCs. Bars, 0.5 μ m.

Asterless regulates centriole length in other proliferative cells

To determine if Asl controls centriole length in other cell types, we examined additional tissues in *asl* mutants. Like the testis, the ovary is complex with many cell types (Fig. 4 A). Similar to mGSCs, remnant centrioles in female germline stem cells (fGSCs) were frequently too long (Fig. 4, B and C; 47/65 centrioles greater than wt mean plus two SDs). We also found that PLP (Fig. 4 D) and Sas-6 (Fig. 4 E) were localized along the entire length of fGSCs centrioles, indicating that centrioles are “proximalized.” These remnant centrioles had a distal end as indicated by the presence of Cep97 and CP110 (Fig. 4, F and G).

To determine if Asl controls centriole length in somatic tissues, we examined remnant, centrioles in *asl* larval wing discs. Long, proximalized centrioles were also identified in these cells (Fig. 4, H–J; 62/118 centrioles greater than wt mean plus two SDs), indicating that centriole length control by Asl is a conserved mechanism among different cell types. There is significant proliferation in the wing disc during larval development (Milán et al., 1996), supporting a model where Asl is required for regulating the length of centrioles that transit through multiple cell cycles. We attempted to examine the affect of loss of Asl on centriole length outside the animal by knocking down Asl in S2 cells. We located a few remnant centrioles in these cells but did not see centrioles that were longer than normal by light microscopy (unpublished data). However, we note that previous studies of core centriole length did not see an effect from knockdown of *cp110* in S2 cells, even by EM (Franz et al., 2013), although they did see an effect on centriole length in the animal in wing discs and we show an effect in mGSCs (see Fig. 6).

Asterless is not required for PCM recruitment and MTOC activity

Asl adopts an extended conformation with its C terminus adjacent to the centriole and its N terminus extended outward (Mennella et al., 2012; Sonnen et al., 2012), ideally positioned to organize PCM around the centriole. However, there are conflicting studies regarding the requirement for Asl in PCM recruitment (see Introduction). To directly test this, we examined several PCM proteins, beginning with PLP, which also extends from the centriole wall. Asl-free centrioles recruit PLP in both mitosis and interphase (Figs. 5 A and S2 E, right columns, blue box), consistent with RNAi studies in S2 cells (Mennella et al., 2012). Importantly, unlike SC centrioles, where PLP is exclusively proximal, PLP localizes along the length of *asl* long centrioles (Figs. 5 A and S2 E, right column, blue box). Because PLP, in *Drosophila*, often behaves more similarly to a centriole protein, we examined the distributions of the more canonical PCM proteins Cnn, Spd2, and γ -tub. Loss of Asl did not prevent the localization of any of these proteins to mGSC mitotic or interphase centrosomes (Fig. 5, B–D; and Fig. S2, F–H, right column, blue boxes), inconsistent with a role for Asl in recruiting PCM (Varmark et al., 2007; Dzhindzhev et al., 2010; Conduit et al., 2014). Furthermore, Cnn, Spd2, and γ -tub were localized along the length of the *asl* long centriole, in contrast to their strict proximal localization on giant SC centrioles (Fig. 5, B–D). These results expose an intimate, possibly physical, link between the core centriole structure and the position of PCM assembly (see Discussion).

The presence of all of the PCM components on *asl* centrioles suggests that these centrosomes could function as MTOCs. We found that *asl* centrioles in mGSCs organize a modest MT array in interphase and a robust array during mitosis along the

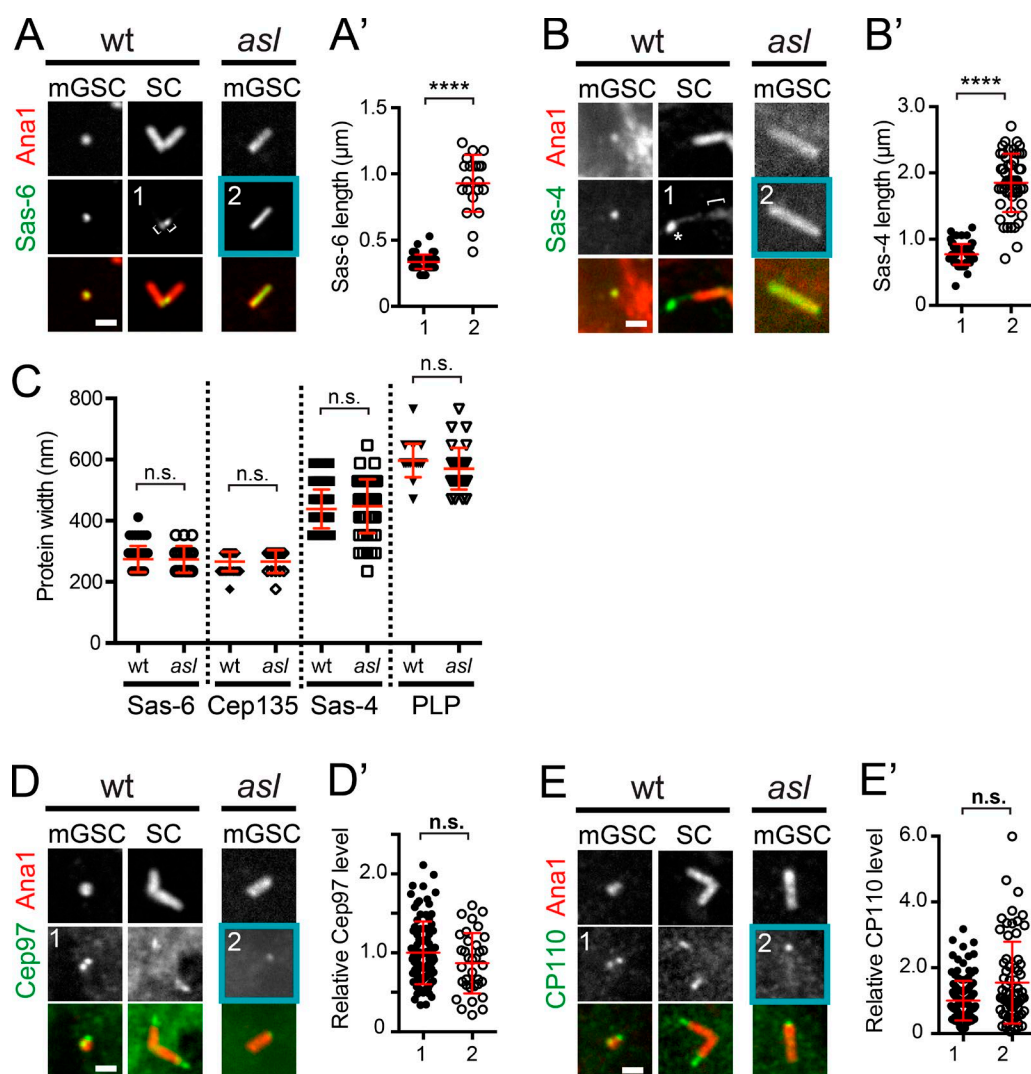


Figure 3. Asl regulates the position of centriole proteins. (A–E) Centrioles from interphase wt (*asl^{mecD/TM6B}*) and *asl* mGSCs or wt mature SCs. (A) GFP::Sas-6 (green) and Ana1::tdTomato (red). Sas-6 extends the length of the centriole in *asl* mGSCs but is only proximal in wt SC centrioles (brackets). (A') Length of Sas-6 signal in wt SCs (1) and *asl* mGSCs (2). (B) Sas-4::GFP (green) and Ana1::tdTomato (red). In SCs, Sas-4::GFP localizes proximally (bracket), where endogenous protein localizes (Gopalakrishnan et al., 2011; Fu and Glover, 2012). Additional localization beyond Ana1 (asterisk) is a consequence of overexpression. In *asl* mGSCs, Sas-4 is uniformly along the entire length of Ana1. (B') Lengths of the Sas-4 signal in wt SCs (1, only the proximal signal) and *asl* mGSCs (2). (C) The radial distribution (diameter) of centriole proteins is unaltered in *asl* mGSCs. (D) GFP::Cep97 (green) and Ana1::tdTomato (red). (D') Intensity of GFP::Cep97, normalized to the mean intensity of Cep97, at centrioles in wt (1) and *asl* (2) mGSCs. Cep97 levels are unaffected in *asl*. (E) CP110::GFP (green) and Ana1::tdTomato (red). (E') Intensity of CP110::GFP, normalized as in D, in wt (1) and *asl* (2) mGSCs. CP110 levels are unaffected in *asl*. Bars, 1 μ m. Mean \pm SD, red. Comparison by unpaired *t* tests, with Welch's corrections when appropriate. ****, $P < 0.0001$; n.s., not significant.

entire centriole (Figs. 5 E and S2 I, right column, blue box), mirroring the extended PCM localization. Live imaging of *asl* mGSCs during mitosis revealed that long centrioles can effectively organize a mitotic spindle pole (Fig. 5 F and Videos 1 and 2). However, because *asl* mGSCs only contain one centriole, they first build a half spindle focused at the long centriole, then an acentrosomal half-spindle on the other side of the chromosomes. These spindles properly proceed through mitosis with only slight delay (Fig. 5 F, AO), comparable mGSCs with no centrosomes (Fig. S2 J and Video 3). Most likely, this delay is a consequence of constructing the acentrosomal half-spindle, not the presence of an unusually long centriole.

To determine if Asl has a role in PCM recruitment in other cells types, we examined remnant centrioles in dividing SCs (Fig. S1 D, zone 4) of *asl* testis. We found that Asl-free centrioles

in SCs recruit γ -tub during meiosis I (Fig. 5 G), consistent with previous findings (Blachon et al., 2008). Note that the example presented does not contain γ -tub along the entire centriole (Fig. 5 G), as it was selected from zone 4 (Fig. S1). These centrioles were constructed before Asl loss and thus do not have an extended luminal (cartwheel) structure (see section Asl is required for basal body function for further details). Together, our results indicate that Asl is not essential to recruit the major PCM components and its loss does not affect the ability of these centrosomes to participate in mitotic or meiotic spindle function.

Asl's binding partner Cep97 is required for mGSC centriole length control

As mentioned, Cep97 and CP110 have been implicated in controlling centriole length, but both are present on *asl* long

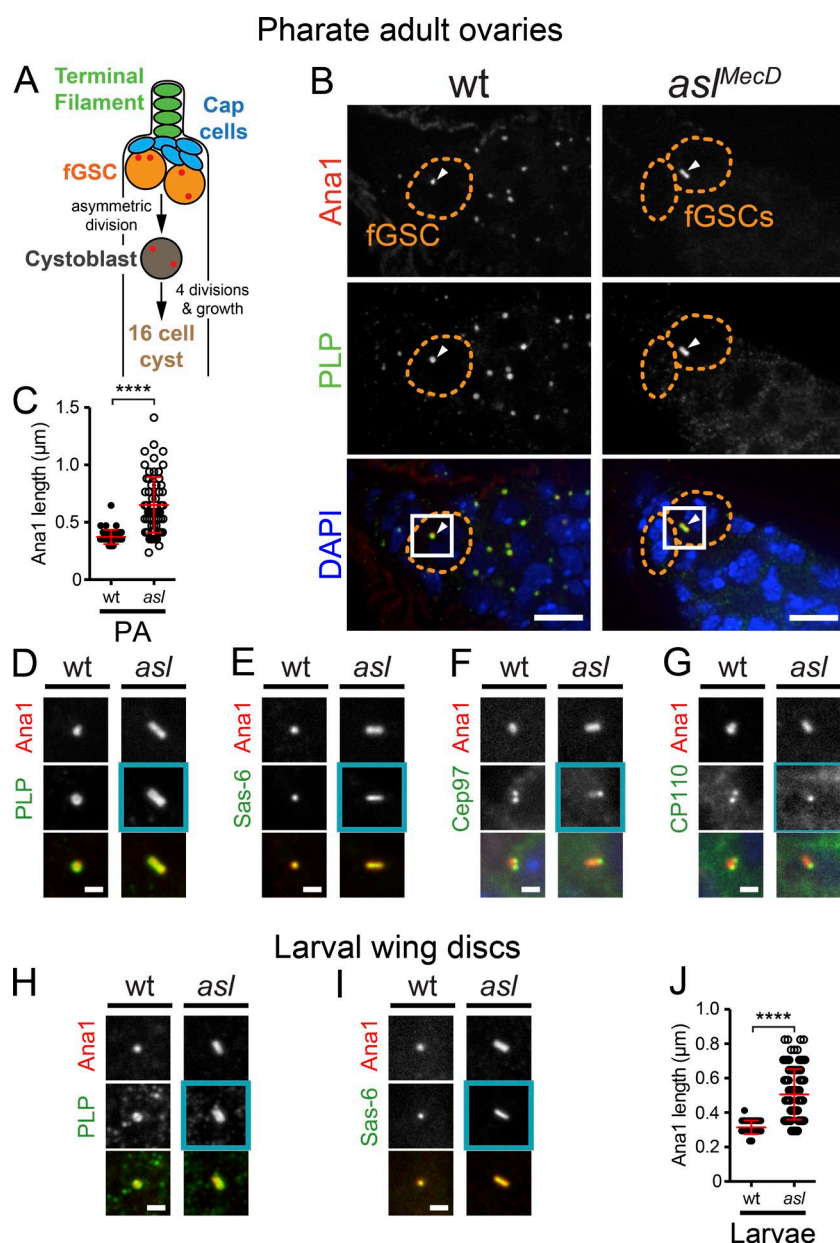


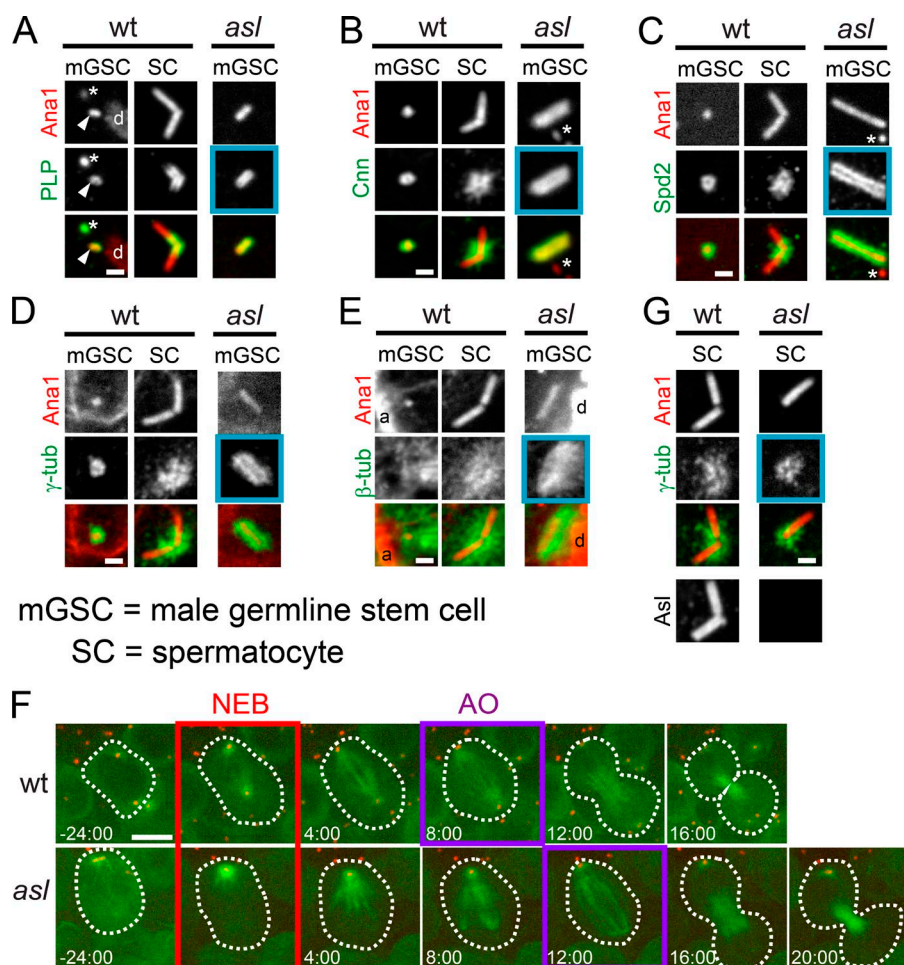
Figure 4. Asl controls centriole length in female germline and somatic cells. (A–G) Remnant centrioles in wt (*asl^{MecD}/TM6B*) and *asl* fGSCs from PAs. (A) Cell types in *Drosophila* ovaries (centrioles, red). (B) Remnant centrioles (Ana1::tdTomato, red; PLP, green) in *asl* fGSCs are longer than in wt. Arrowheads, centrioles in fGSCs. DAPI, blue. Boxes are enlarged in D. (C) Measurements of centriole length (Ana1::tdTomato), in fGSCs. (D) Boxed region from B. PLP is along the length of *asl* centrioles. (E) GFP::Sas-6 (green) extends the length of *asl* centrioles. (F and G) GFP::Cep97 (F, green) and CP110::GFP (G, green) localize to the proximal tip of *asl* fGSC centrioles. (H–J) Centrioles in wt (*asl^{MecD}/TM6B*) and *asl* larval wing discs. (H) Remnant *asl* centrioles (Ana1::tdTomato, red) are longer than wt. PLP (red) is along their entire length. (I) GFP::Sas-6 (green) extends the entire length of *asl* centrioles. (J) Centriole length (Ana1::tdTomato) in wing discs. Mean \pm SD, red. Comparison: unpaired *t* tests, with Welch's correction. ****, $P < 0.0001$. Bars: (B) 5 μ m; (D–I) 1 μ m.

centrioles (Fig. 3, D and E, right column, blue box). However, previous studies have shown that Cep97 and CP110 remain at the distal ends of centrioles in other cases where centrioles grow too long (Kohlmaier et al., 2009; Schmidt et al., 2009; Tang et al., 2009; Korzeniewski et al., 2010; Lin et al., 2013). Thus the localization of these proteins to centrioles alone is not sufficient to ensure proper length control. Therefore, we hypothesized that Asl is required for proper function, not recruitment, of Cep97 and/or CP110 at the centriole. This hypothesis predicts that loss of either of these proteins would result in a similar long centriole phenotype to loss of Asl.

Long centrioles in mGSCs of *cp110* null (*cp110Δ*) flies have not been reported (Franz et al., 2013). We reexamined *cp110Δ* flies and found that ~2.4% of mGSCs/testes contain a long centriole similar to *asl* mutants and never observed in >1,000 control wt mGSCs (Fig. 6, A and E). To examine Cep97 loss, we obtained a transposable element insertion in the first Cep97 intron, which we refer to as *cep97^{LL}* (Schuldiner et al., 2008). We confirmed that the level of Cep97-PB, the longest

isoform, was reduced by >85% in *cep97^{LL}* (Fig. S3 A; LL/LL). Examining homozygous *cep97^{LL}* larval testes showed ~6% of mGSCs/testis contain long centrioles, whereas controls had none (Fig. 6 E and Fig. S3, B–D, LL/Cyo vs. LL/LL). To confirm that this was a consequence of Cep97 loss, we examined *cep97^{LL}* as hemizygotes over a small chromosomal deletion (Materials and methods; *Df(2L)A1*). *cep97^{LL}/Df(2L)A1* flies have long centrioles in ~14% of mGSCs/testis (Fig. 6, B, C, and E, LL/Df), a phenotype abrogated by the introduction of a GFP::Cep97 transgene (Fig. 6, D and E; and Fig. S3 E).

All of the long centrioles in *cep97^{LL}* had PLP and Sas6 decorating their entire length, indicating that they have an extended proximal organization similar to centrioles in *asl* mutants (Fig. 6, B, C, and F). Interestingly, we find that nascent procentrioles (indicated by Sas6) in the *cep97^{LL}* were not always located at one end of the centriole (Fig. 6 F, bottom row). This supports a model where the entire long centriole is of proximal character, as procentriole formation is normally reserved for the proximal end. These data show that Cep97, and



to a lesser extent CP110, work to regulate centriole length in mGSCs in a manner similar to Asl. Because the long centriole phenotype was significantly stronger in *cep97*, we focused subsequent studies on Cep97.

One inconsistency between the *asl* and *cep97* mutant phenotypes is the frequency of long centrioles in mGSCs, with 100% of remnant centrioles being long in *asl* mutant and only 14% being long in the *cep97*. One main difference is that centriole duplication is blocked in *asl*, but not in *cep97* mutants. To test if this accounted for the frequency difference, we blocked centriole duplication in the *cep97* background by introducing a *plk4* mutation. By blocking centriole duplication, the only centrioles that will remain will be those present in mGSC since early embryogenesis. Although remnant centrioles in the mGSCs of *plk4* mutants were normal length (Fig. 6, G and H), in *cep97 plk4* double mutants, 82% of remnant centrioles were long (longer than *wt* mean plus two SDs; Fig. 6, G and H). This is very similar to the 100% seen in *asl* mutants, and is consistent with Cep97 and Asl functioning in the same pathway to control centriole length. In both *plk4* and *cep97;plk4* mutants, centrioles are not found in every mGSC (Fig. 6 G). Combined with our findings that centrioles are not retained in all *asl* and *sas-4* mutant mGSCs (Fig. S1, F and G), it suggests that long centrioles were seen at a low frequency in *cep97* mutants because centrioles are normally lost from mGSCs at a certain frequency and the longest centrioles were the few retained by the mGSC over multiple cell cycles. This is contrary to the prominent model where mGSCs retain their mother centrioles indefi-

nately (Yamashita et al., 2007) but in agreement with analysis of remnant *sas-4* mutant centrioles occasionally found in mGSCs (Riparbelli and Callaini, 2011).

To better link Asl and Cep97, we tested if Asl and Cep97 physically interact. We divided both proteins into three sub-fragments, taking care to not disrupt any predicted motifs or secondary structures (Fig. 7 A). We then tested for direct interaction between these fragments by yeast-two hybrid (Y2H) screening, finding that the C-terminal region of Asl interacts with both the N- and C-terminal regions of Cep97 (Fig. 7, A and B; and Fig. S3 F). We confirmed these interactions by coimmunoprecipitation (coIP) using fragments (Fig. 7, C and D) and full-length proteins (Fig. 7, E and F) overexpressed in S2 cells.

Cep97 functions downstream of Asl to regulate centriole length

A simple model consistent with our findings is that Asl functions through Cep97 to regulate centriole length. In this case, we would predict that the long centriole phenotype in *asl* mGSCs might be overcome if large amounts of Cep97 was forced to the centriole. To accomplish this, we generated a transgenic fly ubiquitously expressing Cep97 fused to GFP and the PACT domain of PLP, a commonly used centrosome-targeting domain (Gillingham and Munro, 2000). We refer to this construct as Cep97^{PACT} (Fig. 8 A). Cep97^{PACT} localized to *asl* mutant mGSC centrioles, and these centrioles were significantly shorter than those in *asl* mGSCs not expressing Cep97^{PACT} (Fig. 8, B–D; and Fig. S4, A and B). Expression of this construct in *wt* did

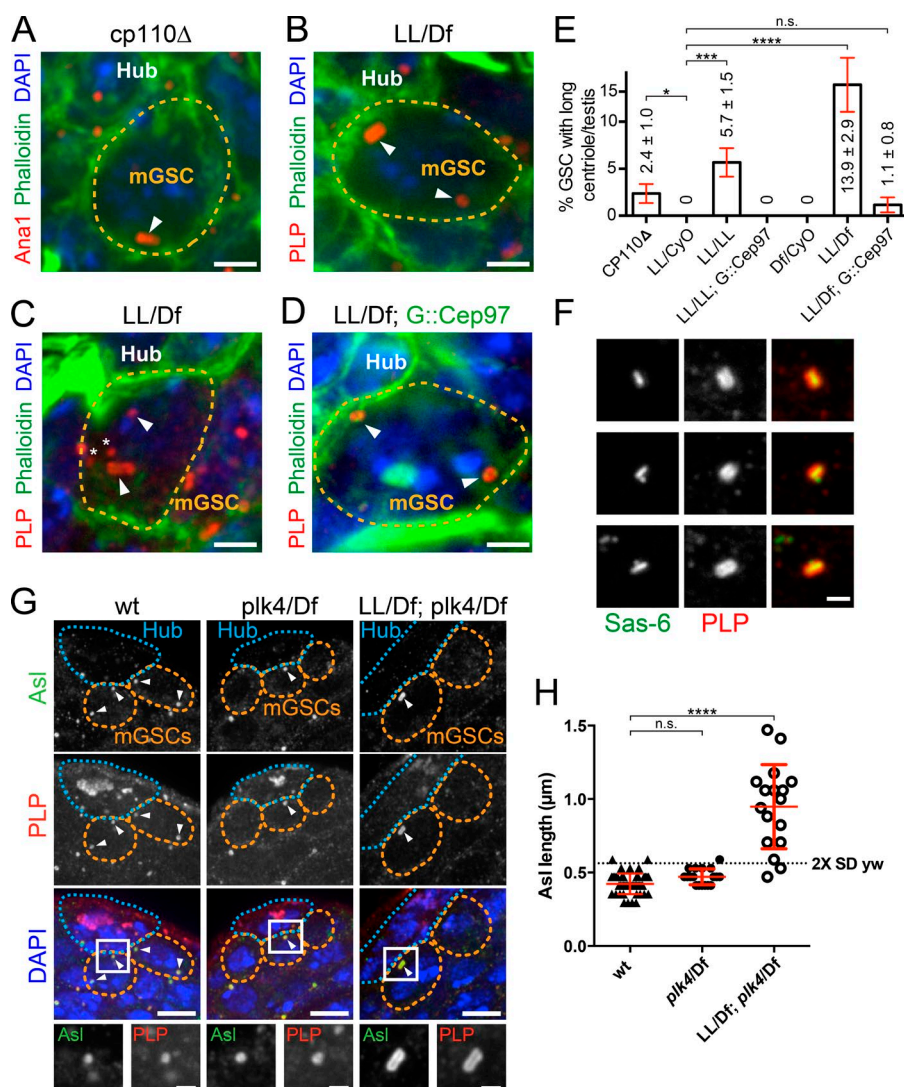


Figure 6. *Cep97* controls mGSC centriole length. (A–D) mGSCs in adult testes of indicated genotypes. Ana1::tdTomato, red (in A); PLP, red (in B–D); phalloidin, green; DAPI, blue. Arrowheads represent centrioles in mGSCs. Asterisks represent centrioles from the hub. (A) An mGSC from a *cp110Δ* testis with a long centriole. The other centriole is not visible in this projection. (B) *cep97^Δ/Df(2L)A1* (LL/Df) mGSC with a long apical centriole. (C) LL/Df mGSC with a long nonapical centriole. (D) An LL/Df mGSC expressing GFP::Cep97 has normal-length centrioles. (E) Percentage of mGSCs with long centrioles per testis ± SEM for indicated genotypes. Testes counted include *cp110Δ*, 36; *LL/CyO*, 17; *LL/LL*, 34; *LL/CyO*; GFP::Cep97, 23; *LL/Df*, 32; and *LL/Df*; GFP::Cep97, 27. Comparison: unpaired *t* tests, with Welch's correction. ****, *P* < 0.0001; ***, *P* < 0.001; *, *P* ≤ 0.05; n.s., not significant. (F) GFP::Sas-6 (green) and PLP (red) extend along the length of *cep97* centrioles. Procentriole can position at one end (middle row) or the middle (bottom row) of the elongated centriole. (G) Remnant centrioles (Asl, green; PLP, red) in *cep97;plk4* double-mutant adult mGSCs are longer than in wt (yw) or *plk4*. mGSCs, orange line; DAPI, blue; arrowheads, centrioles in mGSCs. Boxed region enlarged (bottom). (H) Centriole (Ana1::tdTomato) length (red line, mean ± SD) in mGSCs in adults of the indicated genotypes. Dashed line is the mean of wt plus two SDs. Almost all centrioles in mGSCs are long in *cep97;plk4*. Comparison: ordinary one-way analysis of variance followed by Tukey's multiple comparisons test. ****, *P* < 0.0001; n.s., not significant. Bars: (A–D) 2 μm; (F and G, bottom row) 1 μm; (G) 5 μm.

not prevent the elongation of the normally giant centrioles in SCs (Fig. S4 C). Importantly, neither expression of *Cep97* (Fig. S4 D) nor expression of PACT (Fig. S4 E) could suppress the long centriole phenotype of *asl* mutants, even when expressed at higher levels than *Cep97^{PACT}* (Fig. S4, F and G). This strongly support a model where *Cep97* is a downstream effector of Asl in centriole length control.

We attempted to further link Asl and *Cep97* function by generating a mutation in Asl specifically disrupting the Asl–*Cep97* interaction. However, generating such a mutant proved to be extremely challenging. We produced >1,400 Asl mutant alleles using a reverse Y2H screen (see Materials and methods) and screened them for interaction with *Cep97* and other proteins that interact with Asl (Fig. S5 A). We were unable to generate a mutant that specifically disrupted the interaction with both regions of *Cep97* while maintaining the interaction with all other binding partners. This is consistent with the critical nature of the C terminus of Asl for centriole localization and Plk4-dependent centriole duplication (Klebba et al., 2015), and the large number of protein interactions made by this region (unpublished data). However, the direct interaction between Asl and *Cep97*, the similar mutant phenotypes, and the suppression of the *asl* phenotype by *Cep97^{PACT}* all support a model where Asl and *Cep97* function together to regulate centriole length.

Asl is required for basal body function

The progeny cells resulting from the asymmetric division of mGSCs undergo a differentiation program to become sperm, during which the centrioles undergo major changes. In SCs, centrioles elongate to form giant centrioles by extending the distal end and maintaining the cartwheel protein Sas-6 and PCM at the proximal end. Later, in spermatids (STs), centrioles act as BB that dock at the nuclear membrane and extend the flagellar axoneme from their distal end (Fig. 9 A; Fabian and Brill, 2012). As we have demonstrated in Fig. 3, unlike SC giant centrioles, the long centrioles in *asl* mGSCs adopt a proximal centriole character. Given this vastly different proximal/distal protein arrangement in *asl* mutant centrioles, we hypothesized that they would not be able to function as BBs. Because the SCs and STs are the progeny of the mGSCs, we reasoned that we could test this hypothesis if we could identify centrioles that grew long in the mGSC and were subsequently passed on to its progeny.

In *asl* mutant testes, we find remnant centrioles in all of the cell types derived from mGSCs (Fig. S1 D, zones 3 and 4). These cells could have received their centrioles via two mechanisms. (1) They could inherit centrioles from mGSCs before maternal Asl depletion. We predict these centrioles would have a normal proximal/distal character with Sas-6 restricted to a dot

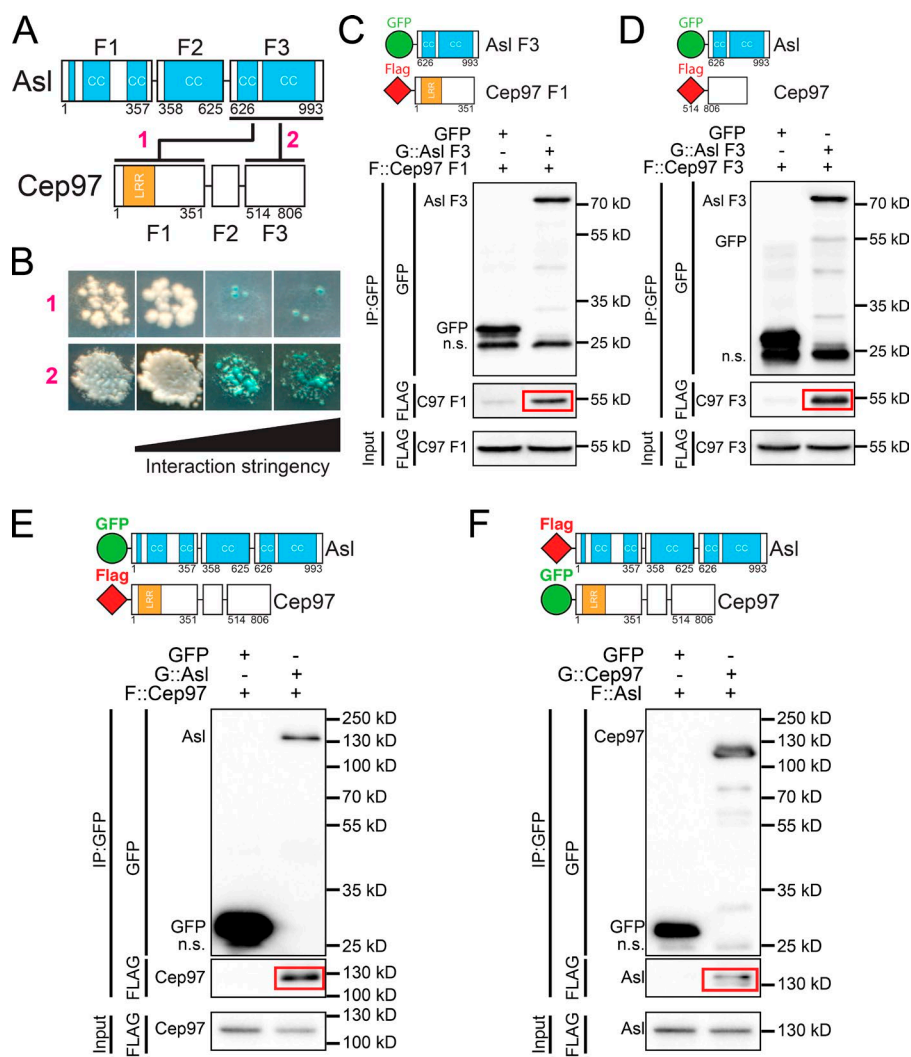


Figure 7. Asterless and Cep97 interact. (A) Schematics of the subfragments (F1–F3) of Asl and Cep97 used for interaction assays. Red numbers refer to B. Numbers are amino acids. Lines, Y2H interactions; CC, coiled coil; LRR, leucine-rich repeat. (B) Asl F3 interacts with Cep97 F1 (1) and Cep97 F3 (2) by Y2H screening. The first column shows the presence of Y2H plasmids. The remaining columns show interaction under conditions of increasing stringency (see Materials and methods). Red numbers refer to A. (C) GFP::Asl F3 colPs Flag::Cep97-F1 from S2 cells. (D) GFP::Asl F3 colPs Flag::Cep97-F3 from S2 cells. (E) GFP::Asl full-length colPs Flag::Cep97 full length from S2 cells. (F) GFP::Cep97 full-length colPs Flag::Asl full length from S2 cells.

at the proximal end. We term these “Sas-6 normal” centrioles. (2) They could inherit a centriole that was in the mGSC when maternal Asl was depleted and grew long in the mGSC before being passed on. These centrioles would have a proximal/distal character similar to the centrioles in *asl* mutant mGSCs and have an extended Sas-6 localization. We term these centrioles “Sas-6 extended.”

We examined testes from pupae and PAs, and found remnant, Asl-free centrioles, in STs. The vast majority of centrioles were Sas-6 normal; however, we identified some Sas-6 elongated centrioles ($n = 16$ centrioles), identical to the centrioles in *asl* mGSCs. We found that 15 out of 16 (from 11 testis of ~30 examined) of these centrioles were defective in nuclear attachment and axoneme formation (Figs. 9 B and S5 B). Interestingly, 14 of the 15 defective BBs formed an extension of Ana1 distal to the elongated Sas-6 signal. This suggests that they underwent the normal elongation of the distal end that takes place in SCs and that defects in BB function might not be a consequence of extended proximal character, but rather a consequence of an independent and as of yet undescribed function of Asl. If this is the case, we would predict that the Sas-6 normal centrioles would show similar BB defects.

When we examined Sas-6 normal remnant centrioles in spermatids, only 37 out of 101 (20 in early and 17 in late spermatids) BBs were attached to a nucleus and had built a normal-

looking axoneme. The remaining 64 had defects in nuclear attachment, improper axoneme formation, or both (Figs. 9 C and S5 C). Most striking, we observed BBs in early spermatids that had forked axonemes as indicated by Ac-tub (Figs. 9 C and S5 C, arrows). Given these defects were seen in *asl* mutant BBs with normal proximal/distal character (Sas-6 normal), they are most likely a consequence of losing yet another unknown function of Asl specific to the BB. Consistent with this idea, Asl protein localization is dramatically rearranged during the final stages of BB formation from coating the centriole wall to a ring-like structure at the base of the axoneme, approximately where the BB to axoneme transition occurs (Fig. S5 D). Thus, we have identified a novel role for Asl in proper BB function during the final stages of spermatogenesis, and this role is independent of its role in centriole length control.

Discussion

Loss-of-function studies have identified proteins critical for the function of centrosomes (Goshima et al., 2007; Dobbelaere et al., 2008; Gönczy, 2012; Balestra et al., 2013). The loss of many of these proteins results in the loss of centrosomes from the cell, precluding analysis of other roles they might play beyond centrosome duplication or maintenance. Here, we investigate

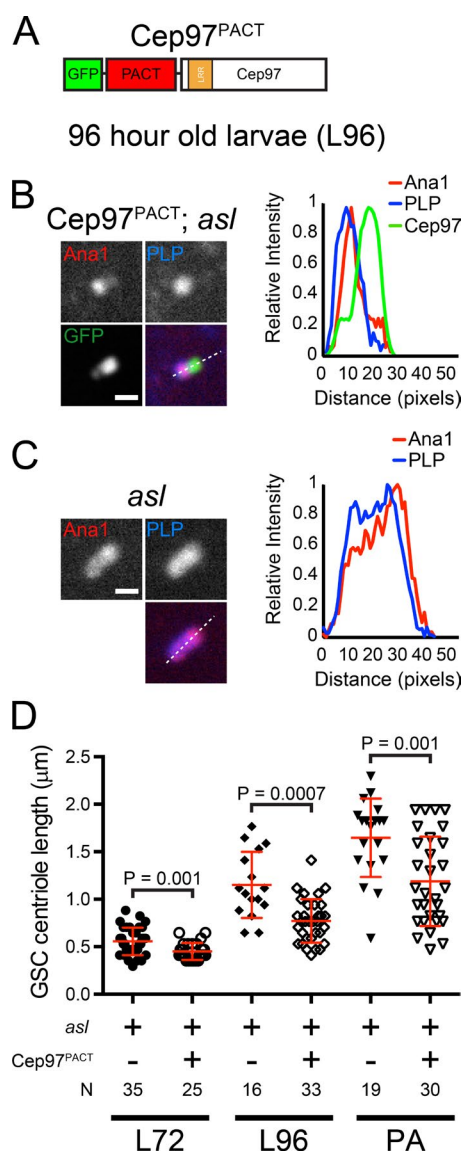


Figure 8. Cep97 functions downstream of Asl in mGSC centriole length control. (A) Centriole targeted Cep97 construct. LRR, leucine-rich repeat; PACT, centriole targeting domain of PLP. (B) A representative centriole in *asl* mGSCs expressing Cep97^{PACT}. PLP, blue; Cep97^{PACT}, green; Ana1::tdTomato, red. Linescans (dashed line) along the long axis of the centriole, normalized to peak intensity. (C) A representative centriole in *asl* mutant mGSCs. PLP, blue; Ana1::tdTomato, red. Linescans as in B. (D) Length of Ana1::tdTomato in *asl* mutant mGSC centrioles at indicated developmental time from individuals with or without Cep97^{PACT} expression. Additional examples are shown in Fig. S4. Error bars are \pm SD. Comparison: unpaired *t* tests with Welch's corrections when appropriate. N, centrioles measured. Bars, 1 μ m.

roles for Asl beyond its well-established function in centriole duplication. We take advantage of *Drosophila* zygotic mutant animals that build centrioles in early development using maternally provided Asl. We then identify these centrioles in larval and adult tissues where they now reside in an environment genetically null for Asl. These remnant centrioles have proven to be a critical tool to investigate Asl, but similar analysis can be performed on remnant centrioles generated using zygotic mutants of other centriole duplication factors.

The most striking finding was that the absence of Asl resulted in the massive (up to 30-fold) elongation of centrioles

in both germline and somatic cells. Although long, these centrioles maintain normal radial distributions of all the tested proteins and a characteristic distal centriole tip (Fig. 10 A). These long *asl* mutant centrioles are formed in a manner that results in proximal centriole proteins along the entire centriole (Fig. 10 A). This is different than the elongation seen in SC centrioles, where these proteins are restricted to the proximal end. Restricting the growth of the centriole proximal domain is a mechanism that has not been previously documented.

Unlike in the other cells examined, remnant centrioles in *asl* hub cells do not grow long. This supports the model that centriole elongation requires cells to be actively proceeding through the cell cycle. Our data suggest that Asl's role in controlling centriole length is most important in highly proliferative cells where a given centriole has the highest chance of being maintained in the same cell over multiple cell cycles. This is a circumstance common to stem cells. Thus, our identification of significant length control defects in two distinct stem cells suggests that length control by this mechanism might be especially critical in stem cells.

Direct protein binding and genetic analysis revealed that Cep97 functions downstream of Asl to exert a significant portion of its centriole length control (Fig. 10 C). The presence of overexpressed Cep97 at the distal end of Asl-free centrioles suggests that the mechanism by which Asl controls length is not simply to recruit Cep97. It remains a formal possibility that at endogenous levels Cep97 localization might be affected by loss of Asl. An attractive alternative model is that Cep97 localizes to centrioles via binding to a different protein but that binding to Asl enhances Cep97's centriole length control activity. The ability of centriole targeted Cep97 to partially compensate for the loss of Asl is consistent with this model. Our data also suggest that Asl does not function exclusively through Cep97 because the long centrioles in *cep97* mutants are shorter than those in similarly aged *asl* mutants.

Cep97 and CP110 have been implicated together in the control of centriole length. However, we found that a hypomorphic *cep97* allele had a stronger effect on centriole length than a null *cp110* mutant, suggesting a role for Cep97, independent of CP110. Additionally, the centrioles in both *asl* and *cep97* mGSCs are morphologically different than the elongated centrioles reported in *Drosophila* cells lacking CP110. These centrioles were only ~ 1.1 times (10%) longer than controls and had a significant MT protrusion (marked by Ac-tub) at one end (Franz et al., 2013); similar results have been reported in mammalian cells (Schmidt et al., 2009). Interestingly, loss of Klp10A in *Drosophila* cells results in both significantly longer centrioles (approximately two times longer [200%]) with additional MT extensions from its distal end, and its function does not require CP110 (Delgehyr et al., 2012; Franz et al., 2013). This supports the conclusion that centriole protein proximal-distal distribution, the length of the centriole as a whole, and the length of centriole wall MTs might all be regulated by distinct, possibly cell type-specific, mechanisms.

One interesting question is why cells have evolved a mechanism to control centriole length. It is possible that evolution of the centriole has selected for a minimum centriole "functional unit" that can efficiently accomplish its task, in this case, serving as an MTOC. Any unessential centriole length might require additional cellular resources, which could provide selective pressure to evolve a centriole length control mechanism. Another possibility is the presence of limited amounts of centriole

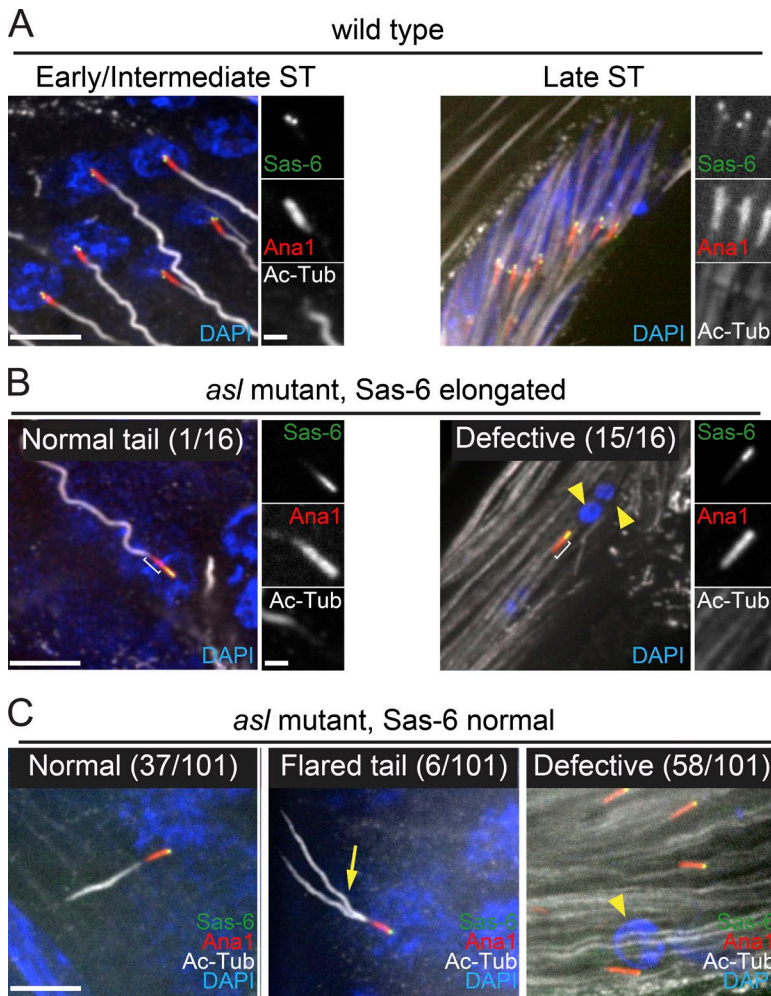


Figure 9. Asl is required for proper function of sperm basal bodies. (A–C) Sas-6 (green) on BBs (Ana1::tdTomato, red) in early/intermediate (left) or late (right) STs. Axonemes, Ac-tub (gray); DNA, DAPI (blue). (A) wt (*asl/TM6*) STs. In early/intermediate STs (left), basal bodies are docked at the nucleus and form axonemes. The nuclei of late STs are elongated and clustered. (B) Remnant centrioles in *asl* mutant STs with “elongated” region of Sas-6 ($n = 16$). ST at all stages were examined. Bracket, distal extension of Ana1 beyond Sas-6. (left) The only remnant centriole (early/intermediate ST) with elongated Sas-6 exhibiting normal nuclear attachment and axoneme nucleation. (right) Representative remnant ST centriole not attached to a nucleus or forming an axoneme. The nuclei of late stage *asl* STs are fragmented and dispersed (yellow arrowheads). (C) Remnant centrioles in *asl* mutants with a “normal” distribution of Sas-6 (restricted to the very proximal end; $n = 101$). (left) Representative remnant centriole in an early/intermediate ST exhibiting normal nuclear attachment and axoneme formation. (middle) Representative remnant centriole in an early/intermediate ST with a disrupted, flared axoneme (arrow). (right) Representative remnant centrioles in late STs not attached to nuclei or nucleating axonemes. Yellow arrowhead indicates nucleus that has not undergone reshaping. Additional examples are shown in Fig. S5. Bars: 5 μ m; (insets) 1 μ m. Defective, no axoneme or no nuclear attachment or both.

proteins in each cell. In this case, limiting the length of the mother centriole would preserve materials needed for growth of the daughter. Yet another possibility is that regulating MTOC size is critical in cells that require movement and positioning of the centrioles, such as in cells dividing asymmetrically or forming cilia.

The effect of Asl loss is not limited to centriole structure; it also influences the distribution of PCM proteins (Fig. 10 A), but not in the predicted way. Based on its position on the outside of the centriole and antibody interference experiments in *Drosophila* embryos, Asl was suggested to be required for PCM recruitment (Conduit et al., 2010, 2014; Mennella et al., 2012). We found that PCM proteins were recruited to Asl-free centrioles in mitotic mGSCs and meiotic SCs, which can organize MTs and build a functional spindle. Interestingly, the location of PCM assembly corresponded to the position of proximal centriole proteins, even in cases where the proximalized *asl* mutant centrioles were $>2 \mu$ m long. An intriguing hypothesis, with broad implications in centrosome assembly, is that the organization of the internal centriole proteins helps establish the position of the PCM through an unknown inside-out communication mechanism. There is precedence for such a mechanism. Previous work suggested that Sas-4/CPAP might help tether PCM proteins located on the outside of the centriole to the centriole via its interactions with Cep135 and Ana2, located inside the centriole (Zheng et al., 2014). Testing this model will be an exciting future direction that will involve the identification and

analysis of the physical links between the centriole core and the PCM, a link that must span the MTs of the centriole wall.

We also identified a role for Asl in sperm BB function. In the absence of Asl in STs, we found significant defects in the attachment of the BB to the nucleus as well as in the assembly and/or maintenance of the flagellar axoneme (Fig. 10 D). Our initial hypothesis was that these defects were a result of the longer, proximalized *asl* mutant centrioles, but our data proved otherwise. We believe these defects uncover a unique and independent role for Asl during the final stages of sperm development. Asl is associated with the outer surface of BB throughout spermatogenesis, and we show it undergoes a dramatic change in localization during the final stages of ST development (Fig. S5 D). The localization of Asl during spermatogenesis is reminiscent of the “centriolar adjunct,” which is wrapped around the length of the basal body in early STs and becomes a ring structure during elongation (Tates, 1971). The defects in flagella assembly seen in *asl* could be the result of defects in the assembly, organization or maintenance of the centriolar adjunct and, thus, the axoneme. The scarcity of BBs in *asl* testis makes a detailed understanding of the role of Asl in flagella assembly especially challenging. Future identification of interactions of Asl with proteins involved in flagella assembly and the generation of separation of function mutations in Asl that retain centriole duplication function, but lose BB functions, will be required to further our understanding of this novel role for Asl.

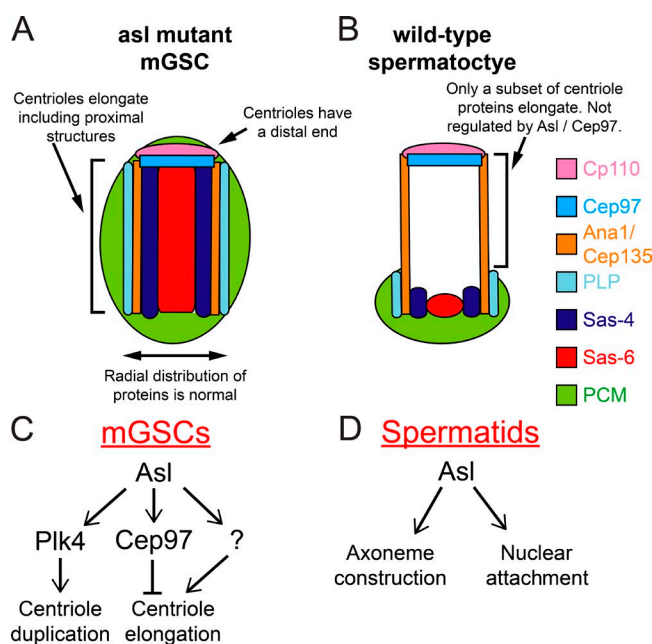


Figure 10. Model for Asl function in centriole organization and length control. (A and B) Not all long centrioles are alike. Centrioles in *asl* mutant mGSCs (A) are organized in a manner distinct from the giant centrioles normally found in SCs (B). This indicates that these two structures, although of similar scale, arise via distinct mechanisms. (C and D) Expanded roles for Asterless at the centriole. In the male germline, Asl plays a critical role in regulating centrosome duplication via Plk4, and controlling centriole length upstream of, and likely directly through, Cep97. (D) In developing STs, Asl is required for normal nuclear attachment and axoneme formation.

Collectively, we have shown that Asl has roles additional to recruiting Plk4 for centriole duplication. Asl functions to control centriole length and the distribution of proximal centriole characteristics and to ensure proper BB functions during spermatogenesis. Insight into additional roles beyond duplication could help shed light on how specific lesions within centriole proteins manifest human diseases. For example, how does the missense mutation Q265P in Cep152 (the human Asl orthologue) lead to the development of human microcephaly (Guernsey et al., 2010)? What exactly does this lesion do to protein function? Our work expands the known roles of Asl, thus allowing one to hypothesize and test which functions are disrupted in specific disease mutations.

Materials and methods

Fly stocks

The following *Drosophila* lines were generous gifts: UAS-Ana1::tdTomato and UAS-Cep135::GFP are transgenic lines with ~2 kb of upstream regulatory elements fused with the cDNA and sequence encoding fluorescent proteins all downstream of the yeast UAS sequence (T. Avidor-Reiss, University of Toledo, Toledo, OH; Blachon et al., 2008). Unc::GFP is under the control of its upstream regulatory elements (Baker et al., 2004). GFP::α-tubulin (T. Avidor-Reiss) and Sas-4::GFP (J. Raff, University of Oxford, Oxford, England, UK; Peel et al., 2007) contain cDNAs, fused to sequences encoding GFP under the control of the ubiquitin promoter. Act5C-Gal4 drives expression of yeast Gal4 under the control of the Act5C promoter (stock #3954; Bloomington *Drosophila* Stock Center). *asl^{mecD}* (T. Avidor-Reiss; Blachon et al., 2008) and *asl²* (M. Gatti, Sapienza University of Rome,

Rome, Italy; Bonaccorsi et al., 1998) are loss-of-function alleles of *asl*. *Df(3R)ED5177* is a deletion that removes the region surrounding *asl* (stock #8103; Bloomington *Drosophila* Stock Center). GFP::Sas-6 BAC carries an insertion from a BAC containing 20 kb of genomic sequence surrounding Sas-6 with GFP sequence recombined in between the start codon of Sas-6 and the rest of its sequence (Lerit and Rusan, 2013). *cep97^{LL01167}* (*cep97^{LL}*; Drosophila Genetic Resource Center at Kyoto Institute of Technology) is a piggyBAC insertion in the *cep97* locus (Schuldiner et al., 2008). This transposable element has strong splice acceptors followed by stop codons in all three reading frames in both orientations. Because the first exon of *cep97^{LL01167}* only contains two complete codons, this insertion should lead to a significant reduction in the longest isoform of the protein, Cep97-PB. *Cep97^{LL}/Df(2L)A1* has long centrioles more frequently than *Cep97^{LL}/Cep97^{LL}*, indicating that *cep97^{LL01167}* is not a genetic null but rather a strong hypomorphic mutant. For *plk4* mutant experiments, *plk4^{c06612}* (a piggyback insertion in the *plk4* locus)/*Df(3L)Pc-2q* (a deletion removing the region surrounding *plk4*) was used as previously described (Bettencourt-Dias et al., 2005).

Transgenic lines were generated by BestGene, Inc. GFP::Cep97, CP110::GFP, and GFP::PACT are under the control of the ubiquitin promoter and constructed in pUGW or pUWG (*Drosophila* Genomics Resource Center). *Cep97^{PACT}* was made by introducing the sequence encoding aa 2,687–2,796 of PLP-PF onto the 5' end of *cep97* cDNA, which was then introduced into pUGW to generate a GFP fusion.

Df(2L)A1 is a Flippase recognition target-mediated deletion (Parks et al., 2004) generated using P[XP]capu^{d05064} and PBac[WH]f02748 (Thibault et al., 2004; the Exelixis Collection at the Harvard Medical School). The removal of the sequence in between the insertions was confirmed by PCR of genomic DNA from *Df(2L)A1* followed by DNA sequencing. Both *cep97^{LL}/cep97^{LL}* and *cep97^{LL}/Df(2L)A1* individuals develop to adulthood. In the case of both heterozygotes and hemizygotes, we found long centrioles less frequently when *cep97^{LL}* heterozygous mothers were used then with *cep97^{LL}* homozygous mothers, suggesting a significant maternal effect. All experiments with *Cep97^{LL}* were performed using individuals derived from homozygous *Cep97^{LL00167}/Cep97^{LL00167}* mothers.

Immunofluorescence methods

Testes, ovaries, and wing discs from the indicated life stage were dissected in *Drosophila* S2 media and fixed in 9% formaldehyde in PBS for 20 min. Samples were washed three times in PBS + 0.3% Triton X-100 (PBST) and blocked for at least 1 h in PBST + 5% normal goat serum. Primary antibodies were incubated with testes overnight at 4°C followed by three 10-min washes in PBST. Secondary antibodies were incubated 4–8 h at room temperature followed by three 10-min washes in PBST. Staining with anti-Ac-tub antibody on pupal testis was performed, with the following modification. Before fixation, testes were treated with 100 μM colchicine in *Drosophila* Schneider's medium for 30 min. Testes from pharate adults were prepared as described previously (Franz et al., 2013) with the following modification. Incubation in 45% and 60% acetic acid was omitted after fixation and testes were immediately transferred to coverslip for squashing. Samples were mounted under a No. 1.5 coverslip in Aquapoly-mount (Polysciences, Inc.) for conventional microscopy or Vectashield (Vector Laboratories) for SIM.

The following primary antibodies were used: guinea pig anti-Asl (gift of G. Rogers, University of Arizona, Tucson, AZ; 1:3,000), mouse anti-Ac-tub (1:500; T6793; Sigma-Aldrich), affinity-purified rabbit anti-PLP (1:4,000; gift of G. Rogers; University of Arizona, Tucson, AZ), rabbit anti-Cnn (1:3,000; this study, raised against aa 698–1,148), rabbit anti-Spd2 (1:2,000, gift of M. Gatti; Giansanti et al.,

2008), mouse anti- γ -tubulin (1:500, T6556, GTU88; Sigma-Aldrich), mouse anti- β -tubulin (1:1,000, MAB3408; EMD Millipore). Secondary antibodies were Alexa Fluor 488, 568, or 647 conjugated (Thermo Fisher Scientific) 1:1500 in PBST plus 5% normal goat serum. DAPI (1:1,000) and Alexa Fluor 488- or 568-conjugated phalloidin (Thermo Fisher Scientific) 1:1,000 were added to some samples.

Microscopy and image analysis

Confocal microscopy was performed on an Eclipse Ti (Nikon) with a 100 \times /1.4 NA objective, a spinning disc confocal head (Visitech International), an interline-transfer cooled charge-coupled device camera (CoolSNAP HQ2; Photometrics), and illumination lasers at 405, 491, 561, and 642 nm. The microscope was controlled by and images were acquired using MetaMorph (Molecular Devices). SIM images were collected using an OMX4 (GE Healthcare). All data analysis was performed using ImageJ (National Institutes of Health). Length and width measurements were performed using line scans across centrosomes and the half maximal width of the line scan is presented. Centrioles were considered long when the measurement exceeded the mean measurement of the control plus 2 \times SD. For intensity measurements, images were collected to avoid saturation and ensure that all pixel intensities fell within the linear response range of the camera. Confocal slices encompassing the volume of the centrosome were summed. The intensities presented are the pixel intensity of a region encompassing the centrosome minus the pixel intensity of an adjacent region of the cell of the same area. Data are normalized to mean intensity of centrosomes in control cells of a single day of imaging.

For live imaging of testes, larval testes were dissected in *Drosophila* S2 media. Whole testes were mounted in a drop of the same media, surrounded by Halocarbon oil 700 (Sigma-Aldrich) on a 50-mm lummo dish (Sarstedt), and covered with a #1.5 coverslip. Live images were acquired on a similar microscope as fixed images equipped with a stage incubator (Binomic System BC-110 Controller; 20/20 Technologies) set to 25°C, a 100 \times /1.49 NA objective, and an ORCA-Flash 4.0 CMOS camera (Hamamatsu Photonics). Stacks, over a 10–15- μ m range, were collected at 1- μ m intervals at 60- or 120-s time intervals. Images in supplemental videos are projections of multiple confocal stacks, displayed at 2 min intervals at 7 frames/s. Mitosis in mGSCs with remnant centrioles is a rare event. mGSCs are in mitosis, which lasts \sim 10 min, about once a day. As such, \sim 1% of mGSCs in a population are in mitosis. Each testis only has \sim 10 mGSCs, and often only three or four are in a good position for live-cell imaging when the testis is mounted. As reported above, only \sim 40% of mGSCs in larvae have a remnant centriole. Imaging dozens of testis for periods of 10–18 h was required to obtain only a few examples. We present representative examples of this small sample.

IP

For IP experiments, fusion proteins were expressed from pUGW and pFAHW plasmids. One well of a six-well plate of S2 cells at \sim 50% confluence was transfected with 0.5 μ g DNA using Effectene (QIAGEN) and allowed to recover for 48 h at 25°C. All subsequent steps were performed at 4°C. 50 μ l of Protein A-conjugated Dynabeads (Thermo Fisher Scientific) was used per IP and resuspended in 1 ml PBS plus 0.01% Tween 20. 1 μ l rabbit anti-GFP antibody (AB290; Abcam) was added and incubated with mixing for 1 h. Cells were harvested and lysed by resuspending in 1 ml RIPA buffer with 1 mM DTT, 1 μ g/ml pepstatin A, 1 μ g/ml leupeptin, and 1 mM PMSF. Lysate was cleared by a 5-min spin in a microfuge at full speed, then incubated with antibody-conjugated beads for 1 h. Beads were washed three times for 5 min each in RIPA buffer. Before the last wash, the beads were moved to a fresh tube. Beads were eluted by boiling in 25 μ l of 2 \times Laemmli

buffer for 5 min, and 10- μ l samples were then analyzed by SDS-PAGE followed by Western blot.

Western blotting

For analysis of Asl protein levels in testes, whole testes dissected from young adult flies of the appropriate genotype were lysed in 2 \times Laemmli buffer (2 μ l/testis) using a microfuge pestle. Samples were boiled for 5 min and SDS-PAGE performed followed by wet transfer to nitrocellulose or polyvinylidene fluoride membrane. After transfer, membranes were blocked with 5% milk in TBS plus 0.05% Tween 20. The following primary antibodies were used: guinea pig anti-Asl (1:5,000; gift from G. Rogers), mouse anti-GFP (JL8, Takara Bio Inc.; 1:10,000), mouse anti-FLAG M2 (1:1,000; Sigma-Aldrich), mouse anti-A ψ -tub (1:5,000, DM1A; Sigma-Aldrich), and chicken anti-Cep97 (raised against the entire protein, this study, 1:2,000). HRP-linked secondary antibodies (1:5,000) were used and SuperSignal West Dura Extended Duration Substrate (1:5,000; Thermo Fisher Scientific) was used for detection.

Y2H screening

Y2H screening to test the interaction between Asl and Cep97 fragments was performed exactly as described previously (Galletta et al., 2014). In brief, the indicated protein fragments were introduced into pDEST-PGADT7 (Rossignol et al., 2007) and pDEST-pGBKT7, modified to carry ampicillin resistance (Galletta et al., 2014) using the Gateway cloning system (Thermo Fisher Scientific). Clones in pGADT7 and pGBKT7 were transformed into Y187 and Y2HGold, respectively (Takara Bio Inc.). Liquid cultures of yeast carrying these plasmids were grown overnight at 30°C in SD –Leu or SD –Trp media, as appropriate, to maintain plasmid selection to an OD₆₀₀ \sim 0.5. Yeast strains of opposite mating types were mated by mixing 20 μ l of a culture of a Y187 strain with 20 μ l of a culture of a Y2HGold strain in 100 μ l 2 \times YPAD media in a well of a 96-well plate. Mating was performed overnight at 30°C with shaking. Mating cultures were transferred to SD –Leu –Trp (DDO) plates using a Multi-Blot Replicator (V&P Scientific). DDO plates were grown for 5 d at 30°C. DDO plates were replica plated to DDO plates, QDO plates (SD –Ade –His –Leu –Trp), DDOXA plates (SD –Leu –Trp with Aurobasidin A [Takara Bio Inc.] and X- α -Gal [Gold Biotechnology]) and QDOXA plates (SD –Ade –His –Leu –Trp with Aurobasidin A and X- α -Gal). Plates were grown for 5 d at 30°C and scored for growth and development of color as appropriate. All Y2H constructs were tested for their ability to activate Y2H reporters in the absence of any binding partner (autoactivation).

The reverse two-hybrid screen to identify mutations in Asl (aa 626–994) that abrogate the Cep97 interaction was performed as described previously (Bennett et al., 2004) with significant modifications described elsewhere (Galletta and Rusan, 2015; Klebba et al., 2015). In brief, Asl (aa 626–994) in pGBKT7 was amplified using low-fidelity PCR conditions (Taq polymerase, in the supplied buffer with 0.05 mM MnCl₂, 0.25 mM dCTP, 0.25 mM dGTP, 0.25 mM dTTP, and 0.06 mM dATP; New England Biolabs, Inc.) and primers designed to complement sequence in the vector (5'-TAATACGACTCACTATAGGGCG-3' and 5'-CGGAATTAGCTTGGCTGC-3'). Amplified PCR products and pGBKT7, linearized with EcoRI and PstI, were gel purified using DEAE cellulose and cotransformed into Y2HGold yeast using standard methods. Yeast carrying the reconstituted plasmid were selected on SD –Trp plates. Approximately 1,400 independent clones containing randomly mutagenized Asl (aa 626–994) were obtained and then individually crossed to Y187 strains carrying Cep97 (aa 1–351) and Cep97 (514–806) in pGADT7. Diploids carrying bait and prey plasmids were selected by growth on DDO plates after 5 d at 30°C. Diploids were replica plated to DDO and QDOXA, as above, and clones that grew

on DDO but failed to grow on QDOXA were selected. These clones were then counterscreened for their interaction with the following Asl interacting fragments (Fig. S5 A): Asl (aa 358–625), Asl (aa 626–994), CP110 (aa 1–325), Plk4 (aa 382–602), Sas-4 (aa 1–347), Sas-4 (aa 348–707), and Spd-2 (aa 664–1,146). No clones disrupted both of the Cep97 interactions without significantly disrupting other interactions.

Statistical analysis

Unpaired *t* tests, with Welch's correction when appropriate, one-way analysis of variance followed by Tukey's multiple comparisons tests, or Fisher's exact tests were used for all statistical comparisons as indicated in figure legends. All statistical analysis was performed with Excel (Microsoft) or Prism (GraphPad Software).

Online supplemental material

Fig. S1 shows an immunoblot demonstrating loss of full-length Asl in *asl^{mecD}*, images and schematics of the centrioles in whole testis from wt and *asl^{mecD}* and demonstration that centrioles in *asl^{mecD}* do not have Asl by immunofluorescence. Fig. S2 shows long centrioles in other allelic combinations of *asl* mutants, the localization of PCM proteins to centrioles during interphase in *asl* mutants, and stills from Video 3. Fig. S3 shows a western blot demonstrating the loss of Cep97 protein in the mutants used in this study, micrographs showing long centrioles in additional allelic combinations of *cep97* mutants and the complete Y2H data demonstrating that Asl and Cep97 directly interact. Fig. S4 shows data that Cep97^{PACT} also rescues centriole length in pharate adult *asl* testis and includes controls for these experiments and those in Fig. 8. Fig. S5 shows examples of reverse Y2H mutants in Asl that failed to disrupt both Asl–Cep97 interactions and not any other Asl interactions. This figure shows additional examples of defects in remnant centrioles/basal body function in *asl* spermatids like those show in Fig. 9. Video 1 shows mitosis in a wt mGSC. Video 2 shows mitosis in an *asl* mGSC with a single, elongated centriole. Video 3 shows mitosis in an *asl* mGSC with no centrioles. Online supplemental material is available at <http://www.jcb.org/cgi/content/full/jcb.201501120/DC1>.

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