KIF4A and PP2A–B56 form a spatially restricted feedback loop opposing Aurora B at the anaphase central spindle

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The mitotic kinase Aurora B is concentrated at the anaphase central spindle by the kinesin MKlp2 during mitotic exit and cytokinesis. This pool of Aurora B phosphorylates substrates including the kinesin KIF4A to regulate central spindle length. In this paper, we identify a counteracting system in which PP2A–B56γ and -ε, but not PP2A–B56α, -β, and -δ, are maintained at the central spindle by KIF4A. Biochemical assays show that PP2A–B56γ can dephosphorylate the T799 Aurora B site on KIF4A and thereby counteract the Aurora B- and microtubule-stimulated ATPase activity of KIF4A. In agreement with these observations, combined silencing of PP2A–B56γ and -ε resulted in increased phosphorylation of KIF4A T799 and decreased central spindle growth in anaphase B. Furthermore, reduced turnover of regulatory phosphorylation on another Aurora B substrate MKlp1 was observed, suggesting that PP2A–B56γ and -ε play a general role opposing Aurora B at the central spindle. KIF4A and PP2A–B56γ and -ε therefore create a spatially restricted negative feedback loop counteracting Aurora B in anaphase.

Introduction
Anaphase central spindle formation is controlled by a group of conserved microtubule-associated proteins and kinesin motors (Barr and Gruneberg, 2007; Glotzer, 2009; Brust-Mascher and Scholey, 2011; Duelberg et al., 2013). Key among these are the antiparallel microtubule-stabilizing factor PRC1 and a microtubule dynamic instability suppressing kinesin KIF4A, which work together to create microtubule overlaps of defined length (Bringmann et al., 2004; Bieling et al., 2010; Subramanian et al., 2010; Hu et al., 2011). These microtubule overlaps are further stabilized by the centralspindlin complex formed by the MKlp1/KIF23 kinesin-like motor protein and the Cyk-4/ MgcRacGAP Rho family GTPase regulator (Mishima et al., 2002; Douglas et al., 2010). Both PRC1 and centralspindlin are inhibited until anaphase onset by Cdk1–cyclin B phosphorylation (Jiang et al., 1998; Mollinari et al., 2002; Mishima et al., 2004; Zhu et al., 2006; Neef et al., 2007). At the onset of anaphase, PP2A–B55 removes the inhibitory Cdk1 phosphorylation on PRC1, triggering central spindle formation only after chromosome separation has commenced (Cundell et al., 2013). Together, these factors can explain the Cdk1-regulated decrease in microtubule catastrophe observed toward the cell equator in anaphase (Brust-Mascher et al., 2004; Cheerambathur et al., 2007). However, Cdk regulation of these components does not provide an obvious means to fine-tune microtubule dynamic instability in a spatially restricted manner and thereby promote anaphase B spindle elongation (Cheerambathur et al., 2007). In addition to negative regulation by Cdk1, all these components are subject to activating phosphorylations by either Aurora B or Plk1 (Polo-like kinase 1). PRC1 and centralspindlin phosphorylation by Plk1 are necessary for division plane specification (Neef et al., 2007; Petronczki et al., 2007; Wolfe et al., 2009). Aurora B phosphorylation of KIF4A at T799 activates its microtubule-stimulated ATPase activity and enhances KIF4A binding to PRC1, and the antiparallel microtubule overlap bundling activity of MKlp1 is promoted by Aurora B phosphorylation at S911 (Douglas et al., 2010; Nunes Bastos et al., 2013). KIF4A and MKlp1 are phosphorylated by a pool of Aurora B transported to the central spindle by the mitotic kinesin MKlp2 (Gruneberg et al., 2004; Neef et al., 2006; Ruchaud et al., 2007; Fuller et al., 2008; Brust-Mascher and Scholey, 2011; Nunes Bastos et al., 2013). Although, in both cases, previous studies
Figure 1. Identification of isoform-specific components of PP2A–B56 complexes. (A) HeLa cells treated with control and B56 regulatory subunit siRNA for 48–96 h were stained for DNA. Arrows indicate mitotically arrested, micronucleated, and binucleated cells. (B and C) Bar graphs show the frequency of binucleate (B) and micronucleated cells (C; means ± SEM, n = 2 measuring >200 cells per experiment). (D) Blots to confirm depletion of endogenous B56 regulatory subunits. KIF4A was used as a loading control and marker for mitosis. (E) Thymidine-synchronized HeLa cells were transfected with GFP-tagged PP2A–B56 regulatory subunits for 18 h, and then, the thymidine was washed out to allow the cells to enter mitosis. After 10 h, these cells were stained for DNA, mouse anti–Aurora B, and sheep anti-MKlp2. PP2A–B56 was visualized by GFP fluorescence. Arrows mark PP2A–B56 associated with the central spindle. (F) PP2A–B56 and –B55α complexes were isolated using GFP antibodies from metaphase (M) and anaphase (A) HeLa cells. Untransfected cells were used as a negative control (Mock). The immune precipitates (IP) were blotted, and the blots are representative of four independent experiments. siControl, control siRNA.
have shown that these phosphorylations have a short half-life (Neef et al., 2006; Nunes Bastos et al., 2013), until now, the KIF4A and MKlp1 phosphatases required for this dynamic behavior have not been identified. Analysis of Aurora B and Plk1 opposing phosphatases in chromosome alignment and spindle checkpoint signaling indicates that PP2A–B56 enzymes fulfill this requirement (Foley et al., 2011; Suijkerbuijk et al., 2012; Kruse et al., 2013). Because of these findings, the potential role of PP2A–B56 family enzymes as Aurora B–opposing phosphatases regulating central spindle dynamics during anaphase was investigated.

### Results and discussion

#### Identification of isoform-specific components of PP2A–B56 complexes

Human cells express five different PP2A–B56 holoenzymes differentiated by the regulatory subunit (Shi, 2009; Wurzenberger and Gerlich, 2011). To determine whether these holoenzymes are associated with anaphase-specific functions, HeLa cells were depleted of the different B56 regulatory subunits. Binucleate cells were observed after silencing of B56γ, and this increased further when B56ε was silenced in combination (Fig. 1, A and B). B56ε- and -β–depleted cells showed increased nuclear abnormalities including micronucleation, and B56β-depleted cells arrested in mitosis with aligned chromosomes (Fig. 1, A and C). Blotting with the available isoform-specific antibodies confirmed depletion of B56α, -γ-, and -β subunits alone and in combination (Fig. 1 D). Other antibodies tested for B56β and -ε failed to show isoform-specific reactivity.

The localizations of the different B56 regulatory subunits were then investigated. Comparison with the anaphase central spindle proteins Aurora B and MKlp2 showed that B56β, -γ, and -ε gave a similar pattern of localization although an additional cytoplasmic distribution was visible (Fig. 1 E and Videos 1–3). B56α- and -β–regulatory subunits showed only a diffuse localization (Fig. 1 E and Video 4). Thus, a subset of PP2A–B56 isoforms localize to the anaphase central spindle, consistent with supplemental data presented in a study of PP2A–B56 function in kinetochore fiber formation (Foley et al., 2011). These results suggest that B56γ- and -ε function in anaphase or cytokinesis, whereas B56α, -β, and -δ may be more critical for chromosome alignment and segregation.

PP2A–B56 complexes were isolated from metaphase and anaphase cells (Fig. S1, A and B). Blotting showed that the different holoenzymes were isolated with approximately equivalent efficiency judged by the enrichment of both PP2A-A scaffold and PP2A-C catalytic subunits (Fig. 1 F). Mass spectrometry of the isolated protein complexes confirmed the equivalence of the different holoenzyme complexes (Fig. S1 C) and revealed the presence of the known PP2A–B56 interaction partners BubR1 and CDC242/Repo-Man (Fig. S1 D; Suijkerbuijk et al., 2012; Kruse et al., 2013; Qian et al., 2013). In addition, two central spindle proteins, KIF4A and GEF-H1, were identified (Kurasawa et al., 2004; Birkenfeld et al., 2007). KIF4A was found in both B56γ and -ε complexes, with a slight enrichment in anaphase compared with metaphase samples (Figs. 1 F and S1 D). This behavior correlates with central spindle targeting of these holoenzymes (Fig. 1 E). The absence of the other major central spindle proteins PRC1, Plk1, MKlp1, and MKlp2 in B56γ and -ε complexes indicates that this is a specific property of KIF4A (Figs. 1 F and S1 D). GEF-H1 was detected in all PP2A–B56 complexes other than those formed by B56ε (Figs. 1 F and S1 D). This does not match the central spindle localization pattern of B56 isoforms and indicates that GEF-H1 is unlikely to be a central spindle–targeting factor for PP2A–B56. The simplest explanation for these observations was that KIF4A may act as a central spindle–targeting factor and substrate for PP2A–B56γ and -ε in anaphase. Both these possibilities were tested.

#### Central spindle localization of PP2A–B56γ and -ε is KIF4A dependent

Cells depleted of KIF4A failed to show the anaphase central spindle staining of PP2A–B56γ and -ε observed in control cells (Fig. 2 A). This was a highly reproducible and penetrant effect seen in nearly 90% of KIF4A-depleted cells (Fig. 2, B and C). Central spindle targeting of PP2A–B56β was not altered in KIF4A-depleted cells (Fig. 2, A and B). This was consistent with the mass spectrometry and Western blot data showing KIF4A was present only in PP2A–B56γ and -ε complexes (Figs. 1 B and S1 D). KIF4A therefore promotes central spindle targeting of PP2A–B56γ and -ε in anaphase, and PP2A–B56β targets to the central spindle by a KIF4A-independent mechanism.

**PP2A–B56γ is a KIF4A pT799 Aurora B site phosphatase**

In anaphase, Aurora B phosphorylates KIF4A on T799 and increases its microtubule-stimulated ATPase activity as part of the mechanism controlling central spindle length (Nunes Bastos et al., 2013). To test whether PP2A–B56γ acts directly on this site and modulates kinesin motor activity, Aurora B phosphorylated KIF4A (KIF4A pT799) was incubated with purified PP2A–B56γ, -B56β, or -B56α (Fig. 3 A). These PP2A complexes purified from interphase cells contain equal amounts of PP2A catalytic and scaffold subunits but lack detectable PP1 (Fig. 3 B). PP2A–B56γ rapidly dephosphorylated KIF4A pT799 but showed little activity toward PRC1 pT481 and pT602 (Fig. 3, A and C). The related enzyme PP2A–B56β, although present at the central spindle, was not active toward KIF4A pT799 (Fig. 3, A and C). In addition, KIF4A pT799 was not dephosphorylated by PP2A–B55α, although this enzyme was capable of dephosphorylating the pT481 Cdk1 site on PRC1 (Fig. 3, A and C), as expected (Cundell et al., 2013). Correlating with their activities toward pT799, PP2A–B56γ, and not B55α, reversed the effect of Aurora B phosphorylation on microtubule-stimulated KIF4A ATPase activity (Fig. 3 D).

The relationship between the two KIF4A-interacting forms of PP2A–B56 and the level of pT799 phosphorylation at the central spindle was then investigated. Because of their roles in error correction during mitotic spindle formation and spindle checkpoint silencing (Foley et al., 2011; Suijkerbuijk et al., 2012; Kruse et al., 2013), cells depleted of all B56 regulatory subunits arrested in a prometaphase-like state with...
that PP2A–B56γ and -ε act together to regulate turnover of Aurora B phosphorylations at the central spindle in vivo. Furthermore, they suggest that PP2A–B56 specificity for KIF4A isn’t solely determined by colocalization of both proteins at the central spindle, indicating that PP2A–B56 isoforms recognize specific determinants in their substrates.

KIF4A dynamics at the central spindle are regulated by PP2A–B56γ and -ε

Because KIF4A ATPase activity and PRC1 interaction are promoted by Aurora B phosphorylation (Nunes Bastos et al., 2013), loss of the pT799 phosphatase should cause increased levels of KIF4A at the central spindle. Moreover, inhibition of Aurora B should no longer result in the release of KIF4A from the central spindle. Live cell imaging of cells stably expressing EGFP-KIF4A confirmed both of these predictions. In cells depleted of PP2A–B56γ and -ε, the level of EGFP-KIF4A at the central spindle was increased by a little over twofold (Fig. 4, A and B). After Aurora B inhibition, EGFP-KIF4A was released from the central spindle in control cells but not in PP2A–B56γ- and -ε-depleted cells (Fig. 4, A and C). Importantly, the chromatin-associated pool of KIF4A, which is not regulated by Aurora B phosphorylation, was not altered by any of these manipulations. FRAP was then performed to analyze the binding of EGFP-KIF4A to the central spindle and chromatin (Fig. 4 D). Because the bleached volumes contain scattered chromatin and failed to enter anaphase. Likewise, cells depleted of B56β arrested in a prometaphase-like state (Fig. 1, A and B), and the time from nuclear envelope breakdown to anaphase onset when observed was 250 ± 25 min (n = 50), compared with 63 ± 12 min (n = 30) for the control. Because of these defects in anaphase entry, it was not possible to analyze the effects of combined depletion of all central spindle isoforms of B56. The KIF4A-dependent central spindle–targeted B56 regulatory subunits γ and ε were therefore silenced individually or in combination. Depletion of B56γ or -ε singly resulted in a slight elevation of pT799 staining, and this was increased when both B56γ and -ε were depleted (Fig. 3, E and F). This effect was not observed in cells depleted of B56δ (Fig. 3, E and F), which is not present on the central spindle but does appear to be required for chromosome segregation because lagging chromatin and micronucleation were observed (Figs. 3 E and 1, A–C). The requirement for B56γ and -ε for turnover of the MKlp1 pS911 Aurora B phosphorylation was then tested. Although pS911 staining was not elevated in B56γ- and -ε-depleted cells, suggesting it is maximally phosphorylated, it failed to turn over after 5 min of Aurora B inhibition (Fig. S2, A and B). Similarly to KIF4A pT799, pS911 turnover involved a combination of both B56γ and -ε. These experiments show that PP2A–B56γ can directly dephosphorylate the pT799 Aurora B site on KIF4A, modulating its microtubule-stimulated ATPase activity, and
both cytoplasmic and central spindle or chromatin-associated EGFP-KIF4A, a two-component model was used to fit the data. In both control and PP2A–B56γ- and -ε-depleted cells EGFP-KIF4A showed rapid recovery at the central spindle with a half-life of 10–11 s (Fig. 4 E). However, the mobile fraction was reduced from 62% to 47% in PP2A–B56γ- and -ε-depleted cells (Fig. 4 E), consistent with the idea that KIF4A binding sites are near saturation in the PP2A–B56γ- and -ε-depleted cells as a result of decreased dephosphorylation and dissociation of KIF4A from the central spindle. Recovery at the chromatin
chromatid separation occurred without any obvious changes in anaphase A (Fig. 5 A and Videos 5 and 6). At this point, 4 min into anaphase, sister chromatids were separated by \( \approx 4.0 \pm 1.0 \) µm in both cases (Fig. 5 B). In anaphase B, from 4 to 11 min, the central spindle extended by a further 6 µm in control cells and by 3 µm, half this extent, in PP2A–B56γ– and -ε–depleted cells (Fig. 5 B and Videos 5 and 6). Similar findings were made using fixed populations of anaphase-enriched cells, in which combined depletion of B56γ and -ε reduced mean central spindle length from 8.3 ± 0.4 µm to 5.9 ± 0.4 µm (Fig. 5 C). This was not caused by loss of the central spindle because both PRC1 and Plk1 were still present in the expected positions and at the same levels (Fig. 5 C). Depletion of B56γ and -ε alone, or other B56 regulatory subunits, did not result in reduction of central spindle length (Fig. 5 C).

General principles of PP2A–B56 and Aurora B–regulated systems

The concept that coordinated pairs of kinases and phosphatases control key regulatory phosphosites is an emerging theme in cell cycle control (Foley et al., 2011; Suijkerbuijk et al., 2012; Kruse et al., 2013; Xu et al., 2013; Espert et al., 2014). As we show here, it is also important for understanding the dynamic nature of central spindle formation and remodeling in anaphase. Rapid central spindle growth in anaphase A is promoted by PRC1 and other factors and then followed by a slower expansion during anaphase B limited by KIF4A and Aurora B (Brust-Mascher et al., 2013; Nunes Bastos et al., 2013).}

was the same under both conditions with a half-life of 5–6 s and identical mobile fraction (Fig. 4 F), indicating that association and dissociation were unaltered. PP2A–B56γ and -ε may therefore control the release of KIF4A from the central spindle by localized dephosphorylation. Further support for this view was obtained from a mathematical model defining all these biochemical reactions (Fig. S3, A and B). This model describes the accumulation of phosphorylated KIF4A (Cp) from a soluble cytoplasmic pool (K) on spindle microtubules as cells enter anaphase (Fig. S3 C, solid lines). Reduction in the soluble pool of PP2A–B56 had little effect on spindle associated KIF4A (Fig. S3 C, dotted lines). KIF4A levels on spindle microtubules increased after the removal of the spindle-associated pool of PP2A–B56 corresponding to B56γ and -ε (Fig. S3 D), similar to experimental data (Fig. 4, A and B). Aurora B inhibition caused rapid loss of KIF4A (Cp) from spindle microtubules in control conditions (Fig. S3 E) but only a slow decline after reduction of the spindle associated PP2A–B56 activity (Fig. S3 F, enlarged region), fitting well with data obtained in cells (Fig. 4 C).

**PP2A–B56γ and -ε regulation of anaphase central spindle length**

Accumulation of phosphorylated KIF4A at the central spindle would be expected to suppress microtubule dynamics and prevent full extension of this structure (Nunes Bastos et al., 2013). In both control and PP2A–B56γ- and -ε-depleted cells, sister chromatid separation occurred without any obvious changes in anaphase A (Fig. 5 A and Videos 5 and 6). At this point, 4 min into anaphase, sister chromatids were separated by \( \approx 4.0 \pm 1.0 \) µm in both cases (Fig. 5 B). In anaphase B, from 4 to 11 min, the central spindle extended by a further 6 µm in control cells and by 3 µm, half this extent, in PP2A–B56γ- and -ε–depleted cells (Fig. 5 B and Videos 5 and 6). Similar findings were made using fixed populations of anaphase-enriched cells, in which combined depletion of B56γ and -ε reduced mean central spindle length from 8.3 ± 0.4 µm to 5.9 ± 0.4 µm (Fig. 5 C). This was not caused by loss of the central spindle because both PRC1 and Plk1 were still present in the expected positions and at the same levels (Fig. 5 C). Depletion of B56γ and -ε alone, or other B56 regulatory subunits, did not result in reduction of central spindle length (Fig. 5 C).
this negatively regulates Aurora B binding to chromatin (Qian et al., 2011). In anaphase, the KIF4A-targeted pool of B56γ and -ε is ideally placed to counteract Aurora B phosphorylations on other central spindle proteins such as MKlp1. However, this complicates existing pictures of Aurora B function in anaphase, which typically envisage a gradient of Aurora B phosphorylation concentrated at the midpoint of the central spindle and minimal at either end (Fuller et al., 2008; Wang et al., 2011, 2013; Ferreira et al., 2013; Afonso et al., 2014). Although Aurora B and its counteracting phosphatases become spatially separated to permit dephosphorylation at chromatin (Wang et al., 2013; Afonso et al., 2014), as shown here, the presence of a phosphatase in the system with the same spatial distribution as the kinase is in fact necessary to explain the dynamic turnover of phosphorylation sites at the central spindle. Distribution of Aurora B alone is thus not the determining factor for phosphorylation of a given site. Instead, the functional outcome depends critically on the distribution of the counteracting phosphatase.

**Materials and methods**

Reagents and antibodies

General laboratory chemicals and reagents were obtained from Sigma-Aldrich and Thermo Fisher Scientific. Sheep antibodies were raised to MKlp1 (aa 1–444), KIF4A (full-length protein expressed in insect cells),

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Figure 5. PP2A–B56γ and -ε contribute to central spindle length control. (A) HeLa EGFP-tubulin mCherry-histone H2B cells were treated with control or combined PP2A–B56γ and -ε siRNA for 36 h, arrested with 2 mM thymidine for 18 h, and then released for 8 h into fresh medium before imaging every minute as the cells passed through mitosis and cytokinesis. Red and green lines mark the anaphase A–B transition and onset of furrowing, respectively. (B) Central spindle length is plotted as a function of time (means ± SEM, n = 8). (C) HeLa cells were treated with control, combined PP2A–B56γ and -ε, PP2A–B56α, -γ, or -ε siRNA for 36 h, arrested with 2 mM thymidine for 18 h, and then released into fresh medium. After 10 h, these cells were stained for DNA, mouse anti-Plk1, and rabbit anti-PRC1 pT602. Central spindle length in anaphase cells is shown (means ± SEM, n = 6).
and MKlp2 (aa 63–193; Neef et al., 2003, 2007; Bastos and Barr, 2010; Nunes Bastos et al., 2013). Rabbit antibodies were raised to full-length recombinant PRC1, PRC1 pT602 peptide CLIR[p(T)NIGS]KIF4A (aa 738–1,323), KIF4A pT799 peptide CLIR[p(T)FSIT], and MKlp1 pS911 peptide CRKRR[p(S)STVA (Neef et al., 2003, 2006, 2007; Grunenberg et al., 2006; Nunes Bastos et al., 2013). Specific antibodies were purified using the antigens conjugated to Affi-Gel 15, eluted with 0.2 M glycine- HCl (pH 6.0) buffer, and dialyzed against various buffers. Commercially available antibodies were used to α-tubulin (mouse DM1A; Sigma–Aldrich), Pkl1 (mouse SC-17783; Santa Cruz Biotechnology, Inc.), PRC1 pT481 [rabbit SC-11768 [Santa Cruz Biotechnology, Inc.] and rabbit monoclonal EP1514Y [Epitomics, Inc.]], the RGS-His tag epitope (mouse; Qiagen), A1M1 (mouse 61,1083; BD), and cyclin B1 (mouse G5N3; EMD Millipore). Affinity-purified primary antibodies were stored at −80°C for further use. To generate the PRC1-pT602 substrate lacking pT481 phosphorylation, 20 μg PRC1 purified from insect cells that had not been treated with okadaic acid, or nucedocazol was phosphorylated in vitro by incubating with 1.2 μg PIK1 in a final volume of 50 μl of 20 mM Tris-HCl, pH 8.0, 150 mM NaCl, 1 mM DTT, and 3 mM Mg-ATP for 1 h at 37°C. The recombinase Aurora A and the substrate PRC1 (aa 1–650) was expressed and stored at 1 μg/ml final concentration. Donkey anti–mouse, –rabbit, or –sheep/ goat secondary antibodies conjugated to HRP were obtained from Jackson ImmunoResearch Laboratories, Inc. Donkey anti–mouse, –rabbit, or –sheep/goat secondary antibodies conjugated to Alexa Fluoris 555, 647, and 488 were obtained from Invitrogen. DNA was stained with DAPI (Sigma–Aldrich).

Kinase inhibitors were obtained from Tocris Bioscience (ZM447439; 10 mM 1,000x stock) and Axon Medchem (B2536; 1 mM 1,000x stock). A bactophate microfluid (5417R; Eppendorf) was used for all centrifugations unless otherwise indicated.

Molecular biology
Human PP2A-B56α, β, γ, δ, and ε and PP2A-B55α were amplified from human testis cdna (Marathon cdNA; Takara Bio Inc.) using pfu polymerase (Agilent Technologies). Mammalian expression constructs for N-terminally FLAG or EGFP-tagged full-length KIF4A or PP2A-B55 subunits were made using the pcDNA5/FRT/TO vector (Invitrogen). The Baronase module was designed from existing crystal structures (Baylor et al., 2003; Sessa et al., 2005) and created using standard molecular biology procedures to fuse human inner centromere protein aa 834–897 plus an additional glycine residue to the N terminus of Aurora B aa 44–344 (Nunes Bastos et al., 2013). For bacterial expression, this was cloned into the pGEM3 (Qiagen) vector and placed under the control of an isopropyl β-D-1-thiogalactopyranoside-inducible hybrid T5-phae promoter. The pAcSG2 vector and BaculoGold system (Invitrogen) were used to create recombinant baculovirus carrying hexahistidine-tagged full-length PRC1 and KIF4A under the control of the Autographa californica nuclear polyhedrosis virus polyhedrin promoter. Primers were obtained from Invitrogen. The siRNA duplexes used to target P2A and KIF4A listed in Table S1 were obtained from Thermo Fisher Scientific and have been described previously (Cundell et al., 2013; Nunes Bastos et al., 2013).

Human and insect cell culture
HeLa cells were cultured in DMEM containing 10% (vol/vol) bovine calf serum [Invitrogen] at 37°C and 5% CO2. For synchronization, cells were treated for 18 h with 2 mM thymidine and washed three times in PBS and twice with growth medium. For plasmid transfection and siRNA transfection, Mirus LT1 (Mirus Bio LLC) and Oligofectamine (Invitrogen), respectively, were used. Stable HeLa cell lines with single copies of the desired transgene were created using the T-Rex doxycycline-inducible Flp-In system (Invitrogen, Carlsbad, CA). Stable cell lines expressing FLAG-PPP2R2A, or FLAG-PPP2R5C. After harvesting, cells were washed twice with growth medium. For plasmid transfection and siRNA transfection, Mirus LT1 (Mirus Bio LLC) and Oligofectamine (Invitrogen) were used at 1 μg/ml final concentration. Donkey anti–mouse, –rabbit, or –sheep/goat secondary antibodies conjugated to HRP were obtained from Jackson ImmunoResearch Laboratories, Inc. Donkey anti–mouse, –rabbit, or –sheep/goat secondary antibodies conjugated to Alexa Fluoris 555, 647, and 478 were obtained from Invitrogen. DNA was stained with DAPI (Sigma–Aldrich).

Isolation of PPP complexes from metaphase and anaphase cells
Three 15-cm dishes of thymidine-synchronized HeLa S3 cells per condition were transfected for 18 h with GFP-tagged PP2A-B56 regulatory subunits then released for 6 h at 37°C, 100 ng/ml nocodazol was added, and they were incubated for a further 12–14 h. Mitotic cells were collected by shake-off, and nocodazol was removed by washing three times with PBS and twice with growth medium warmed to 37°C. Cells were left for 2.5 min at 37°C to form bipolar metaphase spindles (metaphase sample), and then 30 min after, the 200 μM AZ3146 was added, which causes rapid spindle checkpoint inactivation and progression into anaphase (anaphase cell sample). Cells were pelleted and washed three times with ice-cold PBS. Cell pellets were suspended in 1 ml mitotic lysis buffer [20 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1% [vol/vol] IGEPA, 0.1% [vol/vol] sodium deoxycholate, 40 mM [glycerophosphate, 10 mM NaF, 150 mM NaF-vanadate, 100 mM okadaic acid, and protease and phosphatase inhibitor cocktails [Sigma–Aldrich], left for 15 min on ice, and then clarified by centrifugation at 20,000 g, for 20 min at 4°C. Protein complexes were isolated from 3 mg of cell lysate using 3 μg sheep antibodies to GFP bound to 20 μl protein G Sepharose by incubation for 2 h at 4°C. Isolated complexes were washed once with lysis buffer and then three times with 20 mM Tris-HCl, pH 7.4, 150 mM NaCl, and 0.1% [vol/vol] IGEPA. Sam- ples were then analyzed by SDS-PAGE and blotting.

Isolation of PP2A-B55 and PP2A-B56 for phosphatase assays
HEK293T cells were seeded onto 15-cm dishes (7,325 × 10 6 cells per dish) and grown for 24 h. Seven dishes per condition were transiently transfected with 8 μg pcDNA5/FRT/TO FLAG (empty vector control), FLAG-PPP2R2A, or FLAG-PPP2R5C. After harvesting, cells were washed 3x in 50 ml ice-cold PBS, centrifuged at 400 g for 3 min at 4°C, and the pellets lysed in 2 ml ice-cold T-lysis buffer [20 mM Tris-HCl, pH 7.35, 150 mM NaCl, 1% [vol/vol] Triton X-100, and protease inhibitor cocktail [Sigma–Aldrich]] on ice for 15 min. Extracts were clarified at 20,817 g for 15 min at 4°C, supernatants were isolated, and then 20 μl FLAG M2-agarose beads [Sigma–Aldrich] prewashed in T-lysis buffer was added per condition. After 3 h at 4°C, beads were washed 2x in T-lysis buffer, 4x in TBS [20 mM Tris-HCl, pH 7.35, and 150 mM NaCl] containing 0.1% [vol/vol] Triton X-100, 1x in TBS, and 1x in elution buffer [20 mM Tris-HCl, pH 7.35, 150 mM NaCl, and 1 mM MgCl2]. Bound proteins were eluted from beads in 110 μl elution buffer containing 2 μg FLAG peptide (1 mg/ml Sigma–Aldrich) at 22°C for 30 min. Glycerol [25%[vol/vol]] and DTT [1 [mM][vol/vol]] were added to the eluate before aliquoting and snap freezing.

Mass spectrometry of PP2A-B56 complexes
Protein samples for mass spectrometry were separated on 4–12% gradient NuPAGE gels and then stained using a colloidal Coomassie blue stain. Gels were cut into six slices, and the proteins were digested with trypsin. The resulting tryptic peptide mixtures were then analyzed by online liquid chromatography–tandem mass spectrometry with an ultra-performance liquid chromatography system [Proxeon; Thermo Fisher Scientific] and mass spectrometer [Orbitrap Elite; Thermo Fisher Scientific] fitted with a nano-electrospray source (Thermo Fisher Scientific). The MaxQuant 1.5.0.35 quantitative proteomics software package was used to compile and search the raw data against the human Uniprot database [Cox et al., 2009; Typanova et al., 2014].
Activity assays with purified PP2A holoenzymes

Reactions were set up on ice and performed in tubes treated with a blocking buffer (50 mM Tris·HCl, pH 7.35, 0.1 mM MnCl₂, 1 mM MgCl₂, 1 mM DTT, 0.1% [vol/vol] IGEPAL, 150 mM NaCl, and 2 mg/ml BSA) to limit nonspecific loss of protein components on tube walls. Standard assays used 1.33 µg phosphorylated KIF4A or 3 µg phospho CR1C in 280 µl of reaction buffer (50 mM Tris·HCl, pH 7.35, 1 mM MnCl₂, 1 mM MgCl₂, 1 mM DTT, 0.1% [vol/vol] IGEPAL, 300 mM NaCl, and 0.2 µM/bSA). To start the reaction, PP2A holoenzyme was added, and the samples were incubated either at 4°C or 37°C. Samples were removed at the indicated time points, added to SDS sample buffer, and boiled immediately for 5 min to stop reactions.

Kinesin motor ATPase assays

A commercial enzyme linked inorganic phosphate assay was used to measure kinesin ATPase activity (Cytoskeleton, Inc.). In a brief, a microtubule premix was created at room temperature by mixing 1 ml of reaction buffer (15 mM Pipes-KOH, pH 7, and 5 mM MgCl₂), 10 µl of 2 mM paclitaxel, 80 µl of preassembled microtubules (1 mg/ml tubulin, 15 mM Pipes-KOH, pH 7, 5 mM MgCl₂, 1 mM GTP, and 20 µM paclitaxel), 240 µl of 1 M amino-6-mercaptop-7-methylpurine riboside, and 12 µl of 0.1 U/µl purine nucleoside phosphorylase. Phosphorylated KIF4A was prepared by preincubating the motor protein with 10 mM Baranose in a final volume of 7.5 µl TND containing 1 mM Mg-ATP for 30 min at 30°C and then adding 1 µl ZM447439 to stop the reaction. Dephosphorylation was performed using 1 µl recombinant PP2A-B56-γ for 30 min on ice and stopped by adding 1 µM acetic acid. Reactions were set up in 96-well plates by mixing the protein of interest in a total volume of 7.5 µl TND with 147.5 µl of the microtubule premix at room temperature. To start the assay, 10 µl of 10 mM ATP was added to each well. Final assay volume was 165 µl of 12 mM Pipes-KOH, pH 7, 4 mM MgCl₂, 0.61 mM ATP, and 14.5 mM NaCl. This was then rapidly transferred to a 37°C plate reader (Tecan LS900B; Tecan) and read at absorbance at 360 nm. Readings were acquired every 30 s for 1 h. An inorganic phosphate standard curve was created in the same assay buffer and used to convert absorbance to nanomoles of ATP hydrolyzed.

Fixed and live cell microscopy

For fixed cell imaging, cells were grown on No. 1.5 glass coverslips (Menzel-Gläser; Thermo Fisher Scientific), washed twice with 2 ml PBS, and then fixed with 2 ml of 3% (wt/vol) paraformaldehyde in PBS for 10 min. Fixative was removed, and the cells were quenched with 2 ml of 50 mM NH₄Cl in PBS for 10 min. For the localization of GF-tagged PP2A-B56 and −B55a, the cells were fixed in −20°C methanol for 7 min. Coverslips were washed three times in 2 ml PBS before permeabilization in 0.2% (vol/vol) Triton X-100 for 5 min. In all cases, primary and secondary antibody staining was performed in PBS for 60 min at room temperature. Affinity-purified antibodies were used at 1 µg/ml, whereas commercial antibodies were used as directed by the manufacturers. DAPI was added to the secondary antibody staining solution at 0.3 µg/ml. Coverslips were mounted in Mowiol 4–88 mounting medium (EMD Millipore). Coverslips used for imaging on slices were added using a 60x, NA 1.35 oil immersion objective on an upright microscope (BX61; Olympus) with filter sets for DAPI, GFP/Alexa Fluor 488, Alexa Fluor 555, Alexa Fluor 568, and Alexa Fluor 647 (Chroma Technology Corp.), a camera (Cool SNAP HQ2, Roper Scientific), and MetaMorph 7.5 imaging software (Molecular Devices). Visualization was provided by a metal halide light source (Lumen 200 Watt; Prior Scientific Instruments, Ltd.). Image stacks of 17 planes with a spacing of 0.6 µm through the cell volume were taken. Image stacks were maximum intensity projected and then merged to create 4-bit RGB TIFF files in MetaMorph. Images in 24-bit RGB format were then cropped in Photoshop CS3 and placed into Illustrator CS3 (Adobe) to produce the figures.

For live cell imaging using spinning-disk confocal microscopy, cells were plated in 35-mm dishes with a 14-mm No. 1.5 thickness coverslip window in the bottom (Matskef Corporation). For imaging, the dishes were placed in a 37°C and 5% CO₂ environment chamber (Tokai Hit). Imaging was performed at 37°C in 5% CO₂ using an inverted microscope (IX81; Olympus) with a 60x, 1.42 NA oil immersion objective coupled to a spinning-disk confocal system (UltraVIEW Vox; PerkinElmer) fitted with an electron-multiplying charge-coupled device camera (CY1000-13; Hamamatsu Photonics). Exposure times were 30 ms for GFFKIF4A using 4% laser power. Image stacks of 24 planes spaced 0.6 µm apart were taken at 1–4 stage positions every minute for 50 s. A brightfield reference image was also taken to visualize cell shape. Maximum intensity projection of the fluorescent channels was performed in Velocity (PerkinElmer) to create 24-bit RGB TIFF files.

KIF4A FRAP data acquisition and analysis

Prebleach five image stacks of four planes with 0.4-µm spacing were acquired at 1-s intervals. Bleach was performed using an UltraVIEW Photokinesis Device (PerkinElmer) with the 488-nm laser set at 10% with following settings: 5 cycles, step size of 1, spot period of 10, spot period of 10, 1 spot cycle, small spot size, and no attenuation. Recovery images were recorded for 60 time points and analyzed as the FRAP of the FRAP data were performed using ImageJ (National Institutes of Health). Because the bleached volume contained both soluble cytoplasmic and central spindle central EGFP-KIF4A, data were analyzed using a two-component model. The normalized fluorescence intensity (I/I₀) as a function of time (t) for both components equals A x (1 − e⁻⁻⁻⁻ ⁄ kₜₚ), in which A is the mobile fraction and 1 is the rate constant. Images in 8-bit grayscale or 24-bit RGB TIFF format were then placed into Illustrator (Adobe) to produce the figures.

A mathematical model for KIF4A regulation in anaphase

The biochemical reactions providing the basis for the mathematical modeling are described in Fig. S3 A. KIF4A is present in soluble cytoplasmic (K) and central spindle associated fractions (C), which can also be phosphorylated (Kp and Cp). Aurora B kinase is described by the term A, kₚ p and kₚ p1 are rate constants for phosphorylation and dephosphorylation of KIF4A, and Kdp and Kdp1 are rate constants for microtubule binding and release of KIF4A at the spindle. In the model, the time-dependent rate of change of each component is described by a nonlinear ordinary differential equation (Fig. S3 B). Each term on the right-hand side of the ordinary differential equation corresponds to a biochemical reaction. Because the values of the parameters (rate constants and relative protein levels) are largely unknown, they have to be estimated within reasonable bounds based on known details of the system. Finally, it can be assumed that kₚ p >> kₚ p1 because Aurora B is predominantly spindle associated, and kdp > kdp1 because B56 is enriched at spindle but also present in cytoplasm. A global parameter A represents the total activity of Aurora B in the system. Phosphorylation does not alter apparent microtubule binding in pelleting assays, so kₚ p = kₚ p1; however, phosphorylation increases microtubule residence time (ln[2]/kd), so kₜ p < kₜ p1. These parameters are set to A = 1, kₚ1 = 0.2, kₚ = 5, kₚ = 10, kₚ1 = kₚ1 = 0.2, kₙ1 = 0.2, kₚ1 = 0.001, and kₜ p1 = 0.2 for the control conditions. To mimic 95% depletion of B56 y and c, kₚ is reduced to 0.05, and because it is assumed to contribute 50% of soluble PP2A−B56, kₚ1 is reduced to 0.1. To simulate the requirement for Aurora B, A was reduced to 0.1, equal to 90% inhibition. Because all available KIF4A is cytoplasmic at anaphase onset, K = 1 provides the starting condition for the simulation. Time series simulation and analysis of the model were performed with the freely available software XPPAUT (http://www.math.pitt.edu/~bard/xpp/xpp.html).

Online supplemental material

Fig. S1 shows additional characterization and mass spectrometry data for PP2A-B56 complexes. Fig. S2 shows the analysis of Mknlp1 p53P1 turnover after Aurora B inhibition. Fig. S3 provides details of the model explaining control of KIF4A by Aurora B and PP2A−B56. Table S1 lists siRNA duplexes used to target PP2A and KIF4A. Videos 1−4 show the localization of the different GF-tagged PP2A−B56, -γ, -ε, and -δ subunits, respectively, in cells passing from metaphase into anaphase and cytokinesis. Videos 5 and 6 show central spindle differences in control and PP2A−B56−depleted cells, respectively. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201409129/DC1. Additional data are available in the JCB DataViewer at http://dx.doi.org/10.1083/jcb.201409129.dv.

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