Interphase centrosome organization by the PLP-Cnn scaffold is required for centrosome function

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Pericentriolar material (PCM) mediates the microtubule (MT) nucleation and anchoring activity of centrosomes. A scaffold organized by Centrosomin (Cnn) serves to ensure proper PCM architecture and functional changes in centrosome activity with each cell cycle. Here, we investigate the mechanisms that spatially restrict and temporally coordinate centrosome scaffold formation. Focusing on the mitotic-to-interphase transition in Drosophila melanogaster embryos, we show that the elaboration of the interphase Cnn scaffold defines a major structural rearrangement of the centrosome. We identify an unprecedented role for Pericentrin-like protein (PLP), which localizes to the tips of extended Cnn flares, to maintain robust interphase centrosome activity and promote the formation of interphase MT asters required for normal nuclear spacing, centrosome segregation, and compartmentalization of the syncytial embryo. Our data reveal that Cnn and PLP directly interact at two defined sites to coordinate the cell cycle–dependent rearrangement and scaffolding activity of the centrosome to permit normal centrosome organization, cell division, and embryonic viability.

Introduction

Centrosomes are composed of a pair of centrioles embedded in pericentriolar material (PCM) and function as microtubule (MT) organizing centers (MTOCs; Gould and Borisy, 1977). In mitosis, centrosomes organize the bipolar spindle, while in interphase they direct cell migration, traffic cargoes, and build cilia (Doxsey et al., 2005). These functional changes are linked to oscillations in PCM levels. Centrosomes gain MTOC activity by increasing PCM levels, or maturing, before mitosis. The process is then reversed during mitotic exit (Khodjakov and Rieder, 1999; Palazzo et al., 2000). Elucidating the regulation of PCM dynamics is critical to understanding how the centrosome function is normally modulated and deregulated in disease (Nigg and Raff, 2009).

Super-resolution microscopy has revised our view of PCM from an amorphous cloud to a structured architecture (Fu and Glover, 2012; Lawo et al., 2012; Mennella et al., 2012; Sonnen et al., 2012). PCM organization into distinct zones appears conserved across taxa (Lüders, 2012; Mennella et al., 2014), and some proteins, such as Pericentrin (Pcnt)-like protein (PLP; Kawaguchi and Zheng, 2004; Martinez-Campos et al., 2004), and its mammalian orthologue, Pcnt, radially extend across zones (Lawo et al., 2012; Mennella et al., 2012). Understanding how proteins function within these subdomains is key to understanding the cell cycle dynamics, regulation, and function of PCM.

One question that emerges from the discovery of the PCM organization is the identification of the molecular glue, or scaffold, that holds the structure together. A centrosome scaffold was first proposed upon resolving Pcnt and γ-Tubulin (γTub) to a reticular, tubelike lattice (Dictenberg et al., 1998). Much of our understanding of the scaffold comes from studies of the syncytial Drosophila melanogaster embryo, where a constitutively active MTOC serves several essential functions: proper nuclear migration/spacing, actin organization, rapid progression through abridged nuclear cycles (NCs) that lack gap phases, and cellularization (Callaini and Riparbelli, 1990; Rothwell and Sullivan, 2000). Increasing evidence suggests that Centrosomin (Cnn) forms an oligomerized scaffold required to recruit other PCM proteins (Megraw et al., 1999; Vaizel-Ohayon and Schejter, 1999; Zhang and Megraw, 2007; Kao and Megraw, 2009; Conduit et al., 2010, 2014a,b). Thus far, a conserved motif at the N terminus of Cnn (CM1; Zhang and Megraw, 2007), a direct interaction between Cnn and Spd2 (Conduit et al., 2014b), and phosphorylation of Cnn by Polo kinase (Conduit et al., 2014a) have all been implicated in Cnn scaffold assembly. However, how the Cnn scaffold efficiently assembles with each rapid NC remains poorly understood.

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Abbreviations used in this paper: Asl, Asterless; Cnn, Centrosomin; γTub, γ-Tubulin; MT, microtubule; MTOC, MT organizing center; NB, neuroblast; NC, nuclear cycle; NUF, nuclear fallout; PCM, pericentriolar material; Pcnt, Pericentrin; PLP, Pcnt-like protein; SIM, structured illumination microscopy; Y2H, yeast two-hybrid; WT, wild type.

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In humans, mutations in Pcnt and the Cnn orthologue, Cdk5rap2/Cep215, are associated with microcephaly (Bond et al., 2005; Rauch et al., 2008). Because a functional interaction between Pcnt and Cep215 may provide a mechanistic link between these disorders (Buchman et al., 2010), further understanding the interplay between these molecules may contribute to our understanding of disease etiology. Work in *Drosophila* larval neuroblasts (NBs) shows that PLP plays a minor role in organizing Cnn (Martinez-Campos et al., 2004; Galletta et al., 2014); however, a functional role for PLP has not been examined outside of NBs. Thus, it is currently unknown if PLP is required to organize the Cnn scaffold in embryos.

We use live imaging and structured illumination microscopy (SIM) to detail a cell cycle–dependent reorganization of the embryo centrosome, where PCM is expanded during interphase and compact during mitosis. We identify interphase-specific PLP satellites, novel structures that localize exclusively to the tips of Cnn flares. Furthermore, we show a Cnn and PLP direct interaction at two domains. Mutant analysis supports a role for PLP in coordinating the cell cycle–dependent rearrangement of the Cnn scaffold that is essential for normal centrosome function. Our data indicate that proper interphase centrosome organization by the PLP-Cnn scaffold is required for centrosome separation, mitotic progression, genome stability, and embryonic viability.

**Results**

Centrosomes remain active throughout interphase in the early *Drosophila* embryo

Centrosome composition, organization, and activity undergo striking cell cycle oscillations (Palazzo et al., 2000). Previous work has identified Cnn as a key regulator of centrosome dynamics through its scaffolding of PCM proteins (Megraw et al., 1999; Vaizel-Ohayon and Schejter, 1999; Terada et al., 2003; Lucas and Raff, 2007; Zhang and Megraw, 2007; Dobbelkaere et al., 2008; Conduit et al., 2010, 2014a,b). At embryonic centrosomes, Cnn dramatically rearranges from a compact sphere in mitosis to an expanded interphase structure that is defined by a central bolus of PCM and extended radial fibers (Li and Kaufman, 1996; Megraw et al., 2002); for clarity, we refer to these fibers as Cnn flares. In addition, Cnn particles are released into the cytoplasm primarily during interphase (Megraw et al., 2002). This leads to an attractive model whereby the cell cycle–dependent modulation of Cnn organization directly regulates centrosome activity. However, mechanisms regulating this reorganization remain poorly understood, and the physiological significance of centrosome shape changes is currently unknown.

To study Cnn during the rapid early divisions, it is critical to precisely track the timing of its reorganization. By imaging recombinerred Cnn in embryos coexpressing the nuclear marker H2A-RFP, we find that Cnn is a compact structure during mitosis (Fig. 1 A and Video 1). Upon mitotic exit and throughout interphase, Cnn becomes expansive and particulate, ultimately extending reticulated flares that greatly increase centrosome volume and eject particles into the cytoplasm (Fig. 1 A, 4:06). These findings are in agreement with a seminal study of Cnn dynamics (Megraw et al., 2002). To monitor the conspicuous changes to centrosome size and shape, we visualized Cnn distribution by SIM. SIM resolves mitotic Cnn as a hollow sphere with a discernable strand-like substructure (Fig. 1 B–B′) with few cytoplasmic particles (Fig. 1 B″). In contrast, interphase centrosomes form a more elaborate shape with extended flares of uneven thickness and length (Fig. 1, C–C″). These flares are sites of Cnn particle release (Fig. 1 C″), which appear analogous to mammalian centriole satellites (Rattner, 1992; Balczon et al., 1994; Zhang and Megraw, 2007), as both are ejected from the centrosome and bi-directionally move during interphase (Kubo et al., 1999; Megraw et al., 2002). In addition, SIM resolves Cnn particles as rings, similar to the structure of mammalian centriole satellites revealed by EM (de-Thé, 1964; Kubo et al., 1999). Although mammalian centriole satellites facilitate protein trafficking to and from centrosomes and cilia, the overall functional significance of mammalian satellites requires further study (Mahjoub and Tsou, 2013).

To observe the mitotic-to-interphase transition of Cnn in finer detail, we imaged embryos with greater temporal resolution. Unexpectedly, the compact mitotic centrosome gradually appears to unfurl as flares seemingly unfold and extend from the center mass (Fig. 1 D and Video 2), ultimately forming an interphase centrosome lattice. Retention of Cnn at the interphase centrosome is consistent with the constitutive activity of embryonic centrosomes throughout the cell cycle, which is thought to permit efficient NC progression (Callaini and Riparbelli, 1990). These findings suggest that embryonic Cnn does not follow the paradigm of PCM shedding and total centrosome dematuration during mitotic exit, and only a small amount of Cnn is released into the cytoplasm as particles. Understanding this rearrangement and how Cnn remains anchored at the centrosome is fundamental to understanding how centrosome function is modulated during the rapid divisions of early development.

Centrosomes add a flare zone in interphase

Recent studies using SIM to define the PCM structure in cultured cells revealed that the ostensibly amorphous PCM is, indeed, organized into zones (Fu and Glover, 2012; Lawo et al., 2012; Mennella et al., 2012; Sonnen et al., 2012). To understand how interphase and mitotic centrosome shape is determined, we conducted a SIM-based survey to demarcate centrosome zones in the early *Drosophila* embryo.

As expected, we found that the distributions of centriole proteins, such as SAS6 and Asterless (Asl), remain constant in mitosis and interphase (Fig. 2, A and B). These data were used to define a centriole zone with an ~200-nm radius (Fig. 2 B, blue shading; and Fig. S1, A and B). In contrast, the PCM zone, defined by γTub, expands from a radius of 450 nm in mitosis to 600 nm in interphase (Fig. 2 B, orange shading; and Fig. S1, A and B). Thus, the outer edge of the PCM zone expands as embryos enter interphase. The major structural change to the interphase centrosome is the addition of extensive Cnn flares, which protrude well beyond the PCM zone. We term this zone the interphase flare zone (Fig. 2 B, brown shading; and Fig. S1, A and B). Notably, these interphase-specific flares extend ~1,380 nm, with some reaching >2 μm, more than twice as far as the PCM zone (Figs. 2 B and S1 A). In sum, our analysis of embryonic centrosome organization shows that interphase centrosomes form a distinct flare zone in addition to the centriole and PCM zones (Fig. 2 B′). Moreover, our imaging (Fig. 1 C) reveals that the mitotic Cnn bolus appears to unfold to provide the source of interphase Cnn flares.

**Interphase PLP satellites define the margin of interphase Cnn flares**

Reasoning that factors enriched in the interphase flare zone might regulate or contribute to Cnn scaffold function, we as-
Figure 1. Rearrangement of Cnn at the mitotic-to-interphase transition. (A) Live Cnn-GFP and H2A-RFP in a pseudo-cell (broken line) of a WT embryo through three cell cycles. Arrows show the enlarged regions below (Cnn, white). Cnn flares (red arrowheads) and separating daughter centrosomes (white arrowheads) in interphase are indicated. Bars in top panels, 5 µm. (B and C) SIM images of mitotic (B) and interphase (C) embryos stained for...
sayed the localization of many other centrosome proteins (Fig. 2 C). As expected, Sas6, Asl, and the centriole marker, PACT-GFP, are restricted to the centriole zone. Sas4 and Bld10/Cep135 are also enriched in the centriole zone, but show low levels at discrete foci within the flare zone (Fig. S1 C, enhanced image). Polo, a key regulator of maturation, radially extends into the PCM zone, while low levels of PLK4/SAK, a key regulator of centriole duplication, localize to small foci within the flare zone (Figs. 2 C and S1 C). Because we detect only low levels of Sas4, Cep135, and PLK4 in the interphase flare zone upon overexpression of GFP fusion constructs, and could not verify the endogenous localization of these proteins, we cannot speculate on their function within this zone.

The defining molecule of the PCM zone, γTub, is found at low levels in the interphase flare zone (Fig. 2, A and C). However, even at a single centrosome, γTub levels vary greatly between flares (Fig. 2 A, arrows). This heterogeneity is also true for TACC (Fig. 2 C), as previously reported (Megraw et al., 2002). We find endogenous Spd2 primarily within the centriole and PCM zones (Figs. 2 C and S1 C), in agreement with GFP-tagged Spd2 localization at centrioles and PCM with low amounts in “peripheral” areas (Conduit et al., 2014b), which we presume is the flare zone.

In contrast, we find endogenous PLP at high levels in nearly every Cnn flare (94.7%; n = 190 flares, 82 centrosomes) in addition to its known centriole localization, which suggests that PLP may coordinate centrosome shape changes with Cnn, while almost no PLP is detected in the PCM zone (Figs. 2, B–D’). More than two decades ago, Pcnt was localized to “PCM-like bodies (satellites)” in mammalian cells using immuno-EM (see Fig. 4; Doxsey et al., 1994). Subsequent work proved that Pcnt forms a biochemical complex with the canonical mammalian centriole satellite marker, PCM-1 (Li et al., 2001; Dammermann and Merdes, 2002). Although a functional orthologue of PCM-1 has not been identified in Drosophila, the discrete localization of PLP foci at flare tips is reminiscent of satellites orbiting a central body (the centriole); therefore, we refer to PLP localized within the interphase flare zone as PLP satellites.

PLP satellites are found at the tips of Cnn flares (Fig. 2, D and D’) and extend ~1,208 nm from the centriole center (Figs. 2 B and S1 A), 10-fold farther than the predicted ~110-nm length of a PLP molecule (Mennella et al., 2012). Thus, PLP localizes to two distinct pools (centrioles and satellites) separated by a considerable distance. SIM resolves PLP satellites as rings with a diameter of 187 ± 22.9 nm (Fig. 2 D, n = 40). Based on the unique localization of PLP satellites to the tips of the extended interphase centrosome, we hypothesized that PLP functions as a molecular scaffold that stabilizes Cnn flares and contributes to centrosome shape and, in turn, centrosome function.

**PLP satellites and Cnn flares share similar dynamics and coassemble**

To investigate the function of PLP satellites in detail, we examined live embryos expressing full-length PLP-GFP (PLPFL), which mirrors endogenous PLP distribution (Fig. 3, A and B) and fully rescues plp− viability (Galletta et al., 2014). Live imaging shows that PLPFL remains closely apposed to centrioles throughout the cell cycle, but expands into the flare zone as PLP satellites in interphase (30/31 embryos; Fig. 3 C and Video 3). Satellites diminish upon nuclear envelope breakdown (Fig. 3 C, 3:30), are absent in mitosis (Fig. 3 C, 7:00), reemerge during interphase when centrosome separation (Fig. 3 C, 10:30), and become prominent once centrosomes fully separate (Fig. 3 C, 19:00). Therefore, PLP satellite assembly and disassembly are entrained with the cell cycle where satellite-permissive conditions are present during interphase. Moreover, the timing of PLP satellite addition and removal bears a striking resemblance to that of Cnn flares (Fig. 1 A).

Imaging individual embryos progressing from NC 10 or 11 (early blastoderm) through NC 14 (cellularization) reveals that PLPFL satellites become brighter and more structured with each NC, reaching a peak in NC 14 (n = 5/5 embryos; Fig. 3 C, 0:00 vs. 19:00; Fig. S2 A, 97:30). PLP satellites are no longer detected after cellularization (Fig. S2 A’), once the interphase centrosomes are inactivated (Harris and Peifer, 2007). Thus, PLP satellite formation is also developmentally regulated, suggesting that their function may be critical during the increasingly prolonged interphases of later syncytial NCs (Foe and Alberts, 1983).

To examine PLP satellite dynamics in detail, we imaged PLPFL at higher temporal resolution. This reveals that PLP satellites undergo bidirectional, linear runs (Fig. 3, D–D’; and Video 4) with an average velocity of 0.33 ± 0.13 µm/s (n = 27; Fig. 3 E), which is in agreement with velocities reported for cytoplasmic Cnn flare particles (Megraw et al., 2002). More than 35% (n = 21/58) of PLP satellites underwent directed runs >0.5 µm (Fig. 3 F), with an average run length of 0.77 ± 0.22 µm and some runs exceeding 1.5 µm or interspersed with pauses. We conclude that PLP satellites share many of the same dynamic properties as Cnn.

A model where a PLP–Cnn complex can assemble into a stable scaffold is supported by experiments that show that overexpression of either mammalian Cep215 or Pcnt is sufficient to form a super scaffold that recruits high levels of the other (Lawo et al., 2012; Pagan et al., 2015) and by the biochemical association of PLP and Cnn within embryonic extracts (Conduit et al., 2010; Gopalakrishnan et al., 2011). To investigate whether PLP satellites and Cnn are assembled into a complex, we performed live imaging of PLP-GFP and Cnn-mCherry, which revealed an enrichment of Cnn at flare tips that is coincident with PLP satellites (Figs. 4 A and 2 D). Visualizing the release of a Cnn particle shows that Cnn and PLP are packaged together and co-traffic within the cytoplasm (Fig. 4 A and Video 5). Likewise, mobile cytoplasmic particles containing both Cnn and PLP can associate with existing flares (Fig. 4 B and Video 5). In contrast, live imaging of embryos expressing PLP-GFP and another dynamic PCM component, TACC-RFP, reveals an uneven distribution within PLP satellites (Fig. S2 B). Therefore, Cnn and PLP share a specific relationship within the centrosome and likely function together to direct local changes to its size and activity.

Given these coordinated movements, we sought to test if Cnn flare extension is required for PLP satellite formation. Pre-
viorious studies show that treatment with the MT drug colchicine causes Cnn flares to collapse into a compact, mitotic-like configuration (Megraw et al., 2002). Injecting embryos expressing PLP-GFP and Cnn-mCherry with colchicine results in the concomitant compaction of Cnn and PLP (Fig. S2 C). Thus, while MTs are dispensable for the localization of Cnn to the PCM zone (Megraw et al., 1999; Vaizel-Ohayon and Schejter, 1999) and PLP to the centriole zone (Kawaguchi and Zheng, 2004),
we find that MTs are required for the extension of Cnn and PLP into the interphase flare zone. Another correlation is observed upon examination of mother and daughter centrosomes. It was previously noted that higher levels of Cnn accumulate on the mother as compared with the daughter (Conduit et al., 2010). Similarly, we note that greater amounts of PLP associate with

Figure 3. \( \text{PLP}^{\text{FL, +}} \) localizes to dynamic satellite structures. (A and B) Embryos were stained for the indicated proteins. Arrows show PLP at centrioles (blue) and satellites (brown). The boxed sections are enlarged below. (C) Live \( \text{PLP}^{\text{FL, +}} \) shows interphase PLP satellites (arrowheads) in NC 12 and 13. Time is given in minutes:seconds. (D) Anterograde (green) and retrograde (blue) satellite run relative to centriole (asterisk). Time is given in seconds. (D') Corresponding kymograph and plot of distance over time (D''). (E) Average velocity of directed runs; n.s., not significant; \( n = 27 \) runs. (F) Frequency of satellites with directed \( \geq 0.5 \mu \text{m} \) runs; \( n = 58 \) centrosomes. Bars: (A and B, top) 5 µm; (A and B, bottom) 1 µm; (C) 5 µm; (D) 2 µm.
Cnn and PLP form an interphase scaffold

The larger, Cnn-rich mother centrosome than the smaller daughter centrosome (Fig. S2 D). These data further suggest that Cnn and PLP are coregulated and may cooperate to determine centrosome size and activity.

PLP organizes the Cnn scaffold

To date, a functional role for PLP has not been examined in the Drosophila embryo. However, a recently reported mouse model shows that Pcnt organizes Cep215 and is required for cardiovascular and neural development (Chen et al., 2014). To investigate whether PLP regulates centrosome size or function, and whether this regulation is important for early Drosophila development, we generated plp− null embryos (Fig. S3 A). Loss of PLP leads to 100% lethality; 32% die as embryos and 68% as first instar larva (see Materials and methods), which indicates that PLP is essential for viability. To determine if PLP is required to maintain the Cnn interphase lattice, we visualized Cnn in live wild-type (WT) and plp− embryos. Compared with WT embryos expressing Cnn-GFP (Fig. 5 A and Video 6), plp− centrosomes show Cnn disorganization and dispersal (Fig. 5 B). In plp− embryos, Cnn appears to ooze away from the centrosome, forming extended flares that eject an increased number of cytoplasmic particles. In addition, large voids interrupt the normally uniform distribution of Cnn within the PCM zone (Fig. 5 B, bracket).

To quantify these abnormalities, we imaged endogenous Cnn in plp− embryos (Fig. 5, C and D). Strikingly, Cnn flares in plp− mutants are significantly longer by ~30% (1.80 µm vs. 1.38 µm in WT; Fig. 5 E) with some flares reaching >3 µm. The presence of long Cnn flares and the degree of Cnn disorganization at plp− centrosomes made it difficult to accurately measure Cnn within the PCM zone. Therefore, we also investigated endogenous γTub behavior in plp− embryos, which confirmed the significant PCM disorder (Fig. S3 B, brackets). Live imaging of GFP-γTub throughout the cell cycle further shows that PCM dispersal in plp− mutants is most evident in interphase (Fig. S3, C and D, green boxes; and Video 7); γTub during mitosis appears less disrupted and more similar to controls (Fig. S3, B–D).

Our analysis also revealed a significant ninefold increase in cytoplasmic Cnn particles in interphase plp− embryos (13.8/100 µm² vs. 1.6/100 µm² in WT; Fig. 5, C, D, and F). Moreover, these flares remain in the cytoplasm and fail to consolidate into the centrosome in mitosis (13.5/100 µm² vs. 1.0/100 µm² in WT). Our data show that PLP is required for normal Cnn lattice arrangement and for Cnn retention at interphase centrosomes (Fig. 5, B–D). Our findings that γTub dispersal is more severe in interphase plp− embryos (Fig. S3, B and D), and that PLP satellites are only present in interphase hint that PLP satellites might function as structural elements that physically
scaffold and anchor the PCM to properly determine interphase centrosome shape. This positive regulatory role for PLP-mediated regulation of interphase centrosome activity agrees with results from mammalian and Drosophila cultured cells (Loncarek et al., 2008; Kim and Rhee, 2011; Lawo et al., 2012; Mennella et al., 2012; Wang et al., 2013; Pagan et al., 2015), as well as mitotic Drosophila neuroblasts (Martinez-Campos et al., 2004; Galletta et al., 2014), but contrasts with a negative regulatory role for PLP in interphase neuroblasts (Lerit and Rusan, 2013). Thus, modulation of centrosome activity by PLP is differentially regulated in various cellular contexts.

PLP mediates efficient MT radial organization and centrosome segregation

Expanded localization of Pcnt to centrosomes has previously been correlated with MT organization in mouse epithelial cells (Mogensen et al., 1997), and knockdown of Pcnt is associated with MT disorganization in cultured cells (Zimmerman et al., 2004), indicating that Pcnt contributes to proper MT organization. While immunodepletion experiments previously suggested that PLP plays a role in MT nucleation (Kawaguchi and Zheng, 2004), mutant analysis within NBs indicates that plp cells are efficient MT nucleators (Martinez-Campos et al., 2004; Lerit and Rusan, 2013; Singh et al., 2014). Indeed, similar to cnn mutants (Megraw et al., 1999; Vaizel-Ohayon and Schejter, 1999), we found that embryonic plp centrosomes maintain robust MTOC activity comparable to WT (Fig. 6 A). In addition, the cytoplasmic Cnn particles found in plp mutants (Megraw et al., 1999; Vaizel-Ohayon and Schejter, 1999), we found that embryonic plp centrosomes maintain robust MTOC activity comparable to WT (Fig. 6 A). In addition, the cytoplasmic Cnn particles found in plp− mutants serve as a platform for MT organization independent of the centrosome (80%; n = 33/40 of cytoplasmic foci; Fig. 6 A′), which is consistent with ectopic cytoplasmic Cnn foci organizing MT asters within unfertilized eggs (Conduit et al., 2014a). In interphase, it was difficult to determine if these cytoplasmic particles altered the density of the MT network, but there is a clear reduction in

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Figure 5. PLP organizes the Cnn scaffold. (A and B) Live Cnn-mCherry in WT (A) and plp− (B) embryos in interphase. Released particles (arrows), extending flare (arrowheads), and disrupted PCM (bracket) are shown. Time is given in seconds. (C and D) Embryos stained for Cnn. (E) Cnn flare length in interphase embryos; n > 80 centrosomes. Mean ± SD is indicated. (F) Cytoplasmic Cnn particles in a 100-µm² area [interphase embryos: WT n = 87, plp− n = 18, PLPΔ5 n = 24, cnnB4 n = 21; mitotic embryos: WT n = 40, plp− n = 18, PLPΔ5 n = 9, cnnB4 n = 15]. Data are mean ± SD [error bars]. ***, P < 0.001; ****, P < 0.0001 by a Student's two-tailed t test relative to WT. Data shown are from a single representative experiment out of two or more repeats. Bars: (A and B) 2.5 µm; (C and D) 5 µm.
radial MT symmetry in \textit{plp}^{-} embryos, as evident by MT crossovers and more randomized MTs (55%; \(n = 11/20\) embryos; Fig. 6 B). Similarly, disorganized astral MTs were recently reported in a novel \textit{Pcnt} mouse model (Chen et al., 2014), which suggests that MT organization is a conserved function of PLP. Live imaging of MTs shows that \textit{plp}^{-} mutants (\(n = 3/5\) embryos) display MTOC inactivation followed by failed and abortive spindle formation (Fig. 6 C, broken line), indicating that PLP is required to maintain MTOC activity throughout the cell cycle. Further, loss of PLP also disrupts mitotic spindle orientation (Fig. 6 D), and collisions of neighboring nuclei result in uneven nuclear spacing and association of more than two MTOCs with a single nucleus (Fig. 6 C, arrows; and Video 8).

Given this MT disorganization, we reasoned that centrosome separation, a process known to require properly arranged anti-parallel MTs, would be impaired. Analysis confirms that 41% of \textit{plp}^{-} embryos show centrosome separation defects (vs. 1.9% in WT), resulting in two centrosomes at some mitotic spindle poles (Fig. 6, E and F). In addition, 38% of \textit{plp}^{-} embryos show centrosomes detached from nuclei/chromosomes, an event never seen in WT (Fig. 6, E and F). Similar centrosome segregation and detachment defects occur in embryos deficient in dynein (Robinson et al., 1999), Polo (Archambault et al., 2008), and Cnn (Lucas and Raff, 2007; Zhang and Megraw, 2007). These centrosome separation and detachment defects, coupled with the abnormal mitotic spindles observed in our live imaging, may account for the elevated centrosome segregation defects, where 40% (\(n = 14/31\)) of \textit{plp}^{-} embryos contain pseudo-cells with more than two centrosomes (Fig. 6, G and H).

Defects in centrosome positioning and interphase MT arrangement may contribute to aberrant nuclear spacing and cell cycle length. Indeed, polyplloid pseudo-cells were observed in \approx 20\% of \textit{plp}^{-} embryos (Fig. 6, G and H). Additional analysis shows that cell cycle progression is altered in some cells, as nuclei in \textit{plp}^{-} mutants lose the stereotyped mitotic synchronization characteristic of WT syncytial divisions (Fig. 7, F and F'). Together, these data support a role for PLP in organizing the symmetric, compact PCM necessary for proper MT organization, centrosome separation, and efficient cell cycle progression. Furthermore, the extensive similarities observed in \textit{plp}^{-} and \textit{cnn}^{-} loss-of-function studies further support a model where PLP and Cnn function in a common pathway to modulate centrosome size and activity.

**PLP is required for genome stability**

\textit{Drosophila} embryos have evolved a mechanism, nuclear fall-out (NUF), to eject damaged nuclei from the cortex in order to protect the developing embryo from propagating aberrant chromosomes (Sullivan et al., 1993; Rothwell et al., 1998; Takada et al., 2003). In \textit{cnn}^{-} mutant embryos, the accumulation of progressive mitotic failures leads to nuclear collisions, irregular nuclear spacing, and NUF, which prohibit cellularization and lead to embryonic lethality (Megraw et al., 1999; Vaizel-Ohayon and Schejter, 1999; Zhang and Megraw, 2007). To detect chromosome segregation defects and NUF in \textit{plp}^{-} mutants, we imaged successive cell cycles in embryos expressing H2A-RFP. While nuclei synchronously divided without chromosome segregation errors in WT embryos, \textit{plp}^{-} embryos revealed lagging chromosomes followed by NUF (Fig. 7 A and Video 9). Fixed analysis showed that NUF is significantly increased in \textit{plp}^{-} embryos (80% of embryos vs. 16% in WT; Fig. 7, B and C) and is more severe, as more nuclei per embryo are removed from the cortex (13% vs. <2% in WT; Fig. 7 D). Further analysis of ejected nuclei revealed the accumulation of \textit{yH2Av}, a marker of double-stranded DNA breaks (Fig. 7 E), which suggests that chromosome segregation errors trigger DNA damage, leading to NUF. Similarly, NUF is evident in 100% of hypomorphic \textit{cnn}^{-} embryos (\textit{cnn}^{−/−}; \(n = 25\)) with nearly 25% of nuclei removed from the cortex (Fig. 7 D), indicating that both PLP and Cnn function to ensure chromosome fidelity. We speculate that the accumulation of DNA damage and resultant NUF contributes to embryonic lethality.

Previous studies show nuclei that have experienced chromosome missegregation or DNA damage fail to progress through the cell cycle efficiently (Hayashi and Karlseder, 2013; Poulton et al., 2013). In cultured mammalian cells, Pent disruption is linked to aberrant mitotic progression (Tibellius et al., 2009; Wang et al., 2013). Quantification of this defect in fixed embryos indicates there is a 10-fold increase in mitotic asynchrony in \textit{plp}^{-} mutants (45% of embryos vs. <5% in WT; Fig. 7, F and F'). We suspect that these patches of mitotic asynchrony correspond to the patches of NUF observed in \textit{plp}^{-} mutants (Fig. 7 B), as they are of roughly the same size and occur during the same embryonic stages. These data indicate that PLP ensures efficient progression through the cell cycle to confer genomic stability.

**Cnn and PLP directly interact at two distinct domains**

Our collective work strongly suggests that Cnn and PLP function within a complex to regulate centrosome size and activity. To date, a direct interaction between \textit{Cep215}/Cnn and \textit{Pcnt}/PLP has not been reported. Immunoprecipitation experiments from mammalian and \textit{Drosophila} extracts, however, indicate that these proteins comprise a biochemical complex through a conserved motif located in the C terminus of \textit{Cep215}/Cnn (Fig. 8 A), termed Conserved Motif 2 (CM2; Kao and Megraw, 2009; Buchman et al., 2010; Conduit et al., 2010; Wang et al., 2010; Gopalakrishnan et al., 2011; Kim and Rhee, 2014). To test whether Cnn and PLP interact directly, we truncated Cnn and PLP proteins into a series of fragments (Fig. 8 B) and conducted yeast two-hybrid (Y2H) analysis (Fig. 8 C). These studies identified two sites likely to mediate direct interaction. One occurs between Cnn fragment 1 (Cnn-F1) and PLP fragment 5 (PLP-F5; Fig. 8, B and C). A second was detected between Cnn fragment 3 (Cnn-F3) and PLP fragment 2 (PLP-F2; Fig. 8, B and C). Previous work indicates that CM2 interacts directly with Calmodulin (Wang et al., 2010) and Centrocortin (Cen; Megraw et al., 2009), which suggests that this conserved motif mediates several protein interactions. Further Y2H analysis confirms that the CM2 domain within Cnn-F3 is necessary and sufficient to mediate the direct interaction with PLP-F2 (Fig. 8, C and D).

**Cnn and PLP are mutually required for proper localization and function**

The power of \textit{Drosophila} genetics allows us to test the significance of these interactions in the context of an intact organism and to test whether the interactions between Cnn and PLP are required to influence centrosome activity and regulate the cell cycle-dependent rearrangement of the centrosome. To investigate a role for the PLP(F5)–Cnn(F1) interaction in PLP satellite and Cnn flare formation, we generated animals expressing a GFP-tagged PLP transgene that lacks the F5 region. Fixed and live studies of this construct showed it localizes

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Figure 6. PLP is required for MT organization. (A and B) WT and plp⁻ mitotic (A) and interphase (B) embryos stained for the indicated proteins. (A') Acentriolar Cnn particle (inset) organizes MTs. Boxes in B show radial MT array, enlarged in the insets on the right. (C and D) Live GFP-MT in embryos. Broken circles show MTOC inactivation. Arrows show nuclear collisions and the arrowhead shows an orthogonal spindle. (E) Centrosome separation (closed
to centrioles and satellites in both WT and *plp*− backgrounds (PLPΔ5; *plp*Δ, hereafter PLPΔ5; Fig. S4, A–C; and Video 10). The centriole localization was unexpected because PLP-F5 contains the highly conserved PACT domain that is sufficient for centriole targeting (Gillingham and Munro, 2000; Kawaguchi and Zheng, 2004; Martinez-Campos et al., 2004). Our data show that motifs outside of PLP-F5 are sufficient to generate PLP satellites and maintain colocalization with Cnn within flares and cytoplasmic particles.

Notably, PLPΔ5 satellites are less structured and extend farther from the centriole (1.5 ± 0.5 µm vs. 1.2 ± 0.3 µm for PLPΔ1 and 1.2 ± 0.3 µm for endogenous PLP; n = 20 centrosomes). PLPΔ5 also exacerbates the release of Cnn cytoplasmic particles found in *plp*Δ mutants (25/100 µm² vs. 13.8/100 µm² for *plp*Δ and 1.6/100 µm² for WT; Figs. 5 F and S4 D, arrows). Surprisingly, live imaging reveals that PLPΔ5 satellites persist throughout the cell cycle (n = 6/6 embryos; Fig. S4 C and Video 10), which is unlikely due to overexpression of PLP, as PLPΔ5 (Fig. 3 A) or PLPΔ5 in the WT background do not perturb the normal centrosome structures (Fig. S4 A). Importantly, the PLPΔ5 gain-of-function phenotype in mitosis lends support to the hypothesis that PLP satellites act to organize Cnn, because the abnormal mitotic PLPΔ5 satellites also organize abnormal interphase-like Cnn flares (Fig. S4, B and E). Even when only one mitotic centrosome at a nucleus displays PLPΔ5 satellites, it also extends Cnn flares (Fig. S4 D, solid box), whereas the PLP satellite-free centrosome shows compact Cnn that is characteristic of normal mitotic centrosomes (Fig. S4 D, broken box), implicating local regulation of centrosome organization by the PLP–Cnn complex. Overall, these data suggest that PLP-F5 is dispensable for PLP satellite and Cnn flare formation, but is required for their proper organization and mitotic compaction. Furthermore, we conclude that the PLP(F5)–Cnn(F1) interaction is dispensable for PLP–Cnn colocalization, and possibly complex formation.

These data raise the intriguing possibility that the second interaction defined between Cnn CM2 and PLP-F2 may be important for the normal formation of Cnn flares and PLP satellites in interphase. Unfortunately, despite testing 16 truncations of PLP-F2, our efforts to narrow the interaction region were unsuccessful (Fig. S5 A). Likewise, despite screening many candidate transgenic lines, we were unable to generate an animal expressing a PLP transgene that lacks the F2 region (see Materials and methods). Therefore, to abrogate the interaction between Cnn CM2 and PLP-F2, we used a previously published allele of *cnn*, *cnn*Δ5, which is defined by a single point mutation at an invariant arginine residue (R1141H; Fig. 8 A) within CM2 (Vaizel-Ohayon and Schejter, 1999; Kao and Megraw, 2009). Remarkably, introduction of R1141H into Cnn-F3 is sufficient to abolish the interaction with PLP-F2 (Fig. 8, C and D), which suggests that *cnn*Δ5 mutants are particularly useful to probe the significance of the Cnn(CM2)–PLP(F2) interaction in vivo.

Despite producing normal levels of Cnn protein, prior detailed analysis of *cnn*Δ5 embryos indicates pronounced similarities in the PCM dispersion and NUF phenotypes we observe in *plp*Δ mutants (Kao and Megraw, 2009). Consistent with these studies, all interphase *cnn*Δ5 embryos display highly disordered Cnn and γTub (Figs. 9 A and S5 B; n = 34). PLP localization was not previously examined in *cnn*Δ5 mutants. Thus, to assay the significance of the Cnn CM2 interaction with PLP-F2, we examined PLP distribution in control and *cnn*Δ5 embryos. Significantly, localization of PLP to the satellites is eliminated in interphase *cnn*Δ5 embryos, whereas PLP at centrioles is unperturbed (Fig. 9 A; n = 34). The effect of Cnn on PLP satellite formation is specific, as disruption of other PCM factors, such as TACC and MSPS, previously localized to flare-like particles (Lee et al., 2001), did not alter PLP satellite formation (Fig. 9 A).

SIM confirms that interphase centrosomes in *cnn*Δ5 embryos lack PLP satellites but maintain PLP at centrioles (Fig. 9 B), which signifies that the two PLP pools are regulated by distinct mechanisms. Furthermore, we conclude that the Cnn-n(CM2)–PLP(F2) interaction is dispensable for PLP centriole localization but is essential for PLP satellite formation. The lack of PLP satellites in *cnn*Δ5 embryos corresponds with a loss of Cnn flares and an increase in Cnn cytoplasmic particles, similar to the *plp*Δ phenotype (Figs. 5 F and 9 B). While it remains formally possible that the *cnn*Δ5 mutation abrogates the structure of the Cnn molecule and/or its overall function, mitotic *cnn*Δ5 centrosomes are much less disrupted (Kao and Megraw, 2009), suggesting that the pathway that organizes Cnn during mitosis remains intact and that the Cnn molecule is not completely disrupted. Therefore, we propose that PLP satellites function to form a Cnn scaffold in the interphase flare zone that organizes and confines Cnn to interphase centrosomes (Fig. 10). Our data demonstrate that Cnn and PLP are mutually required for the proper localization and function of the other, thereby demonstrating the presence of a positive feedback loop that ensures the proper formation of the interphase centrosome scaffold.

**Discussion**

Our identification of a positive feedback loop between Cnn and PLP that coordinates the cell cycle–dependent reorganization of the centrosome structure adds to the growing recognition that the centrosome is subject to sophisticated layers of regulation to ensure proper function. The interdependence of three major PCM components (Pcnt/PLP, Cep215/Cnn, and Cep192/Spd2) and two kinases (PIK1/Polo and Aurora A) in generating a fully functional mitotic centrosome is well established (for a recent review see Mennella et al., 2014; Fu et al., 2015). However, we demonstrate a novel role for PLP in building an elaborate interphase centrosome scaffold in the early embryo. Our data show that PLP undergoes a major rearrangement that is entrained to the cell cycle and is critical for proper Cnn dynamics, distribution, and functions upstream of γTub.

Interestingly, our imaging of Cnn during mitotic exit is highly suggestive of Cnn forming a reticular, strand-like structure that is compact in mitosis, but unravels as the cell enters interphase to form a giant centrosome with constitutive MTOC activity. The mechanisms that control these Cnn dynamic movements will require further studies, including the use of fluorescence photoconversion methods. What mediates this cell cycle–dependent transition in centrosome architec-
Interphase 
Prophase 
Metaphase 
Anaphase 
Telophase 
Interphase

Figure 7. PLP maintains genome stability. (A) Live H2A-RFP in embryos. Broken circles show mitotic asynchrony. Arrowheads show lagging chromosomes (9:00) followed by NUF (16:00). (B) NUF [broken circle] detected by DAPI. (C and D) The frequency [C] and amount (D) of NUF is quantified. (E) γH2Av labels nuclei (DAPI, blue) ejected from the cortex. Arrowheads show nuclei that have undergone nuclear fallout and stain positive for γ-H2A. The negative sign indicates distance below the embryo surface. (F) Embryos stained with DAPI [red; all nuclei] and pH3 [green; mitotic nuclei] to detect mitotic asynchrony. Results are quantified in F'. Data are mean ± SEM (error bars) for C, all other data are mean ± SD. Time is given in minutes:seconds. *, P < 0.01; ***, P < 0.0001; n.s., not significant. Data shown are from a single representative experiment out of two or more repeats. Bars: (A) 10 μm; (B, E, and F) 20 μm.

Moreover, we uncovered PLP localized to the tip of each Cnn flare. These PLP satellites are distinct from a PCM pool of PLP predicted by FRAP analysis of PACT-GFP (a GFP fusion with a motif from the C terminus of PLP; Martinez-Campos et al., 2004), as we find that PACT-GFP is absent from PLP satellites (Fig. 2 C). Thus, PACT is not an accurate predictor of PLP localization, at least not beyond the centriole pool (see also Fu and Glover, 2012). More recently, FRAP of PLP-GFP also showed nonuniform recovery within the centrosome, but a
Cnn and PLP form an interphase scaffold

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A second major peak corresponding to PLP satellites at distances beyond the PCM zone at 650 nm was not reported (Conduit et al., 2014b). Thus, our work is the first to describe the dynamic behavior of PLP satellites, which define the end of Cnn flares and the outer margin of the interphase centrosome.

Several recent studies support a model of Cnn scaffold formation in Drosophila. Polo phosphorylates Cnn at sites within Cnn-F2 near the centriole wall, which is required for the formation of higher-order Cnn structures (Conduit et al., 2014a). The Cnn scaffold then migrates into the PCM zone (between 200 and 600 nm; Fig. 10). Spd2 is critical for this process (Fu and Glover, 2012; Conduit et al., 2014b), where it functions in a positive feedback loop with Cnn to form a Spd2–Cnn PCM scaffold (Fig. 10, gray box, 1), which is consistent with earlier studies showing that Cep192/Spd2 recruits PCM (Dix and Raff, 2007; Gomez-Ferreria et al., 2007; Giansanti et al., 2008; Haren et al., 2009; Fu and Glover, 2012). Our work suggests that PLP satellites and Cnn form a second, spatially distinct scaffold (Fig. 10, outside of gray box) located in the interphase centrosome flare zone (Fig. 10, 600–1,400 nm). These two scaffolds are likely interactive, as our live imaging suggests that Cnn emerges from the PCM zone and extends into the flare zone in an MT-dependent manner (Fig. 10, 2). Additionally, a conserved motif at the N terminus of Cnn, CM1, is required for the extension of Cnn flares (Zhang and Megraw, 2007). We speculate that CM1 may link Cnn to MTs and drive scaffold expansion in embryos and/or centrosome dematuration in other cell types.

Although related, the two scaffolds are likely functionally unique. The Spd2–Cnn scaffold is the main anchor of γTub within the PCM zone, whereas the PLP–Cnn scaffold functions to anchor Cnn within the flare zone, effectively preventing interphase centrosome dematuration by resisting MT-dependent particle release forces (Fig. 10, 4 and 5). Critically, unlike the Spd2–Cnn scaffold, the PLP–Cnn scaffold is exclusive to interphase centrosomes. We conclude that each architectural subdomain that comprises the centrosome structure is subject to multiple layers of regulation, some being zone specific. This stratification of regulatory organization defines the function of each zone.

Our Y2H studies show that PLP and Cnn likely interact at two defined domains. Analysis of PLPΔ5 suggests that the Cnn(F1)–PLP(F5) interaction is required to properly organize the PCM zone and extends into the flare zone in an MT-dependent manner (Fig. 10, 2). Additionally, a conserved motif at the N terminus of Cnn, CM1, is required for the extension of Cnn flares (Zhang and Megraw, 2007). We speculate that CM1 may link Cnn to MTs and drive scaffold expansion in embryos and/or centrosome dematuration in other cell types.

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Cnn flares and PLP satellites, and is required to dampen Cnn particle release (Fig. 10, 5). However, this large truncation includes the PACT domain and may disrupt other PLP interactions. Nevertheless, we can conclude that the Cnn(F1)–PLP(F5) interaction is not essential for PLP–Cnn scaffold formation, because Cnn and PLP still colocalize at flares and cytoplasmic particles. The second interaction between Cnn-F3 and PLP-F2 requires the Cnn CM2 domain (Fig. 10, 6) and is disrupted by the \( \text{cnn}^{B4} \) mutation. However, this mutation also disrupts an interaction with Cen required for actin furrow assembly (Kao and Megraw, 2009). Formally, we cannot rule out the possibility that the \( \text{cnn}^{B4} \) mutation may disrupt Cnn function due to protein destabilization. In addition, because gross actin aberrations were not observed in \( \text{plp}^{-} \) mutants (unpublished data), it is likely that Cnn CM2 mediates several interactions involved in distinct pathways that are disrupted in the \( \text{cnn}^{B4} \) mutant background.

Nevertheless, the \( \text{cnn}^{B4} \) mutation does lead to a loss of PLP satellites, with no effect on the PLP centriole pool (Fig. 10, 3). This intriguing result supports a model where PLP satellites and Cnn flares are functionally interdependent within the interphase flare zone, analogous to the interdependence of Spd2 and Cnn within the PCM zone. Moreover, the \( \text{cnn}^{B4} \) mutation demonstrates a specific separation of PLP function at satellites versus centrioles. Our data show the disordered localization of Cnn and \( \gamma \)Tub, as well as NUF and embryonic lethality, is more severe in \( \text{cnn}^{B4} \) than in \( \text{plp}^{-} \) mutants. These data suggest that Cnn functions to organize interphase centrosome structure through multiple pathways, with only one requiring PLP activity.

Previous work on the regulation of Cnn by Polo and Spd2, combined with our functional analysis of direct interactions with PLP, paints a complex portrait of the Cnn molecule. We propose that the coregulation of PLP satellites and Cnn flares is required for their anchorage to the centrosome, preventing interphase centrosome inactivation, and maintaining robust MT asters throughout the cell cycle. Disruption of this interphase PLP–Cnn scaffold leads to increased cytoplasmic PCM particles that act as platforms for MTs, leading to the formation of an aberrant interphase MT array in the embryo. These effects on MTs then cause defects in centrosome separation, nuclear spacing, and spindle assembly, culminating in chromosome missegregation, DNA damage, and embryonic lethality. Together, our data emphasize the fundamental importance of regulating interphase centrosome form and function for viability.

### Materials and methods

#### Fly stocks

The following mutant strains and transgenic lines were used: the \( \text{plp}^{P56} \) allele is a P-element insertion that interrupts the PLP sequence (Spradling et al., 1999) and was recombined onto the FRT\(^{10} \) chromo-
some to generate plp$^{2172}$; FRT$^{141}$ chromosomes were used for the generation of germ line clones (plp$^-$ embryos) by the PLP/ovoD method (Chou and Perrimon, 1996); the tacc$^1$ allele is a strong hypomorph (Gergely et al., 2000); and Df(3R)Exel$^{6142}$ (Bloomington Stock Center), nsp$^{RNA}$i (Vienna Drosophila Resource Center transformant 21982), and maternal-Tubulin GAL4 expressed GAL4 under the control of the maternal Tubulin promoter (Bloomington Stock Center); cnn$^{B4}$ is a hypomorphic allele caused by an R1141H single amino acid mutation (Vaizel-Oha-yon and Schejter, 1999), Ubi-GFP-γ-Tub23C expresses GFP-γ-Tub23C under the Ubiquitin promoter, and PBac-GFP-SAS6 is a recombineered GFP-SAS6 construct (Lerit and Rusan, 2013); UASp-Bld10-GFP expresses Cep135/Bld10-GFP under UAS (upstream activating sequence) regulator elements (Mottier-Pavie and Megraw, 2009); Ubi-GFP-PACT is a fusion of GFP with the PACT domain of PLP expressed under the Ubiquitin promoter (Martinez-Campos et al., 2004); Ubi-GFP-SAK expresses GFP-SAK/Plk4 under the Ubiquitin promoter (Basto et al., 2008); GFP-Polo expresses GFP-Polo with a Polo promoter (Moutinho-Santos et al., 2008); Ubi-D-TACC-RFP expresses TACC-RFP under the Ubiquitin promoter (Conduit et al., 2010); Ubi-GFP-SAS4 expresses GFP-SAS4 under the Ubiquitin promoter (Dix and Raff, 2007); H2AvD-mRFP expresses H2A-RFP under endogenous regulatory elements (Pandey et al., 2005); Ubi-GFP-α-Tubulin expresses GFP-α-Tubulin under the Ubiquitin promoter (Rebollo et al., 2004); and mCherry-Cnn (a gift from P. Singh and C. Cabernard, University of Basel, Basel, Switzerland), unpublished reagent generated by tagging Cnn with mCherry at the endogenous locus as described in Singh et al., 2014. PLP$^{Δ5}$-GFP was expressed in plp$^{2172}$; F(3L)Brd15 mutants, and all other transgenes (GFP-γ-Tub, H2A-RFP, Cnn-mCherry, and PLP$^{Δ1}$, GFP) were expressed in plp$^-$ germline clones unless otherwise noted.

**Construction of transgenic animals**

All transgenic flies were generated by BestGene, Inc., using standard P-element transformation. For PLP$^{Δ5}$-GFP, a C'-terminal truncation (deleting amino acids 2,539–2,895, which include the PACT domain) of PLPFL was PCR amplified using the primers 5′-CACCATGCCCCATTAATGCTTTATTTACG-3′ and 5′-TTCATTGAAGTGTTCCAACTCTGTTTCGGC-3′, directionally cloned into the pENTR-D vector (Invitrogen), and recombined into the P-element destination vector pUWG (Ubiquitin promoter, C'-terminal GFP; Drosophila Genomics Resource Center) via the Gateway cloning system (Invitrogen). P$_{hsp}$-GFP-Cnn was generated from the bacteriophage artificial chromosome clone CH322-10I24 (BACPAC Resources Center, Children's Hospital Oakland Research Institute) using the following primers to amplify the N-terminal GFP-Kanamycin cassette from the N-EGFP (N-terminal EGFP) template vector (Venken et al., 2008): 5′-TCAAGTGTTAGAATTATTGTGTGCGAAAGTTAACTATTTTGAGGACCCCTCCATGTTGACCAAGGCGAGGAG-3′ and 5′-GTACCATTTGGCGCCGCAATATCCGCCGCAGAAACTGTTAGACTGGTGACTACTACCCCGCTGAGGGACGACGACCAGGC-3′. This modified bacteriophage artificial chromosome was directionally integrated into the genome using the PhiC31 system at site VK33 (Chromosome 3), PLP$^{Δ5}$-GFP was generated by PCR amplification of PLP-F1 flanked by 5′-pEnt and 3′-P-FL homologous arms using the primers 5′-GCAAGGCTCCGGCGCGGCGGCCCTTCCACCAAGGAGATGAACTCTGACTATATACCGATATGACAGC-3′ and 5′-GCAATTCCCGGACATCGCCGCGATCTGCTCTC-3′.
A second fragment consisting of PLP-F3 through PLP-F5 flanked by 5'-PLP-F1 and 3'-pEntr homology arms was PCR amplified using the primers 5'-GAAGAGAGCAGATCCGCTGATCTCCGACCAAGACCGATGCCGAAATGC-3' and 5'-AGCTTGGGTTCGCCGGCCTCACCCTTTGGCTATGATAGTCCGCCGGCATGCCCTC-3'. The two fragments were enzymatically combined by a one-step isothermal DNA assembly (Gibson et al., 2009) into the pENTR-D vector, and then recombined into the P-element destination vector pUWG (Ubiquitin promoter, C'-terminal GFP; Drosophila Genomics Resource Center) via the Gateway cloning system. More than 32 independent PLPΔ25-GFP transgenic lines were screened for fluorescence using confocal microscopy. None of these lines showed GFP localization in cycling embryos.

Assessment of zygotic lethality and hatch rate analysis

Because males zygotically homozygous for the plp3172 mutation are uncoordinated and sterile (Martinez-Campos et al., 2004), we crossed males heterozygous for plp3172 (over the TM6B,Tb balancer chromosome) to females with germline homozygous mutant for plp, thus 50% of the analyzed embryos received a paternal WT plp gene and were paternally rescued. Eggs were collected at 25°C and the percentage of hatched larvae was divided by the total number of fertilized eggs. Embryonic lethality for plpΔ was 16% (n = 64/404 embryos) and is significantly increased compared with WT (2.4%; n = 458 embryos; P < 0.0001 by Fisher's exact test). Because 50% of the embryos were paternally rescued, lethality for plpΔ maternal/zygotic mutant embryos may be estimated to be 32% (64/202). The remaining 68% (138/202) maternal/zygotic mutant embryos hatched, but die as first instar larvae, as we do not detect any second instar larvae that are non-Tubby (Tb).

We conclude that all of the detected viable animals had received a WT copy of plp from the balancer chromosome. PLPΔ flies showed significantly greater embryonic lethality (54.9%; n = 89/162 embryos) than either controls or plp−/− (P < 0.0001). Because 50% of embryos were paternally rescued, this suggests that PLPΔ maternal/zygotic mutants are 100% embryonic lethal. The fact that embryonic lethality of PLPΔ exceeded plpΔ suggests it may exert dominant-negative effects.

Microscopy

Embryos were imaged using 40x, 1.3 NA or 100x, 1.49 NA oil immersion objective lenses on a microscope (Eclipse Ti; Nikon) fitted with a stage incubator (20/20 Technology, Inc.) heated to 25°C, a CSU-22 spinning-disc confocal head (Yokogawa Electric Corporation), a cooled CCD camera (Clara; Andor Technology), and the Perfect Focus System (Nikon) all run by an automated controller (MAC6000; Ludl Electronic Products) using MetaMorph software (Molecular Devices). Laser excitation was supplied by a laser merge module equipped with 491 nm, 561 nm, and 642 nm solid-state lasers (VisiTech International). For super-resolution SIM, immuno-labeled samples were mounted in Vectashield (Vector Laboratories, Inc.) and imaged with the DeltaVision OMX4 SIM Imaging System (Applied Precision).

Embryos (1–2 h old) were prepared for live imaging as described previously (Lerit and Gavis, 2011) with the following modifications: series 700 halocarbon oil (Sigma-Aldrich) was used to cover embryos adhered to a stick¥1.5 coverslip layered with glue extracted from double-sided Scotch tape (3M). The embryos and coverslip were inverted and 50-mm gas-permeable lumox dish (Sarstedt) fitted with two halves of a broken 1 coverslip used as spacers. Images were collected at 0.25-µm intervals over a 5–10 µm volume at 30-s time intervals or imaged at 500-ms intervals in a single optical plane as noted. Images were assembled using ImageJ (National Institutes of Health), Photoshop (Adobe), and QuickTime Player Pro 7 (Apple) software to crop regions of interest, adjust brightness and contrast, separate or merge channels, and generate maximum-intensity projections.

Image analysis

Total fluorescence intensity projections of the entire centrosome/cen- triole volume were used to measure the peak (P) and outer edge (OE) position of Sas6, Asl, and PLP, and the OE position of γTub and Cnn. Single optical sections through the middle of the centrosome were used for P measurements of γTub and Cnn. All line scans were single-pixel-wide line scans of 5 µm generated by the Plot Profile tool in ImageJ. Line scans were converted to a −2.50 to 2.50 scale such that the 0.0 µm position marked the center of the centriole. Average background fluorescence was subtracted from each pixel along the line scan. GraphPad Software was then used to fit each line scan to a Gaussian curve from which the mean (μ) and SD (Gaussian RMS) was calculated. μ represents the position of the Gaussian peak (P), which is the peak fluorescence intensity. The full width at half maximum (FWHM) was calculated for each curve using 2.35482×SD. The protein’s OE was calculated as μ + (0.5xFWHM). The mean P and OE for each measured centrosome protein is reported in Fig. S1 A. For Sas6, the entire −2.5 to 2.5 line scan was used to fit the Gaussian curve, for all other proteins two independent curves were generated along the −2.5 to 0.0 and 0.0 to 2.5 positions. For the γTub and Cnn within the PCM zone, a Gaussian Blur tool in ImageJ was used to smooth the edges of the PCM such that the fitted Gaussian produced an R2 >0.90 (Fig. S1 B). Comparisons of individual native and blurred line scans indicate a negligible impact of the blurring function on position measurements. For OE measurements of PLP satellites and Cnn flares, a half-Gaussian curve was fitted to the distal edge of the flare. A mirror image of this curve was added to generate the full-Gaussian from which μ and SD were calculated (Fig. S1 B, OE flare).

Cytoplasmic Cnn flares (not continuous with the centrosome or satellites) were manually counted from a 20 × 20-µm (400 µm2) box from three random areas in at least five different embryos for each genotype and cell cycle stage. Flare density is reported per 100 µm2. To analyze PLP satellite dynamics, kymographs along a single pixel line scan were generated using ImageJ software, and satellites that remained visible for at least five consecutive frames were manually tracked using the nTrack plugin (Meijering et al., 2012). Mean velocities were determined for individual satellites by dividing the time of transit by the distance between the first and last monitored positions. NUF was quantified as described previously (Poulton et al., 2013); in brief, embryos were stained for actin to label cortical boundaries and DAPI to label DNA. Empty (devoid of nucleus) actin cages were counted as sites of NUF. γH2Av labels double-stranded DNA breaks and was observed in all nuclei undergoing NUF. Data were plotted and statistical analysis was performed using Excel (Microsoft) and Prism software (GraphPad Software). To calculate significance, the normality of the distributions was confirmed with the D’Agostino and Pearson normality test. Data were then subjected to Student’s two-tailed t test or the Wilcoxon rank-sum test and are displayed as mean ± SD unless otherwise noted.

Immunofluorescence

Embryos were fixed in a 1:4 solution of 4% paraformaldehyde/heptane for 20 min before devitellinization in methanol. Samples were rehydrated stepwise into PBST (PBS with 0.1% Tween-20) then blocked in BBT (PBST with 0.1% BSA) for 2 h before an overnight incubation with nutation at 4°C in BBT with primary antibody. Samples were washed, further blocked in BBT supplemented with 2% normal goat serum (NGS), then incubated for 2 h at room temperature in secondary antibody. For visualization of γTub, embryos were blocked in PBST with 1% BSA. For visualization of MTs, embryos were prepared as described previously (Lerit and Gavis, 2011) according to the method of Theurkauf (1994). In brief, embryos were fixed for 3 min in 1:1 37% paraformaldehyde/heptane, rinsed three times in fresh PBS, and
hand devitellinized. After staining, samples were washed in PBST and mounted in Aqua-PolyMount (Polysciences, Inc.) and imaged on a spinning disk confocal microscope (Eclipse Ti; Nikon).

Primary antibodies used were as follows: rabbit and guinea pig anti-PLP (made against amino acids 8–351; 1:3,000; Brownlee et al., 2011), guinea pig anti-Asl (made against full-length Asl; 1:3,000; gift from G. Rogers, University of Arizona Cancer Center, Tucson, AZ), rabbit anti-Cnn (made against amino acids 271–1,034; 1:2,000; gift from T. Megraw, Florida State University, Tallahassee, FL); anti-phospho-Histone H3 Ser10 (pH3; 1:1,000; EMD Millipore), mouse anti-γ Tub ascsites GTU-88 (1:50; Sigma-Aldrich), mouse anti-α-Tubulin DM1a (1:500; Sigma-Aldrich), rabbit anti-γH2A made against QPD-QRKNVILSQAY (1:500; gift from K. McKim, Rutgers University, Piscataway, NJ), and rabbit anti-αSpd-2 (1:2,000; Rodrigues-Martins et al., 2007). Secondary antibodies and counterstains were as follows: Alexa Fluor 488, 568, or 627 (1:500; Molecular Probes); DAPI (4′,6-diamidino-2-phenylindole; 10 ng/ml); and Alexa Fluor 488 phalloidin (1:500; Life Technologies).

Y2H analysis
Y2H was performed as described in Galletta et al. (2014). In brief, fragments of PLP and Cnn were introduced into pDEST-pGADT7 and pDEST-pGBK7 using Gateway technology (Invitrogen), transformed into Y187 or Y2HGold yeast strains (Takara Bio Inc.), and grown in –leu or –trp media to select for plasmids. After mating of the two strains, yeast were grown on –ade, –leu, –trp, –ura (QDO) media to select for plasmids. All plasmids were tested for the ability to drive reporter activity in the presence of an empty vector (autoregulation), are shown. All plasmids were tested for the ability to drive reporter activity in the presence of an empty vector (autoregulation). Plasmids that conferred autoactivity were omitted from further analysis (see Fig. S5 A).

Online supplemental material
Fig. S1 shows analysis of centrosome protein distribution. Fig. S2 shows the dynamics of PLP satellites. Fig. S3 shows disruption of γ Tub in plp−/ embryos. Fig. S4 shows that PLP-F5 is required for compaction of the mitotic centrosome. Fig. S5 shows interaction refinement of PLP-F2. Video 1 shows Cnn-GFP and H2A-RFP through multiple NCs in a control embryo. Video 2 shows Cnn-GFP during the mitotic centrosome. Video 1 shows Cnn-GFP and H2A-RFP through multiple NCs in a control embryo. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201503117/DC1.

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