Cdk1 Phosphorylates *Drosophila* Sas-4 to Recruit Polo to Daughter Centrioles and Convert Them to Centrosomes

**Highlights**

- Cdk1 phosphorylates Sas-4 to initiate Polo/Plk1 recruitment to daughter centrioles
- Polo recruitment promotes Asterless (Asl) incorporation into daughter centrioles
- Asl incorporation licenses new centrioles to duplicate and organize centrosomes
- These observations help explain why centriole conversion is tied to mitosis

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**In Brief**

Newly formed centrioles are converted to duplication-competent centrosomes during their first mitosis. Novak et al. show that in *Drosophila* this centriole conversion is induced by the Cdk1-dependent phosphorylation of Sas-4. Phosphorylated Sas-4 recruits Polo/Plk1 to new centrioles and promotes Asl incorporation, allowing these centrioles to duplicate and assemble a centrosome.
Cdk1 Phosphorylates Drosophila Sas-4 to Recruit Polo to Daughter Centrioles and Convert Them to Centrosomes

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SUMMARY

Centrioles and cilia are organized by a centriole pair comprising an older mother and a younger daughter. Centriole numbers are tightly regulated, and daughter centrioles (which assemble in S phase) cannot themselves duplicate or organize centrosomes until they have passed through mitosis. It is unclear how this mitotic “centriole conversion” is regulated, but it requires Plk1/Polo kinase. Here we show that in flies, Cdk1 phosphorylates the conserved centriole protein Sas-4 during mitosis. This creates a Polo-docking site that helps recruit Polo to daughter centrioles and is required for the subsequent recruitment of Asterless (Asl), a protein essential for centriole duplication and mitotic centrosome assembly. Point mutations in Sas-4 that prevent Cdk1 phosphorylation or Polo docking do not block centriole disengagement during mitosis, but block efficient centriole conversion and lead to embryonic lethality. These observations can explain why daughter centrioles have to pass through mitosis before they can duplicate and organize a centrosome.

INTRODUCTION

Centrioles organize centrosomes and cilia. These organelles have many important functions in cells, and their dysfunction has been linked to several human diseases (Bettencourt-Dias et al., 2011; Conduit et al., 2015a; Nigg and Raff, 2009). Much attention has focused on the mechanisms that regulate centriole (and so centrosome and cilium) numbers, as numerical abnormalities can be highly deleterious to cells. Centriole loss gradually triggers a p53-dependent response that leads to cell-cycle arrest and/or cell death in mammalian cells (Izquierdo et al., 2014; Lambrus et al., 2015; Wong et al., 2015), while centriole amplification has been linked to cancer (Basto et al., 2008; Ganem et al., 2009; Godinho et al., 2014; Gönçzy, 2015) and more recently to microcephaly (Arquint and Nigg, 2014; Martiensens et al., 2013). How centriole amplification influences these pathologies remains unclear (Serçin et al., 2016; Vitre et al., 2015).

Most cells are born with two centrioles that are precisely duplicated in S phase when a new “daughter” centriole is assembled at right angles to each “mother” in a tightly apposed engaged configuration. This configuration appears to be crucial for regulating centriole numbers, as mother centrioles normally cannot duplicate again until they disengage from their daughters during the subsequent mitosis, a process that requires the mitotic kinase Polo/Plk1 (Loncarek et al., 2010; Shukla et al., 2015; Tsou and Stearns, 2006a; Tsou et al., 2009). Thus, mother centrioles normally only duplicate once in S phase, and disengagement functions as a “license” that is acquired during mitosis to enable mother centrioles to duplicate again during the next S phase (Fırat-Karalar and Stearns, 2014; Nigg, 2007; Tsou and Stearns, 2006b).

Another important mechanism that helps regulate centriole numbers is that only mother centrioles are competent to duplicate during S phase (Cunha-Ferreira et al., 2009; Kleylein-Sohn et al., 2007; Loncarek et al., 2009). The restriction that prevents newly assembled daughter centrioles from duplicating during S phase does not simply rely on them being engaged to their mothers: under certain experimental conditions, such as overexpression of Sak/Plk4 (the key protein kinase that initiates centriole duplication), mother centrioles can duplicate again and form multiple daughters, but daughters cannot form daughters of their own. Importantly, it was recently shown that newborn daughter centrioles cannot form centrosomes or duplicate until they have passed through mitosis and are modified by Polo/Plk1 (Wang et al., 2011); this process has been termed “centriole-to-centrosome” conversion (Fu et al., 2016; Izquierdo et al., 2014; Wang et al., 2011), although we sometimes use the more generic “mitotic centriole conversion” here (see Discussion).

Great progress has been made in understanding the molecular mechanisms of centriole assembly (Conduit et al., 2015a; Gönçzy, 2012; Jana et al., 2014), and recent studies have shed some light on the process of mitotic centriole conversion. In human cells, Cep295 is required for this process: in the absence of Cep295, new daughter centrioles disengage from their mothers during mitosis but they cannot recruit any pericentriolar material (PCM) and these centrioles are destabilized (Izquierdo et al., 2014). In flies, the conserved centriole protein Asterless (Asl) is essential for both centriole duplication and for mitotic PCM recruitment (Blachon et al., 2008; Bonaccorsi et al., 1998; Dobbelare et al., 2008; Varma et al., 2007). Asl helps recruit Sak/Plk4 to the mother centriole to initiate centriole duplication (Dzhindzhiev et al., 2010; Novak et al., 2014), and to recruit Spd-2 and Cnn to the mother centriole to initiate mitotic centrosome assembly (Conduit et al., 2014a, 2014b). It was recently
shown that Asl is only recruited to daughter centrioles during mitosis, at about the time that they are converted to centrioles that can recruit PCM and duplicate (Fu et al., 2016; Novak et al., 2014). Thus, the recruitment of Asl to newly disengaged centrioles during mitosis appears to be a crucial step in centriole conversion in flies. Interestingly, a recent study showed that the fly homolog of Cep295, Ana1, also plays an important part in centriole conversion in flies (Fu et al., 2016). Ana1 is recruited to centrioles by Cep135/Bld10 in interphase, and is subsequently required to recruit Asl to the newborn centrioles during mitosis; a similar mechanism appears to operate in human cells (Fu et al., 2016). These previous studies, however, provide no insight into the crucial question of how Asl recruitment is regulated during the cell cycle so that it only occurs during mitosis.

We previously showed that another conserved centriole protein, Sas-4, has an important role in initially recruiting Asl to newborn centrioles during mitosis in flies (Novak et al., 2014), and a direct interaction between Sas-4 and Asl (human CPAP and Cep152, respectively) has been seen in both flies and humans (Cizmecioglu et al., 2010; Dzhindzhev et al., 2010; Hatch et al., 2010). Unlike Asl, Sas-4 is recruited to daughter centrioles during their assembly in S phase (Conduit et al., 2015b; Novak et al., 2014). If Sas-4 helps recruit Asl to newborn centrioles, this function must be strictly regulated so that it only occurs during mitosis. In this study we set out to identify the mechanism that might regulate the ability of Sas-4 to promote the recruitment of Asl to daughter centrioles during mitosis. We find that Sas-4 is phosphorylated during mitosis on Thr200 by the master mitotic regulator Cdk1 to create a Polo-docking site that appears to be required to recruit Polo, and subsequently Asl, to new centrioles. Remarkably, single point mutations that perturb Sas-4-Thr200 phosphorylation or the creation of the Polo-docking site strongly perturb Polo and Asl recruitment to the new centriole during mitosis. Although these newborn centrioles can separate from their mothers during mitosis, they are not converted to mother centrioles that can duplicate or recruit PCM, and embryos expressing these mutant Sas-4 proteins die very early in development.

**RESULTS**

**Sas-4 Thr200 Is Required to Recruit Asl to New Centrioles**

To investigate whether the ability of Sas-4 to recruit Asl to centrioles might be regulated by cell-cycle-dependent phosphorylation we searched for regions within, or adjacent to, the Asl-interacting region of Sas-4 (amino acids [aa] 101–650) (Dzhindzhev et al., 2010) that contained consensus phosphorylation sites for either Cyclin-dependent kinases (Cdks) or Polo/Plk kinases and that were conserved in *Drosophila* species. We identified four regions for further analysis: I (aa 198–244), II (aa 257–304), III (aa 378–406), and IV (aa 654–686) (Figure S1A).

We synthesized mRNA in vitro encoding GFP fusions of either wild-type (WT) Sas-4 (Sas-4-GFP) or mutant versions in which these regions were individually deleted. We injected these mRNAs into fly embryos expressing Asl-mCherry; the Sas-4-GFP proteins are gradually synthesized from the injected mRNA and incorporated into centrioles in competition with endogenous, unlabeled, Sas-4. We then assayed the effect of each fusion protein on Asl-mCherry incorporation (see Figure 1A for a schematic summary of this assay). In all cases, the Sas-4-GFP signal appeared as a tight, symmetrical focus at newly separated centriole pairs, demonstrating that none of the deletions blocked Sas-4 targeting to centrioles (Figures 1B and S1B). Asl normally localizes asymetrically to newly separated centriole pairs because it only starts to be incorporated into daughter centrioles during mitosis, so new mother centrioles contain very little Asl (Novak et al., 2014), and Asl-mCherry behaved in this way in all embryos expressing Sas-4-GFP, Sas-4-[ΔI]-GFP; Sas-4-[ΔII]-GFP; Sas-4-[ΔIII]-GFP, or Sas-4-[ΔIV]-GFP (Figures 1B and S1B). In contrast, the recruitment of Asl-mCherry to new centrosomes at the end of mitosis was greatly reduced in embryos expressing Sas-4-[ΔI]-GFP (note that Asl-mCherry localization in all these experiments was scored blindly) (Figures 1B and S1B).

Although Sas-4-[ΔI]-GFP expression severely disrupted the recruitment of Asl-mCherry to newly disengaged daughter centrioles, it did not detectably disrupt the localization of Asl-mCherry at newly disengaged mother centrioles (Figures 1B and S1B). This is expected, as fractions of both Sas-4 and Asl are stably incorporated into centrioles (Conduit et al., 2015b; Novak et al., 2014) and Asl-mCherry was initially recruited to these older centrioles during earlier rounds of mitosis: as Sas-4-[ΔI]-GFP is being gradually translated in these experiments, the ratio of Sas-4-[ΔI]-GFP/endogenous-Sas-4 at centrioles gradually increases during successive rounds of centriole duplication, presumably until it reaches a critical level that is sufficient to perturb Asl-mCherry recruitment to the new daughter centrioles (see Figure 1A).

Region I of Sas-4 is required to promote Asl recruitment to newly disengaged daughter centrioles and contains several Ser/Thr residues that are highly conserved in *Drosophila* species (Figure S2A). We tested whether mutating any of these residues to Ala would lead to a defect in Asl-mCherry recruitment (Figure S2). The individual expression of several forms of Sas-4-GFP containing combinations of multiple Ser/Thr-to-Ala substitutions within region I, including a form in which nine Ser/Thr residues were mutated to Ala (Sas-4-9A), did not detectably perturb Asl-mCherry recruitment (Figures 1B, S2B, and S2C; and data not shown). In contrast, expressing a form of Sas-4-GFP in which the single residue Thr200 was mutated to Ala (Sas-4-T200A-GFP) gave a strong defect in Asl-mCherry recruitment (Figures 1B and S2C). We conclude that Sas-4-Thr200 is required to promote Asl recruitment.

**Sas-4 Thr200 Can Be Phosphorylated by Cdk1 to Create a Polo-Docking Site In Vitro**

In flies, Sas-4-Thr200 is followed by a conserved proline residue (Pro221; Figure 2A), suggesting that it might be a substrate for Cdks (Endicott et al., 1999); it is also preceded by a conserved serine residue (Ser199; Figure 2A), suggesting that Thr200, when phosphorylated, could act as a docking site for Polo kinase (Plk1 in humans), which can interact with phosphorylated SpS/pT motifs through its conserved Polo-box domain (PBD) (Lowery et al., 2005). A peptide containing the fly Sas-4-Thr200-SPS motif was efficiently phosphorylated by recombinant human Cdk1/Cyclin B in vitro, and the phosphorylation was dramatically reduced if either Thr200 or Pro221 (but not Ser199) was mutated (Figure 2B). The peptide also functioned as an efficient docking
site for a recombinant human GST-PBD fusion protein in vitro, but only when Thr200 was phosphorylated; binding was also dramatically reduced when Ser199 was mutated (Figure 2C).

We conclude that the Sas-4-Thr200-STP motif can be phosphorylated by Cdk1/Cyclin B to create a Polo-docking site in vitro.

Interestingly, although the Asl/Cep152-interacting region of Sas-4/CPAP is generally not well conserved between flies and vertebrates, vertebrate CPAP proteins also contain a highly conserved STP motif in this region—surrounding human CPAP Thr616—and this motif can also be phosphorylated by Cdk1/Cyclin B to create a Plk1-docking site in vitro (Figures S3A–S3C).

The Sas-4-Thr200-SP Motif Is Required to Recruit Asl to Centrioles
To test whether the recruitment of Asl to new centrioles required Cdk1 phosphorylation (dependent on Pro201, in vitro) and/or the

Figure 1. Sas-4-Thr200 Is Required to Localize Asl to New Centrioles
(A) Schematic and example images illustrate the RNA injection assay used for screening the role of various mutant Sas-4-GFP proteins (green) in the recruitment of Asl-mCherry (red) to new centrioles. Arrows in micrographs highlight a positive hit, where Asl-mCherry has not detectably incorporated into the new centriole that has just separated from its older mother. Scale bars, 2 μm.

(B) Asl-mCherry (red) localization at newly separated centrioles in living embryos expressing various Sas-4-GFP fusions from injected mRNA (green, as indicated). Note that Asl-mCherry localizes normally at new centrioles in embryos expressing WT Sas-4-GFP or Sas-4-9A-GFP, in which nine conserved Ser/Thr residues in the vicinity of Thr200 were substituted to Ala (see Figure S2B). The localization of Asl-mCherry is disrupted in embryos expressing Sas-4-[Δ]GFP or embryos expressing full-length Sas-4-GFP carrying either T200A, P201G, or S199G point mutations. Arrows indicate new centrioles that have not incorporated Asl-mCherry. Scale bar, 5 μm.

See also Figures S1 and S2.
creation of the Polo-docking site (dependent on Ser199, in vitro), we expressed several forms of Sas-4-GFP in which either Pro201 or Ser199 were mutated. Both Sas-4-P201G-GFP and Sas-4-S199G-GFP produced a strong defect in Asl-mCherry recruitment (Figure 1B), supporting the hypothesis that Asl recruitment is dependent on both Cdk1 phosphorylation and the creation of a Polo-docking site. Moreover, the conservative substitution of Ser199 to Thr (Sas-4-S199T) also strongly disrupted Asl-mCherry recruitment (Figure S2C), suggesting that the crucial function of Ser199 is not to be phosphorylated, but rather to recruit the PBD. Finally, we also tested the effect of substituting Thr200 for a “phospho-mimicking” Glu. The negative charge of Glu can mimic the negative charge of the phosphate group, potentially allowing Glu to engage in electrostatic interactions in a similar manner to phospho-Thr. Importantly, however, the interaction between a phospho-peptide and the PBD is only partially mediated by electrostatic interactions, as it also relies on specific hydrogen-bonding interactions between the phosphate group and the PBD (Elia et al., 2003), so PBD binding to phospho-Thr should not be mimicked by Glu. The expression of Sas-4-T200E-GFP strongly disrupted Asl-mCherry recruitment (Figure S2C), demonstrating that Glu cannot mimic the function of phospho-Thr in Asl recruitment. Taken together, these studies support the hypothesis that the Sas-4-Thr200-STP motif functions as a Cdk1-dependent Polo-docking site that is required to recruit Asl to newborn centrioles during mitosis in vivo.

Sas-4-Thr200 Appears to be Phosphorylated during Mitosis In Vivo
To test whether Thr200 is phosphorylated in vivo, we generated antibodies against a phospho-Thr200-containing peptide and
purified antibodies that recognized the phosphorylated, but not the non-phosphorylated, peptide (Figure S3D). In fixed embryos these antibodies did not detectably stain centrosomes (not shown), perhaps because phosphorylated Thr200 rapidly binds Polo, which then masks the phosphorylated epitope. In an attempt to overcome this potential problem, we fluorescently labeled the antibodies and injected them into living embryos that expressed the microtubule (MT) marker Jupiter-mCherry to allow the accurate staging of cell-cycle progression (Figure 2D).

We previously showed that fluorescently labeled antibodies raised against non-phosphorylated Sas-4 bind to centrosomes throughout the cell cycle when injected into embryos (Novak et al., 2014). In contrast, the anti-Sas-4-pThr200 antibodies did not detectably bind to centrosomes in early S phase (Figure 2D, t = −300 s) but started to accumulate there in late S phase, reaching maximal levels (Figure 2D, t = −40 s) just before nuclear envelope breakdown (Figure 2D, t = 0 s). Antibody levels at centrosomes remained high until metaphase (Figure 2D, t = 100 s) and then fell during anaphase (Figure 2D, t = 260 s), and the antibody was essentially undetectable at centrosomes by telophase (Figure 2D, t = 280 s). Importantly, the antibodies started to bind to centrosomes again toward the end of the next S phase (Figure 2D, t = 1,420 s), demonstrating that they were not simply degraded or inactivated in the embryo. Thus, Sas-4 Thr200 appears to be phosphorylated in vivo from late S phase to late mitosis. This strongly suggests that Sas-4-Thr200 phosphorylation is catalyzed by the M-phase-specific Cdk1, rather than by the S-phase-specific Cdk2.

The Sas-4-Thr200-1STP Motif Is Required to Recruit Polo to New Centrioles but Is dispensable for Centriole Disengagement

When we injected the anti-Sas-4-pThr200 antibodies at higher levels, we noticed that the antibody perturbed the recruitment of Asl-GFP to new centrosomes at the end of mitosis (Figure 3A).

This strongly argues (although it does not conclusively prove) that the antibody is specifically binding to the phospho-Sas-4-Thr200 epitope in embryos, rather than non-specifically binding to a different centrosomal protein, as the antibody can elicit the same very specific phenotype as mutating Thr200. Moreover, we noticed that high concentrations of antibody also appeared to block the recruitment of Polo-GFP to the newly converted mother centrioles without interfering with Polo-GFP localization at the older mother centriole (Figure 3B). Thus, remarkably, interfering with a single putative Polo-docking domain on Sas-4 appears to be sufficient to block the recruitment of Polo to new centrosomes in embryos.

To test this possibility further, we assayed the effect of separately expressing three mutant forms of the Sas-4-Thr200-1STP motif (S199G, T200A, and P201G) on Polo recruitment to new centrosomes. Like the phospho-specific antibody, each of these mutant forms of Sas-4 strongly perturbed the recruitment of Polo-GFP to new centrosomes (Figure 3C). Interestingly, however, neither the anti-Sas-4-pThr200 antibodies nor the Sas-4-Thr200-1STP-motif mutants perturbed centriole disengagement at the end of mitosis (Figures 1B, S3E, and S3F). This suggests that centriole disengagement does not require Polo recruitment to the daughter centriole.

Sas-4-Thr200 Is Required for Mitotic Centriole Conversion In Vivo

Polo and Asl both play an important part in mitotic PCM recruitment in flies (Conduit et al., 2014a, 2014b; Dobbelare et al., 2008; Sunkel and Glover, 1988; Varmark et al., 2007). We predicted, therefore, that although the expression of Sas-4-T200A does not block centriole disengagement, it should block the subsequent recruitment of PCM to the disengaged daughter centriole that cannot recruit Polo or Asl. In embryos expressing WT Sas-4-GFP, new centrosomes formed centrosomes after they separated from their mothers (as judged by their ability to organize MTs), which invariably formed spindle poles during the next mitosis (Figures 4A and 4C). In contrast, in embryos expressing Sas-4-T200A-GFP many of the new centrosomes failed to organize MTs after they separated from their mothers, and these subsequently failed to form spindle poles (Figures 4B and 4D). As a result, many of the spindles in these embryos shared spindle poles organized by the original mother centrioles, while the “unconverted” centrioles floated freely in the cytoplasm (arrows in Figure 4D). Thus, the Sas-4-T200A point mutation appears to block mitotic centriole conversion and subsequent centrosome assembly in embryos.

The Sas-4-Thr200-1STP Motif Is Essential for Early Embryonic Development

In these RNA injection experiments we had to express the Sas-4-GFP fusion protein in the presence of the endogenous (but unlabeled) Sas-4 protein, as mutant flies lacking Sas-4 are uncoordinated (unc) due to the lack of cilia and so cannot mate or lay embryos (Basto et al., 2006) (see schematic, Figure S4A). To test whether the Sas-4-Thr200-1STP motif was essential for Sas-4 function in vivo, we generated stable transgenic lines expressing mCherry fusions to either WT or 1STP-motif mutant forms of Sas-4. These fusion proteins were all expressed at similar levels (Figures S4C and S4D), but whereas WT Sas-4-mCherry efficiently rescued the unc defect of Sas-4 mutant flies, Sas-4-T200A-mCherry rescued this phenotype very poorly, indicating that the rescued flies still had significant cilia defects (Figure S4B).

Embryos laid by Sas-4 mutant females rescued by WT Sas-4-mCherry developed normally, but embryos laid by mutant females rescued by any of the three mutated forms of Sas-4-mCherry (S199G, T200A, or P201G) failed to hatch as larvae (>1,000 scored for each genotype; data not shown). A detailed analysis of mutant embryos expressing Sas-4-T200A-mCherry revealed that they all arrested very early in development with only a few nuclei (Figure 5A). The spindles in these embryos were often disorganized, and of 18 optically sectioned spindle poles only seven had detectable Asl and Cnn staining (data not shown). We also expressed Polo-GFP in these embryos to allow us to follow Polo behavior. Of the 24 spindle poles we optically sectioned in these embryos, only five had detectable Asl or Polo staining (Figure 5B). Centrosomes are essential for early embryonic development in flies, and embryos lacking functional centrosomes arrest with a very similar phenotype to that we observe here (Stevens et al., 2007; Varmark et al., 2007). Thus, these observations support the conclusion from our embryo mRNA injection studies that the Sas-4-Thr200-1STP motif is essential for proper centriole duplication and centrosome assembly in embryos.
Figure 3. The Sas-4-Thr200-STP Motif Helps Recruit Both Polo and Asl to New Centrioles

(A and B) Asl-GFP (A, green) and Polo-GFP (B, green) localization in embryos injected with high levels of anti-Sas-4-pThr200 antibodies (red). Embryos are shown during mitosis and the following S phase; two regions from the same embryo are shown in which there is either a low concentration (unbound, left panels) or high concentration (bound, right panels) of the antibodies. Colored arrowheads indicate the same centrosomes in mitosis and then after centrosome separation; arrows highlight examples where binding of the antibody has perturbed the recruitment of Asl-GFP (2/2 embryos) or Polo-GFP (2/3 embryos) to new centrioles. Scale bars, 5 μm.

(C) Polo-GFP (green) localization at newly separated centrioles in embryos expressing WT Sas-4-mKate2 (mKate2 was used as a red fluorescent tag due to its relatively short fluorescence maturation time compared to other red fluorescent proteins; see Supplemental Experimental Procedures) or Sas-4-mKate2-Thr200-SP-motif mutant forms (red; as indicated). Note that Polo-GFP localizes symmetrically at newly separated centriole pairs in embryos expressing WT Sas-4-mKate (arrowheads), however, Polo-GFP localization is strongly disrupted or absent from new centrioles in embryos expressing T200A, P201G, or S199G mutant forms of Sas-4-mKate2. Arrows highlight examples where the mutant fusion protein has perturbed the recruitment of Polo-GFP to new centrioles, while arrowheads indicate the unperturbed older centrioles in these embryos. Scale bars, 5 μm (left panel) and 2 μm (right panel).

See also Figure S3.
The Sas-4-Thr200-STOP Motif Is Required for Efficient Polo and Asl Recruitment and Centriole Duplication and Centrosome Assembly in Somatic Brain Cells

The nuclear division cycles of early Drosophila embryos are extremely short and thus necessitate very rapid and efficient mitotic centriole conversion. We wanted to test whether the Sas-4-STOP motif was important for centriole conversion during the much longer cell cycles in cell types other than embryos. We therefore examined the recruitment of Polo-GFP to centrosomes in living Sas-4 mutant mitotic larval brain cells expressing either WT Sas-4-mCherry or Sas-4-T200A-mCherry. Almost all the mitotic cells expressing Polo-GFP and WT Sas-4-mCherry had two centrosomes that were strongly decorated with both fusion proteins (Figure 5C). In contrast, ~40% of cells expressing Polo-GFP and Sas-4-T200A-mCherry had no detectable centrosomes, ~35% had one centrosome, and only ~25% had two centrosomes (as judged by the presence of Sas-4 foci). In the cells that had at least one centrosome, ~30% of these Sas-4 foci contained no detectable Polo-GFP (suggesting these were not functional centrosomes but rather unconverted centrioles), while the rest had detectable Polo-GFP, although it was often only weakly localized (Figure 5C).

We obtained similar results when we analyzed the distribution of Asl in fixed Sas-4 mutant brains expressing WT or Sas-4-Thr200-STOP-motif mutant mCherry-fusion proteins (Figure 5D). All the brains rescued by the Sas-4-STOP mutant proteins exhibited reduced centrosome numbers (as judged by the number of Asl foci), and, in those cells that contained centrosomes, Asl recruitment was often very weak (Figures 5D, S4E, and S4F). We note that somatic brain cells can occasionally form acentrosomal MT-organizing centers that contain Asl, but not Sas-4 (Baumbach et al., 2015), potentially explaining why the reduction in centrosome numbers in the mutants appears to be greater when we score centrosomes by counting Sas-4 foci (Figure 5C) than when we count Asl foci (Figure 5D). We conclude that Polo and Asl recruitment to centrosomes, and centriole duplication and centrosome assembly, are strongly perturbed in Sas-4 mutant brain cells that are rescued by Sas-4-Thr200-STOP-motif mutants.

DISCUSSION

It has long been known that newly formed daughter centrioles are unable to duplicate during the S phase in which they were born, even under conditions that allow mother centrioles to proceed through multiple rounds of duplication (Cunha-Ferreira et al., 2009; Kleylein-Sohn et al., 2007; Loncarek et al., 2008). How exactly daughter centriole duplication in S phase is prevented is unclear, but it has recently been shown that
Figure 5. The Sas-4-Thr200-STP Motif Is Essential for Sas-4 Function In Vivo

(A) Quantification of nuclei number in 0- to 2-hr-old Sas-4 mutant embryos expressing endogenous levels of either Sas-4-mCherry (n = 349 embryos) or Sas-4-T200A-mCherry (n = 272 embryos).

(B) Micrographs of typical embryos stained to reveal the distribution of MTs (red), nuclei (cyan), Polo-GFP (yellow), and the centrosomal marker Asl (magenta). Note that both Polo-GFP and Asl are enriched at the spindle poles in embryos expressing Sas-4-mCherry, while their localization is severely disrupted in embryos expressing Sas-4-T200A-mCherry, resulting in the formation of mitotic spindles that usually lack detectable centrosomes. The arrowhead in embryo I highlights a pole with a centrosome that contains some Polo and low levels of Asl; the spindles in embryo II do not detectably have centrosomes at their poles. Scale bars, 10 μm.

(C) Micrographs show examples of living mitotic larval neuroblast cells co-expressing Polo-GFP with Sas-4-mCherry or Sas-4-T200A-mCherry. Graphs below show the percentage of mitotic cells that contain the indicated number of Sas-4 foci that are either Polo-GFP positive (darker green shading; categories I and III) or Polo-GFP negative (lighter green shading; categories II and IV); the percentage of cells without any detectable centrosomes is indicated by blue/green shading (category V). The percentages in each category are: Sas-4-mCherry: I, 96.43%; V, 3.57%; Sas-4-T200A-mCherry: I, 17.5%; II, 7.5%; III, 25%; IV, 10%; V, 40%. Scale bar, 5 μm.

(D) Graph shows the quantification of Asl foci (typical examples of each class are illustrated in the micrographs) in mitotic (phospho-histone H3 [PHH3] positive) third instar larval brain cells of the indicated genotypes. Error bars indicate SEM. Scale bar, 5 μm.

See also Figure S4.
Stearns, 2003), while daughter centrioles cannot duplicate at (Loncarek et al., 2008; Tsou and Stearns, 2006a; Wong and during S phase because they are engaged to their daughters et al., 2014). Thus, mother centrioles do not normally reduplicate while daughter centrioles have passed through their first mitosis, whose acquisition during mitosis is required to allow new centri- oles stably (Novak et al., 2014), meaning that once 2014). Importantly, a fraction (Novak et al., 2014) of Asl is incorporated into mitotic PCM (Conduit et al., 2014b; Novak et al., 2014). Polo and Asl recruitment to the new centriole in vivo, leading to severe defects in centriole duplication and centrosome assembly. Moreover, antibodies that specifically recognize the phosphorylated Sas-4-Thr200-STP motif in vitro localize to centro- somes during mitosis and can also specifically block the recruitment of Polo and Asl to newly disengaged daughter centrioles. The simplest interpretation of this collective data is that the Polo-docking site on Sas-4-Thr200 fail to convert into functional cen- trosoles during successive cell generations. (C) New centrioles that are not phosphorylated on Sas-4-Thr200 fail to convert into functional cen- trosoles, which can result in spindle abnormal- ities during the following cell cycle (cycle “n+1”), and potentially complete centrosome depletion. Polo/Pik1 modifies the daughter centriole during mitosis in some way that allows it to subsequently duplicate and form a centro- some (Wang et al., 2011). This process has previously been termed centriole-to-centrosome conversion (Fu et al., 2016; Iz- quierdo et al., 2014; Wang et al., 2011), although we prefer the term “mitotic centriole conversion” because, although in fly embryos the converted centrioles almost immediately form centro- somes and duplicate, in other cell types the converted centrioles may only duplicate or form a centrosome much later in the next cell cycle (when the cells eventually enter S phase or M phase, respectively). Thus, mitotic centriole conversion generates a centriole that is competent to both duplicate and form a centro- some, although these events may occur independently and sometime after the initial conversion event, depending on the cell type. In flies, the recruitment of Asl to the new centriole during mitosis appears to be a critical event in centriole conversion, as Asl incorporation ultimately allows centrioles to both duplicate and recruit mitotic PCM (Conduit et al., 2014b; Novak et al., 2014). Importantly, a fraction (~50%) of Asl is incorporated into centrioles stably (Novak et al., 2014), meaning that once daughter centrioles have passed through their first mitosis, they no longer have to pass through mitosis again to acquire Asl. In this way, Asl incorporation acts as a “primary license” whose acquisition during mitosis is required to allow new cen- trioles to duplicate and form a centrosome for the first time (Novak et al., 2014). Thus, mother centrioles do not normally reduplicate during S phase because they are engaged to their daughters (Loncarek et al., 2008; Tsou and Stearns, 2006a; Wong and Steams, 2003), while daughter centrioles cannot duplicate at all because they lack Asl and so cannot recruit Sak/Pik4 (Dzhindzhiev et al., 2010; Novak et al., 2014). In this study, we have identified a Polo/Pik1-dependent mechan- ism that is required to recruit Asl to newly formed centrioles during mitosis, and we have shown that this mechanism appears to be initiated by the recruitment of Polo to the daughter centriole by the Cdk1-dependent phosphorylation of Sas-4 (Figure 6). Sas-4 is a conserved centriole protein that is essential for centriole duplication in flies (Basto et al., 2006), and we previ- ously showed that it is involved in recruiting Asl to newly formed centrioles during mitosis (Conduit et al., 2014b; Novak et al., 2014). Cdk1 can phosphorylate the Sas-4-Thr200-STOP motif to create a Polo-docking site in vitro, and an analysis of several STOP-motif mutants reveals that mutations that disrupt Cdk1 phosphorylation or PBD binding in vitro invariably perturb both Polo and Asl recruitment to the new centriole in vivo, leading to severe defects in centriole duplication and centrosome assem- bly. Moreover, antibodies that specifically recognize the phosphorylated Sas-4-Thr200-STOP motif in vitro localize to centro- somes during mitosis and can also specifically block the recruitment of Polo and Asl to newly disengaged daughter centrioles. The simplest interpretation of this collective data is that the Polo-docking site on Sas-4 directly recruits Polo to the new centriole, although we cannot exclude the possibility that this Sas-4-Thr200-STOP motif may not bind Polo in vivo and actually performs some other unknown function that indirectly is required for Polo recruitment. Nevertheless, our findings now directly implicate Cdk1 kinase in promoting the centriole duplication cycle through the direct phosphorylation of a centriolar protein, and

Figure 6. Model
(A) Schematic model illustrates how Sas-4 (light gray) is incorporated into new centrioles during S phase and is phosphorylated on Thr200 (brown) by Cdk1 as cells enter mitosis, thus allowing the recruitment of Polo (magenta) and then Asl (blue). The recruitment of Asl allows the new centriole to recruit PCM (which also contains Polo and so is indicated by the magenta cloud around the fully converted centriole). The mother centriole and its associated PCM are indicated semi-transparently. (B) Schematics summarize the importance of the Cdk1- and Polo-regulated conversion of new centrioles to duplication-competent MT- organizing centers (MTOC) during mitosis. (B and C) Schematics summarize the importance of the Cdk1- and Polo-regulated conversion of new centrioles to duplication-competent MT- organizing centers (MTOC) during mitosis. (B and C) Schematics summarize the importance of the Cdk1- and Polo-regulated conversion of new centrioles to duplication-competent MT- organizing centers (MTOC) during mitosis. (C) New centrioles that are not phosphorylated on Sas-4-Thr200 fail to convert into functional cen- trosoles, which can result in spindle abnormal- ities during the following cell cycle (cycle “n+1”), and potentially complete centrosome depletion. (A) Schematic model illustrates how Sas-4 (light gray) is incorporated into new centrioles during S phase and is phosphorylated on Thr200 (brown) by Cdk1 as cells enter mitosis, thus allowing the recruitment of Polo (magenta) and then Asl (blue). The recruitment of Asl allows the new centriole to recruit PCM (which also contains Polo and so is indicated by the magenta cloud around the fully converted centriole). The mother centriole and its associated PCM are indicated semi-transparently. (B and C) Schematics summarize the importance of the Cdk1- and Polo-regulated conversion of new centrioles to duplication-competent MT- organizing centers (MTOC) during mitosis. (C) New centrioles that are not phosphorylated on Sas-4-Thr200 fail to convert into functional cen- trosoles, which can result in spindle abnormal- ities during the following cell cycle (cycle “n+1”), and potentially complete centrosome depletion. Polo/Pik1 modifies the daughter centriole during mitosis in some way that allows it to subsequently duplicate and form a centro- some (Wang et al., 2011). This process has previously been termed centriole-to-centrosome conversion (Fu et al., 2016; Iz- quierdo et al., 2014; Wang et al., 2011), although we prefer the term “mitotic centriole conversion” because, although in fly embryos the converted centrioles almost immediately form centro- somes and duplicate, in other cell types the converted centrioles may only duplicate or form a centrosome much later in the next cell cycle (when the cells eventually enter S phase or M phase, respectively). Thus, mitotic centriole conversion generates a centriole that is competent to both duplicate and form a centro- some, although these events may occur independently and sometime after the initial conversion event, depending on the cell type.
provide an intriguing molecular insight into the mechanism that likely regulates the primary recruitment of the crucial cell-cycle regulator Polo/Plk1 kinase to newly formed centrioles in Drosophila.

In fly embryos the Sas-4-Thr200-STA motif appears to be essential for centriole conversion. In somatic fly cells this motif appears to be important, but it is not essential; STP-motif mutations perturb Asl and Polo recruitment to centrosomes, and centrosome numbers are dramatically reduced, but centrosome duplication and centrosome assembly are not abolished. These data suggest that the Sas-4-Thr200-STA motif plays a critical role in promoting efficient centriole conversion rather than being an indispensable component of this process. We speculate that when the enrichment of Polo at daughter centrioles is perturbed by Sas-4-Thr200-STA mutations then other, suboptimal, Polo-docking sites within the daughter centriole, or the Polo pool present in the cytoplasm or in the PCM surrounding the neighboring mother centriole, can still initiate the molecular cascade leading to the conversion of the daughter centriole, although with far lower efficiency. Such a reduction in efficiency appears sufficient to block conversion during the rapid divisions in the syncytiotum embryo (where daughters must be converted in just a few minutes), but is not sufficient to completely block this process in brain cells, which have much longer cell-cycle times and thereby a greater time window in which to complete centriole conversion.

Surprisingly, disrupting the recruitment of Polo to daughter centrioles did not appear to alter the efficiency of centriole disengagement, even though Polo/Plk1 is of critical importance in this process (Loncarek et al., 2010; Shukla et al., 2015; Tsou et al., 2009). This demonstrates that the recruitment of Polo to daughter centrioles, at least in flies, does not play a key role in timing centriole disengagement; instead the pool of Polo on the mother centriole (or in the cytoplasm) is sufficient to drive this event.

The recruitment of Polo/Plk1 to daughter centrioles during mitosis is likely a key regulatory event in the centrosome cycle. It is perhaps surprising, therefore, that this recruitment relies so heavily on a single phosphorylation site on a single protein. We suspect that the phosphorylation of Sas-4-Thr200 is only required for the initial binding of Polo to new centrioles: once recruited, Polo probably phosphorylates several other nearby proteins to create further Polo-docking sites, thereby driving robust Polo recruitment that ultimately influences many aspects of centriole and centrosome function. How Polo docking to Sas-4 might enable Asl recruitment to daughter centrioles is unclear, but an attractive possibility is that Polo subsequently phosphorylates either Asl or other sites on Sas-4 to increase the efficiency of the previously characterized and conserved direct interaction between Sas-4 and Asl (Cizmecioglu et al., 2010; Zhidnizhev et al., 2010; Hatch et al., 2010). An alternative, and not mutually exclusive, possibility is that the centriole proteins Cep135 and Ana1 are phosphorylated by Polo, as both proteins also help to recruit Asl to centrioles (Fu et al., 2016), although there is some evidence that Ana1 may primarily help to generally maintain Asl at all mother centrioles, rather than specifically help to initially recruit Asl to new mother centrioles (Saurya et al., 2016). Future studies will doubtless identify the crucial substrates of Polo that allow Asl to be recruited to new centrioles during mitosis.

**Experimental Procedures**

**Transgenic Drosophila Lines**

The Asl-mCherry, Sas-4-mCherry (Conduit et al., 2014b), Sas-4-GFP (Novak et al., 2014), Asl-GFP (Blachon et al., 2008), Polo-GFP (Buszczak et al., 2006) and Jupiter-mCherry (Callan et al., 2010) lines used in this study have been described previously. In all experiments, Asl-mCherry or Asl-GFP was expressed at near-endogenous levels in the aslGFP (Baumbach et al., 2015) homozygous mutant background. Sas-4-GFP (when supplied transgenically, not in mRNA injection experiments) was expressed in the Sas-4GFP (Basto et al., 2006) homozygous mutant background. To generate Sas-4–T200A-mCherry, Sas-4–P201G-mCherry, and Sas-4–S199G-mCherry lines, we introduced the respective point mutations into the Sas-4–mCherry P element transformation vector (Conduit et al., 2014b) using a QuikChange II XL Site-Directed Mutagenesis Kit (Agilent Technologies). The transgenic lines were generated by the Fly Facility in the Department of Genetics, University of Cambridge. Sas-4–mCherry, Sas-4–T200A-mCherry, Sas-4–P201G-mCherry, or Sas-4–S199G-mCherry were analyzed in the Sas4GFP/Df(3R)SBSC221 genetic background. OregonR was used as the WT control.

**RNA Synthesis and Microinjection**

In vitro RNA synthesis was performed using a T3 miniMESSAGE mMACHINE kit (Ambion) and RNA was purified using an RNeasy MiniElute kit (Qiagen). All RNA constructs were injected at a concentration of 2 mg/ml into 0–30-min-old embryos. Sas-4-GFP constructs were injected into either Asl-mCherry, aslGFP or Jupiter-mCherry embryos (both lines are homozygous for the WT Sas-4 allele), and Sas-4–mKate2 constructs were injected into polo-gfp (Trap)/TM4 embryos (also homozygous for the WT Sas-4 allele). Microinjected embryos were incubated at 22°C and imaged after 60–120 min (when GFP fusions were injected) or after 120–150 min after injection (when mKate2 fusions were injected; as the fluorescence maturation time of mKate2 appears significantly longer than that of GFP in flies), always within the syncytial blastoderm stage of development. Live imaging was performed using the spinning-disc confocal system described below.

**Live Imaging**

Live syncytial blastoderm stage embryos were imaged on a PerkinElmer ERS spinning-disc confocal system on a Zeiss Axiovert microscope, using a 63 x 1.4NA oil-immersion objective. Thirteen confocal sections were collected from each embryo (0.5-μm steps) every 20 s.

Living third instar larval brains were also imaged on the spinning-disc confocal system specified above. Twenty-one confocal sections (0.5-μm steps) were collected of mitotic neuroblasts every 50 s.

**Quantitative Analysis of Living Embryos and Brains**

Asl-mCherry and Polo-GFP localization in embryos injected with WT or mutant Sas-4 RNA constructs was scored blind (movies were renamed and mixed post acquisition, and the entire dataset generated for this study was scored blindly at the same time). Embryos in which multiple new centrosomes lacked any detectable levels of Asl-mCherry were classified as showing defective Asl-mCherry recruitment, while in the case of Polo any embryos in which multiple centrosome pairs displayed strongly asymmetric Polo-GFP levels were counted as showing defective Polo-GFP recruitment. Embryos were scored qualitatively this way, and fluorescence levels were not quantified due to the variable nature of the RNA injection process. In embryos injected with anti-Sas-4-α-TbPol200, Asl-GFP or Polo-GFP localization was compared within each embryo on the side closest to the injection site (centrosomes bound by antibody) and the side furthest away (unbound centrosomes). Timing of centrosome separation was also assessed this way in Sas-4–GFP embryos injected with anti-Sas-4–α-TbPol200.

Quantification of Sas-4–mCherry and Polo-GFP foci in living larval brains was also performed blind. Movies of mitotic larval neuroblasts (identified by clustered, Polo-GFP-labeled kinetochores) were renamed and mixed post acquisition, and the entire dataset was scored blindly at the same time.

**Fixed Analysis of Embryos and Larval Brains**

0- to 2-hr-old embryos were fixed and stained as described previously (Stevens et al., 2009), Guinea-pig anti-Asl (Rloque et al., 2012) and mouse

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anti-α-tubulin (Sigma) primary antibodies were used at 1:500 dilution to immunostain embryos. GFP-Booster (atto 488, ChromoTek), Alexa Fluor antimouse 568, and anti-guinea-pig 647 (Life Technologies) were used at 1:500 dilution as secondary antibodies. Hoechst 33258 (Life Technologies) was used at 1:5,000 dilution to stain DNA. Images of fixed embryos presented in this study were collected using an Olympus confocal microscope (FV1200 IX83; Olympus) with Fluoview software, using a 60 × 1.3NA silicon oil-immersion Super Apochromat lens (UPLSAPO 60XS). Nineteen confocal sections were collected (0.2-μm steps). Nuclei numbers were quantified in fixed embryos using a Zeiss Axioskop 2 microscope with a 10 × 0.3NA dry objective.

Third instar larval brains were dissected, fixed, squashed, and stained as described previously (Stevens et al., 2009). The following primary antibodies were used at 1:500 dilution to immunostain brains: rabbit anti-phospho-His-tone3 (Cell Signalling), rat anti-Asl (Franz et al., 2013), guinea-pig anti-Cnn (Lucas and Raff, 2007), Alexa Fluor anti-rat 488, anti-rabbit 568 and anti-guinea-pig 647 (Life Technologies) were used at 1:500 dilution as secondary antibodies. DNA was stained with Hoechst 33258 (1:5,000 dilution). All samples were blinded following slide preparation, prior to imaging and quantification. The samples were imaged and centrosome numbers were counted on a Zeiss Axioskop 2 microscope with Metamorph software, using a 63 × 1.4NA oil-immersion objective. Eleven confocal sections were collected (0.2-μm steps). Samples were unblinded after the quantification was completed for the full dataset. Percentage values were calculated for each brain (each data-point represents a brain). Statistical comparisons of centrosome frequencies (shown in Figure S3F) were performed using the Mann-Whitney test (for pairwise comparisons) or the Kruskal-Wallis test (for simultaneous comparison of three or four genotypes).

**In Vitro Cdk1 Kinase Assay**

Synthetic peptides (50 μM final concentration) were incubated with 40 units of recombinant human Cdk1-cyclin B (New England Biolabs) in 1 x NEBuffer for Protein Kinases (New England Biolabs) with 100 μM cold ATP and 5 μCi γ-[32P]ATP in a reaction volume of 20 μl. Reactions were incubated at 30 °C for 30 min, then terminated by the addition of 10 μl 7.5 M guanidine-hydrochloride. 1.2 μl of each reaction was spotted onto an avidin-coated membrane (SAM2 biotin oil-immersion objective. Eleven confocal sections were collected (0.2-μm steps). Nuclei numbers were quantified in fixed embryos using a Zeiss Axioskop 2 microscope with a 10 × 0.3NA dry objective.

**In Vitro GST-Plk1 PBD Binding Assay**

Streptavidin beads (30 μl/reaction; Dynabeads MyOne Streptavidin T1, Life Technologies) were washed three times with Binding Buffer (50 mM Tris [pH 8.0], 150 mM NaCl, 1 mM EDTA, 1 mM DTT, 1 μM protease inhibitor cocktail [Roche], 1 μM phosphatase inhibitor cocktail [Sigma]) and incubated with a 10-fold molar excess of biotinylated peptide in 1 ml of Binding Buffer for 30 min at room temperature. For analysis of the Drosophila Sas-4-Thr200-STOP motif, the synthetic phosphopeptides and their non-phosphorylated counterparts were coupled directly to the beads. For analysis of the human CPAP-Thr616-STOP motif, the synthetic non-phosphorylated peptides were pre-incubated in 1 x NEBuffer for Protein Kinases (New England Biolabs) and 200 μM ATP (New England Biolabs) with or without 120 units of recombinant human Cdk1-cyclin B (New England Biolabs) in a final volume of 60 μl (3 nmol peptide/reaction) for 30 min at 30 °C before coupling to the streptavidin beads as described above (the entire reaction volume was added to the beads together with 1 ml of Binding Buffer). This approach was taken because synthetic production of the phosphorylated version of the human CPAP-Thr616-STOP peptide could not be completed by the manufacturer within the required time frame. The beads were rinsed four times by resuspension in 200 μl of Capture Buffer (50 mM Tris [pH 8.0], 150 mM NaCl, 1 mM EDTA, 1 mM DTT, 1 μM protease inhibitor cocktail [Roche], 1 μM phosphatase inhibitor cocktail [Sigma], 0.01% Tween 20, 0.01% BSA), then incubated with 1 ml of PBD solution (0.1 μM recombinant human GST-Plk1-PBD [Sigma; #SRP0360, GST-Plk1a aa 367–603] in 50 mM Tris [pH 8.0], 150 mM NaCl, 1 mM EDTA, 2 mM DTT, 1 μM protease inhibitor cocktail [Roche], 1 μM phosphatase inhibitor cocktail [Sigma], 0.01% Tween 20, 0.01% BSA) for 3 hr at 4 °C. The beads were rapidly washed four times in 250 μl of Wash Buffer (50 mM Tris [pH 8.0], 150 mM NaCl, 1 mM EDTA, 2 mM DTT, 1 μM protease inhibitor cocktail [Roche], 1 μM phosphatase inhibitor cocktail [Sigma], 0.1% Tween 20) then resuspended and boiled in 30 μl of SDS loading dye. 2 × 1.5 μl of each sample was spotted onto nitrocellulose membranes (Bio-Rad). The membranes were air-dried at room temperature and incubated in milk solution (PBS + 4% milk + 0.1% Tween 20) for 1 hr. One of the two membranes was probed using rabbit anti-GST antibody (Abcam; 1:500 dilution) and anti-rabbit-terhesardis peroxidase (HRP) (GE Healthcare; 1:3,000 dilution) for GST-Plk1-PBD detection, and the other membrane was probed with streptavidin-HRP (Thermo Scientific; 1:3,000 dilution), as the peptide loading control. Three technical repeats were performed.

**Rescue Experiments**

Sas-4-mCherry, Sas-4-T200A-mCherry, Sas-4-P201G-mCherry, and Sas-4-S199G-mCherry transgenes were crossed into the Sas4Δ2214/Df3R/BSC221 genetic background for all rescue experiments. All flies analyzed (embryonic, larval, or adult phenotypic assessments) were homozygous for the transgene they carried. For assessment of the importance of Sas-4-Thr200 in Polo recruitment, polo-gfp(Trap) was crossed into each of the Sas-4-mCherry, Sas4Δ2214/Df3R/BSC221 and Sas-4-T200A-mCherry, Sas4Δ2214/Df3R/BSC221 genetic backgrounds (Polo-GFP was only expressed in the rescue experiments where its presence is indicated in the respective figure panel).

Embryonic phenotypes were assessed in the progeny of rescued females that were mated with OregonR males. Several hundred such embryos were collected from mutant females rescued by each Sas-4-mCherry, Sas-4-T200A-mCherry, Sas-4-P201G-mCherry, or Sas-4-S199G-mCherry. None of the embryos laid by mothers rescued by any of the three STP mutant constructs hatched as larvae and 100% of these embryos arrested before gastrulation, as judged by their homogeneous white color even following >7 days of incubation at 25°C. Detailed immunofluorescence analysis was then performed on the embryos laid by Sas-4-T200A-mCherry, Sas4Δ2214/Df3R/BSC221 mothers (both with or without co-expression of Polo-GFP) to confirm that eggs were fertilized and to analyze the nature of the early embryonic arrest.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures and four figures and can be found with this article online at http://dx.doi.org/10.1016/j.devcel.2016.05.022.

**AUTHOR CONTRIBUTIONS**

Z.A.N. designed and performed the experiments and wrote the manuscript. A.W. performed some of the immunofluorescence imaging. L.G. optimized the use of the mKate2 fluorescent tag in the RNA injection screen system. J.W.R. designed experiments and wrote the manuscript.

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REFERENCES


Supplemental Information

Cdk1 Phosphorylates *Drosophila* Sas-4 to Recruit Polo to Daughter Centrioles and Convert Them to Centrosomes

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Inventory of Supplemental Materials:

- **Figure S1**, related to Figure 1. Identifying mutations in Sas-4 that interfere with the loading of Asl onto new centrioles.
- **Figure S2**, related to Figure 1. Sas-4-Thr200 is required to load Asl onto new centrioles.
- **Figure S3**, related to Figures 2 & 3. The Sas-4-Thr200-STM motif is required to load Polo onto new centrioles.
- **Figure S4**, related to Figure 5. Analysis of Sas-4-Thr200-STM motif mutant transgenic flies.
- **Supplemental Experimental Procedures**
- **Supplemental References**
Supplemental Figures

Figure S1, related to Figure 1.

**Identifying mutations in Sas-4 that interfere with the loading of Asl onto new centrioles.**

**A**  
*Dr. Drosphila melanogaster Sas-4:*  
- Amino acid sequence of *Drosophila melanogaster* Sas-4, indicating the Asl-interacting region (grey) (Dzhindzhev et al., 2010) and the four regions of interest we identified that contain multiple Ser/Thr residues that are highly conserved across fly species (I-IV, highlighted in different colors).  
- Micrographs illustrate Asl-mCherry (red) localization at newly separated centrioles in living embryos expressing full length Sas-4-GFP or the GFP-fusion of one of the four deletion constructs (green), as indicated; arrows highlight new centrioles that have separated from their mothers, but have not incorporated Asl-mCherry. The deletion of regions II, III or IV does not detectably interfere with Sas-4 localisation or Asl incorporation; deletion I does not detectably interfere with Sas-4 localisation, but dramatically perturbs Asl incorporation at new centrioles.

**B**  
Embryos displaying defective Asl-mCherry recruitment.
Figure S2, related to Figure 1.

Sas-4 Thr200 is required to load Asl onto new centrioles. A) Amino acid sequence alignment of Sas-4 region I from several Drosophila species (alignment was performed using the full protein sequence from each species); Thr200 is highlighted by an asterisk. B) The amino acid sequence of Sas-4 region I from Drosophila melanogaster highlighting three highly conserved putative Cdk phosphorylation sites—Thr200 (pink/maroon; asterisk), Ser217 and Ser244 (blue)—and 9 other conserved Ser/Thr residues (yellow). C) Micrographs show Asl-mCherry (red) localization at newly separated centrioles in living embryos expressing either full length Sas-4-GFP or the indicated mutated forms of Sas-4-GFP (green) in which either Thr200 (pink label), Ser217 and Ser244 (blue label), or all 9 of the other conserved Thr/Ser residues highlighted in (B) (yellow label), have been mutated to Ala. Note that, of these three constructs, only the T200A mutation appears to perturb the loading of Asl onto new centrioles (arrows). Asl-mCherry localization appeared normal in embryos injected with Sas-4-S199T-GFP or Sas-4-9A-GFP, but was disrupted in embryos injected with Sas-4-T200A-GFP. Micrographs also show that each the Sas-4-GFP S199T (grey label) and the T200E mutation (maroon label) disrupt the recruitment of Asl-mCherry to new centrioles similarly to the T200A mutation (arrows).
The Sas-4-Thr200-TP-motif is required to load Polo onto new centrioles. A) A schematic representation of human CPAP and *Drosophila* Sas-4 highlighting the conserved TCP domain (green) that interacts with STIL/Ana2 (Cottee et al., 2013; Hatzopoulos et al., 2013) and the region shown to interact directly with Cep152/Asl (purple) (Cizmecioglu et al., 2010; Dzhindzhev et al., 2010; Hatch et al., 2010). This latter region is not generally well conserved, and the dotted lines delineate regions where an amino acid sequence alignment between a number of vertebrate and *Drosophila* species is shown below. These regions contain a short stretch of reasonable homology between the fly and vertebrate proteins (red box); the conserved Sas-4-Thr200-TP-motif (asterisk) in flies is located just upstream of this region, while the vertebrate sequences contain a conserved TP motif (asterisk) located slightly downstream from this region. B) *In vitro* assay of Cdk1/Cyclin B-dependent phosphorylation of the human CPAP-Thr616-TP-motif. The dot blot shows the loading of the biotinylated peptides (left panel); the autoradiogram shows the incorporation of $^{32}$P (right panel). C) *In vitro* assay of the human CPAP-Thr616-TP-motif binding to recombinant GST-Polo-Box protein. The peptides were pre-phosphorylated with Cdk1/Cyclin B prior to incubation with GST-Polo-Box as indicated (see Experimental Procedures). The dot blots show the loading of peptide (top panel), and the binding of GST-Polo-Box-domain (bottom panel). D) Western blots confirm the specificity of the anti-Sas-4-pThr200 antibody: this antibody recognizes a peptide containing the Sas-4-Thr200 region only if Thr200 is phosphorylated (right panel). Streptavidin-HRP (StAv) was used to confirm the equal loading of the biotinylated peptides (left panel). E) Anti-Sas-4-pThr200 antibodies do not block centriole disengagement at the end of mitosis. Images show the timing of centrosome separation in a transgenic Sas-4-GFP (green) embryo that has been injected with anti-Sas-4-pThr200 (red). Two regions from the same embryo are shown, in which there is either a low concentration (unbound) or high concentration (bound) of antibody, and the embryo is shown during mitosis (left panels) and the following S-phase (right panels). Note that binding of the antibody to centrosomes during mitosis does not appear to alter the timing of centrosome separation, demonstrating that centriole disengagement has not been detectably perturbed. F) Images show the timing of centrosome separation in embryos expressing Jupiter-mCherry (red) and either wild type Sas-4-GFP or Sas-4-T200A-GFP (green), as indicated. Centrosome separation occurs with normal timing during telophase/early S-phase in embryos expressing Sas-4-T200A-GFP, confirming that the expression of this protein does not detectably perturb centriole disengagement, even though the loading of Polo-GFP onto new centrioles is strongly perturbed.
**Figure S4, related to Figure 5.**

**Analysis of Sas-4-Thr200-STOP-motif mutant transgenic flies.**

A) Schematics indicate the life cycle stages and the specific genotype-phenotype relationships in *Drosophila* that correspond to the different Sas-4-STOP transgenic rescue experiments presented in this study. Green, orange and red boxes indicate increasingly severe phenotypical defects (green: minimal defects – red: lethality).

B) Climbing assay of adult *Drosophila* measuring the proprioception of flies expressing transgenic Sas-4-mCherry or Sas-4-T200A-mCherry in a Sas-4 null genetic background. Sas-4 null flies are severely uncoordinated and completely unable to stand up therefore they were not subjected to the climbing assay. The graph shows that Sas-4-mCherry expression fully rescues this proprioception defect of Sas-4 null flies (as measurable with this type of assay) while Sas-4-T200A-mCherry expression can only marginally rescue the Sas-4 null phenotype. Error bars show SEM. Statistical analysis was performed using the Mann-Whitney test.

C) Western blot shows that Sas-4-mCherry and Sas-4-T200A-mCherry are expressed at near-endogenous levels in pre-cellularised embryos and that levels of Asl are not detectably perturbed. Actin is shown as a loading control. Note that the multiple bands of Sas-4 visible on the blot are not phosphorylation-dependent (bands appear resistant to phosphatase treatment; data not shown).

D) Western blot shows that in 3rd instar larval brains Sas-4-mCherry and the Sas-4-mCherry T200A, P210G and S199G mutant proteins are expressed at similar levels. The levels of Asl are also not detectably perturbed, although Asl levels are reproducibly slightly downregulated in Sas-4 mutant brains that are not rescued by any of these transgenes. Actin is shown as a loading control.

E) Images show examples of 3rd instar larval brain cells that were used for quantifying Asl foci in mitotic cells; genotypes are indicated. The centrosomal levels of Asl were often variable in flies expressing Sas-4-T200A-mCherry, Sas-4-P201G-mCherry or Sas-4-S199G-mCherry in the absence of wild type Sas-4 (an example of 2 cells from the same Sas-4-P201G-mCherry, Sas-4 brain is shown—both cells have two centrosomes, but one has normal Asl levels, the other has low Asl levels, although both cells were scored as having two centrosomes).

F) Scatter plot representation of graph shown in Figure 5D. Each data point represents an average value obtained from one brain (30-50 cells counted in each brain).
bars show SEM. Statistical analysis was performed using the Mann-Whitney test (pairwise comparisons; indicated with solid lines) or the Kruskal-Wallis test (comparison of 3 or 4 categories; indicated with dashed lines). Bold lines highlight significant differences between categories. Note that the variability in centrosome frequencies is far greater in flies expressing Sas-4-T200A-mCherry, Sas-4-P201G-mCherry or Sas-4-S199G-mCherry when compared to WT Sas-4-mCherry, suggesting that each of these mutations induces a stochastic failure in centriole conversion.

Supplemental Experimental Procedures

Generation of Sas-4 constructs for RNA microinjection
To generate the various Sas-4-GFP constructs used for RNA injection experiments the Sas-4 coding region was PCR amplified from w+ DNA and subcloned into the pRNA-EGFP-CT (Cottee et al., 2015) vector using Gateway technology. The single intron of Sas-4 was removed using QuickChange II XL Site-Directed Mutagenesis Kit and all described deletions or point mutations were subsequently introduced with QuickChange II XL or QuickChange Multi Site-Directed Mutagenesis Kits. To generate the Sas-4-mKate2 constructs the mKate2 coding sequence was PCR amplified from the pDONR P2R-P3-mKate2 vector (Buj et al., 2013) with flanking homology arms to the sequence surrounding the eGFP coding region within the pRNA-Sas-4-EGFP vector. An Nhel restriction site was introduced into the pRNA-Sas-4-EGFP vector immediately upstream of the GFP start codon and In-Fusion cloning (Clontech) was used to swap the eGFP coding sequence for the mKate2 sequence using the Nhel and SpeI restriction sites within pRNA-Sas-4-EGFP (the Nhel site was not included in the homology arms when amplifying mKate2 and therefore was removed during the In-Fusion reaction to restore the original linker sequence). The T200A, P201G and S199G point mutations were introduced into pRNA-Sas-4-mKate2 as described in Experimental Procedures.

Phospho-antibody generation
The anti-Sas-4-pThr200 antibody was generated by Pocono Rabbit Farm & Laboratory Inc. A polyclonal antibody was raised in rabbit against a synthetic peptide with the sequence of SS{pThr}PDLKSSYASSC. Following affinity purification using the phospho-peptide, the antibody was purified against the non-phosphorylated peptide sequence SSTPDLKSSYASSC to eliminate any antibodies that are not specific to the phospho-epitope.

Antibody labelling and microinjection
The anti-Sas-4-pThr200 antibody was covalently coupled to Alexa488 or Alexa568 (succinimidyl ester forms, Life Technologies) by 1-hour incubation of the antibody and dye in 50mM HEPES pH 8.6, 50mMKCl, 1mM MgCl, 1mM EGTA. The labelled antibody was separated from unbound dye and eluted in 50mM HEPES pH 7.5, 50mMKCl, 1mM MgCl, 1mM EGTA using size exclusion chromatography and concentrated to approximately 5-10mg/ml using Amicon Ultra filters (50kDa MWCO). The antibodies were injected into syncytial blastoderm stage embryos and imaged immediately following injection using the spinning disc confocal system described in Experimental Procedures.

Image processing and presentation
All fluorescent images presented that were collected on the Perkin Elmer Spinning Disc confocal system (living embryos and brains) were obtained by 3D rendering of image data using Volocity software. The Volocity ‘smooth zoom’ filter was applied to all images. All immunofluorescence images of fixed embryos and brains were processed using Fiji software (Schindelin et al., 2012).

Peptides
For the kinase assays and Polo-Box binding assays, peptides (and, where appropriate, their phosphorylated equivalents) were synthesised by GenScript. The full peptide sequences were the following: biotin-Ahx-G-Ahx-GGAKNACS-{S/G}-T/P/A-{P/G}-DLKSAAKK (representing the Drosophila Sas-4-Thr200-STP-moitif) and biotin-Ahx-G-Ahx-GGAGHRMSS-{T/A}-{P/G}-VKAVAKK (representing the human CPAP-Thr616-STP-motif). Ahx indicates aminohexanoic acid. Peptides were resuspended and stored in 0.1M phosphate buffer pH7.4, 150mM NaCl, 2mM DTT.
**Antibodies used for Western blotting**
Rabbit anti-Sas-4 (Basto et al., 2006), rabbit anti-Asl (Conduit et al., 2010), mouse anti-actin (Sigma) and rabbit anti-Sas-4-pThr200 (this study) primary antibodies were diluted 1:500 in milk solution for Western blot/dot blot experiments. HRP-conjugated anti-rabbit or anti-mouse (GE Healthcare; diluted 1:3,000 in milk solution) were used as secondary antibodies. Streptavidin-HRP (Thermo Scientific; diluted 1:3,000 in milk solution) was used for detecting biotin, as a loading control, for all in vitro experiments.

**Adult Drosophila climbing assay**
The climbing ability of wild type, Sas-4-mCherry,Sas4S2214/Df(3R)BSC221 and Sas-4-T200A-mCherry,Sas4S2214/Df(3R)BSC221 was analysed blind. Groups of five 1-day-old flies were place in a 15ml measuring cylinder and knocked to the bottom. After 6 seconds the distance climbed by each of the five flies was measured and averaged. Each group of flies was measured 3 times and a single data point was calculated as the average value of the three repeats. 8-11 groups of flies were scored for each genotype (total of 40-55 flies, 3 runs each), and groups were unblinded after quantification was completed. Statistical comparisons between the different genotypes were performed using the Mann-Whitney test.

**Supplemental References**


