Tension promotes kinetochore–microtubule release by Aurora B kinase

To ensure accurate chromosome segregation, interactions between kinetochores and microtubules are regulated by a combination of mechanics and biochemistry. Tension provides a signal to discriminate attachment errors from bi-oriented kinetochores with sisters correctly attached to opposite spindle poles. Biochemically, Aurora B kinase phosphorylates kinetochores to destabilize interactions with microtubules. To link mechanics and biochemistry, current models regard tension as an input signal to locally regulate Aurora B activity. Here, we show that the outcome of kinetochore phosphorylation depends on tension. Using optogenetics to manipulate Aurora B at individual kinetochores, we find that kinase activity promotes microtubule release when tension is high. Conversely, when tension is low, Aurora B activity promotes depolymerization of kinetochore–microtubules while maintaining attachment. Thus, phosphorylation converts a catch-bond, in which tension stabilizes attachments, to a slip-bond, which releases microtubules under tension. We propose that tension is a signal inducing distinct error-correction pathways, with release or depolymerization being advantageous for typical errors characterized by high or low tension, respectively.

Introduction

To maintain genome integrity during cell division, kinetochores of sister chromatids attach to opposite spindle poles—bi-orientation—to ensure accurate segregation. This process relies on robust mechanisms to identify errors by distinguishing correct from incorrect attachments and to correct errors by changing the connections between kinetochores and microtubules. To identify errors, tension from opposite spindle poles is widely accepted as a signal indicating correctly bi-oriented sister kinetochores, whereas lack of tension signals an error, based on findings in several model systems (Nicklas, 1997; Sarangapani and Asbury, 2014; Lampson and Grishchuk, 2017; Mukherjee et al., 2019). Correcting errors depends on destabilizing kinetochore–microtubule interactions so that new attachments can form. Aurora B kinase is a key regulator that destabilizes these interactions by phosphorylating kinetochore substrates that bind microtubules (Lampson and Cheeseman, 2011; Carmena et al., 2012; Krenn and Musacchio, 2015). To couple error identification and correction, current models propose that phosphorylation of Aurora B substrates depends on tension through several possible mechanisms: separation of kinetochores from Aurora B at the inner centromere (Liu et al., 2009; Yoo et al., 2018; García-Rodríguez et al., 2019; Zaytsev et al., 2016), regulation of Aurora B activity (Asai et al., 2019), or Aurora B localization to a kinetochore binding site (Campbell and Desai, 2013; Broad et al., 2020; Broad and DeLuca, 2020).

An alternative model that has not been tested is that tension could regulate the downstream response to Aurora B substrate phosphorylation. Two models by which Aurora B destabilizes chromosome attachments have been proposed: release or depolymerization (Fig. S1 A; Lampson and Grishchuk, 2017). The release model, in which kinetochores detach from microtubules, is supported by abundant in vitro evidence of Aurora B substrate phosphorylation lowering the kinetochore–microtubule affinity (Carmena et al., 2012; Krenn and Musacchio, 2015; Sarangapani and Asbury, 2014). Alternatively, cellular studies suggest that Aurora B kinase induces depolymerization of kinetochore-attached microtubules without full detachment. When sister kinetochores are attached to a single pole—syntelic attachment—depolymerization pulls both toward the pole (Lampson et al., 2004), where they subsequently detach due to pole-localized activities (Ye et al., 2015; Chmátal et al., 2015). Furthermore, in the presence of mutations that mimic phosphorylation of the outer...
kinetochore protein Hec1 by Aurora B, kinetochores track the plus-ends of depolymerizing microtubules with minimal detachment (Long et al., 2017).

**Results and discussion**

To distinguish between the release and depolymerization models, we acutely recruited Aurora B kinase to kinetochores using a photocaged small molecule that heterodimerizes HaloTag and *Escherichia coli* dihydrofolate reductase (eDHFR) fusion proteins (coumarin-TMP-Halo [CTH]; Figs. 1 A and SI B; Ballister et al., 2014; Zhang et al., 2017). We fused HaloTag to the kinetochore protein SPC25 (the anchor) and eDHFR to the INBox subdomain of INCENP (the effector). INBox recruits and activates Aurora B kinase as part of the chromosome passenger complex (Carmena et al., 2012) and thus can be used to control localized Aurora B activity (Fig. 1 B; Zaytsev et al., 2016; Banigan et al., 2015). The release and depolymerization models predict different outcomes of Aurora B kinase activation. Microtubule release reduces or eliminates forces exerted by kinetochore fibers (K-fibers), whereas depolymerization increases the force pulling the kinetochore toward the attached pole. Using kinetochore movement as a readout, we tested these predictions with both monopolar and bipolar spindles, representing low- and high-tension conditions, respectively.

Inhibition of kinesin-5 (Egs) with S-trityl-L-cysteine (STLC) generates monopolar spindles so that kinetochores are under low tension with frequent syntelic attachments (Kapoor et al., 2000; Skoufias et al., 2006). Normally, the combination of K-fiber dynamics and polar ejection forces dictates chromosome positions relative to the poles (Fig. 1 C). After Aurora B kinase activation, the release model predicts chromosome movement away from the pole due to reduced forces from K-fibers, whereas the depolymerization model predicts movement toward the pole due to increased pulling forces from K-fibers. High-amplitude oscillations on monopolar spindles make it difficult to distinguish activation-induced movement from noise. To improve the signal-to-noise ratio in our analysis, we activated the entire cell to average movements over all kine-tochores and observed net poleward movement, as expected for increased Aurora B kinase activity, as recruitment of a mutant INBox in the bipolar spindle assay compared with the monopolar spindle assay that higher activity promotes more microtubule release. After an initial lag phase, the released kinetochores moved with a steady-state velocity (~1.5 µm/min) consistent with kinetochore ablation assays mimicking release (Figs. 2 F and S2 E; Sikirzhytski et al., 2014; Long et al., 2017). Furthermore, inter-kinetochore distances decreased after Aurora B kinase recruitment (Fig. S2 F), consistent with the relaxed tension predicted by microtubule release from one kinetochore. We found lower INBox recruitment in the bipolar spindle assay compared with the monopolar spindle assay (Fig. S2 G), indicating that the high frequency of release events on bipolar spindles was not due to higher Aurora B activity. Together, these results indicate that bi-oriented kinetochores under tension primarily release their attached microtubules upon phosphorylation.

To directly visualize microtubules after single kinetochore activation, we performed correlated serial-section EM. Cells were fixed after the activated kinetochore moved out of the metaphase plate, indicating release (Fig. 3 A). We defined the activated kinetochore as lagging and the sister kinetochore as leading because it moves toward its attached pole (Figs. 3 B and S2 H). We found microtubules bound to the leading kinetochore (Kb), but not to the activated lagging kinetochore (Ka; Figs. 3 B’ and S2 H’). In contrast, kinetochore pairs that were not activated aligned properly on the metaphase plate, with K-fibers attached on both sides (Figs. 3 B’ and S2 H and H’). Thus, the activated kinetochore releases its attached microtubules, losing the tug of war to the sister.

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Figure 1. Recruitment of Aurora B to syntelic kinetochores triggers microtubule depolymerization. (A) Light-induced dimerization schematic. The dimerizer has a Halo ligand (blue) linked to the eDHFR ligand trimethoprim (TMP; red), which is protected by a photoactivatable cage (purple). The kinetochore protein SPC25 anchors HaloTag at the outer kinetochore, and effectors are fused to eDHFR. Effectors are recruited to kinetochores by uncaging the HaloTag-bound dimerizer with light. (B) Constructs for this study. Dimerizer uncaging recruits eDHFR-tagged effector proteins (or Void as a negative control). The PACT domain targets to centrosomes to label spindle poles (yellow triangles; Gillingham and Munro, 2000). (C) Schematic of monopolar spindle assay, with chromosomes under a tug of war between K-fiber depolymerization and polar ejection forces. Red circles indicate Aurora B–activated kinetochores. After full or partial release of kinetochore microtubules, polar ejection forces dominate and chromosomes move away from spindle poles. In contrast, depolymerization increases poleward forces. (D–F) Representative images before and after uncaging at t = 0 (Videos 1, 2, 3, and 4). (G) Kinetochore displacement over time. For each cell, kinetochore-pole distances are measured and averaged at every time point. Displacements are defined relative to t = 0, with poleward movement defined as the positive direction (Void: n = 41 cells; MCAK: n = 46; INBox: n = 42; 2×INBox: n = 39; mean ± SEM). Scale bars, 5 μm.
Figure 2. Recruitment of Aurora B to a single kinetochore of a bi-oriented pair triggers microtubule release. (A) Schematic of bipolar spindle assay. The bi-oriented sisters are under a tug of war between the two attached K-fibers. Microtubule release or depolymerization at the activated kinetochore (red circles) leads to movement in opposite directions. (B and C) Images from representative experiments showing INBox (B; Video 6) or 2×INBox (C; Video 8) recruitment after activation of a single kinetochore (yellow triangles) at t = 0. Insets show the targeted kinetochore pair at higher magnification. (D and E) Displacement of the activated kinetochore from the metaphase plate over time: Each trace represents a single kinetochore after INBox (D; n = 64) or 2×INBox (E; n = 79) recruitment, with the starting location defined as zero. Dashed lines show the range of chromosome dynamics covering 96% of control (Void recruited) kinetochores (Fig. S2, A and B; and Video 5). Histograms show maximal displacement for each trace. (F and G) Analyses of released kinetochores after INBox or 2×INBox recruitment. Example trace of 2×INBox recruitment (F) shows waiting time after activation, followed by movement at steady-state velocity (Fig. S2 E). (G) Waiting time distribution (2×INBox: median = 1.5 min, n = 41; INBox: median = 3.5 min, n = 20). *, P < 0.005. Scale bars, 5 µm or 1 µm in insets.
The coexistence of release and depolymerization events after Aurora B kinase activation (Fig. 2, B–E) suggests an underlying kinetic race, with the release rate dominant on bipolar spindles. In addition, we found a shorter waiting time during the lag phase after recruitment of 2×INBox (median, 1.5 min) vs. INBox (median, 3.5 min), indicating that higher Aurora B activity further increases the microtubule release rate and the fraction of release events (Figs. 2 G and S2 D). To explain the differences between monopolar and bipolar spindles, one possibility is that time in mitosis affects the outcome, as Eg5-inhibited cells are arrested in mitosis for up to 2–6 h in our experiments. To test this possibility, we activated individual kinetochores in the presence of the anaphase-promoting complex/cyclosome (APC/C) inhibitor proTAME to delay mitotic exit in cells with bipolar spindles (Fig. S3, A and B). The fraction of release vs. depolymerization events and the release kinetics remained unchanged under these conditions (Fig. S3, C–E), indicating that time in mitosis is not a key variable.

We next tested whether differences in tension, which is high on bipolar spindles but low on monopolar spindles, might explain the different outcomes in these two contexts. If tension promotes microtubule release upon Aurora B kinase activation, then we predict that experimentally reducing tension on bipolar spindles would inhibit microtubule release. Reducing microtubule cross-linkers leads to a ∼10% decrease in interkinetochore distance (Elting et al., 2017; Polak et al., 2017), suggesting that interpolar microtubule arrays are mechanically coupled to K-fibers. We therefore tested whether the microtubule–cross-linking motors Eg5 or KIF15 (kinesin-12) mediate interkinetochore tension. Eg5 motors cross-link and slide microtubules apart to maintain proper pole-to-pole distance (Kapitein et al., 2005; Shimamoto et al., 2015; Uteng et al., 2008), while KIF15 motors additionally cross-link K-fibers to control chromosome movement (Sturgill and Ohi, 2013; Drechsler et al., 2014). Blocking nucleotide binding of these motors entraps them in a rigor state, which maintains cross-linking activity but diminishes the powerstroke (Elting et al., 2017; Needleman et al., 2010; Dumas et al., 2019). We found that both the Eg5 rigor inhibitor, BRD9876, and the KIF15 rigor inhibitor, KIF15-IN-1, reduce interkinetochore distances by ∼10%, indicating reduced tension (Figs. 4 A and S3, F and G). These inhibitors can stabilize microtubules against depolymerization (Chen et al., 2017); therefore, we did not use the fraction of release vs. depolymerization events after single kinetochore activation as our readout (Fig. S3 H). Instead, we measured the waiting time from activation to release. Addition of either inhibitor increased the median waiting time from 1.5 to 2.5 min (Fig. 4, B–D). These findings indicate that reduced tension slows the Aurora B–induced microtubule release rate, although microtubule stabilization by the inhibitors could also contribute.

Figure 3. Activated kinetochores lack end-on microtubule attachments. (A) Live imaging showing 2×INBox recruitment and chromosome movement after activation of a single kinetochore (yellow triangles) at t = 0, as in Fig. 2 C. (B) The same cell was fixed at 3.5 min and examined by fluorescence and serial-section EM. Top: A single focal plane shows 2×INBox overlaying SPC25 at the activated kinetochore (white arrow) and chromosomes by fluorescence, and the corresponding section from the 3D EM dataset. Kinetochores of the activated chromosome pair (B’) and a nonactivated aligned chromosome (B’’) are shown at higher magnification below (blue: DNA; green: microtubules; magenta: kinetochores). Microtubules are bound to the leading kinetochore (Kb), but not the activated lagging kinetochore (Ka). Full EM series through these two kinetochores are shown in Fig. S2 H.
Overall, our results demonstrate contrasting responses to Aurora B activity: depolymerization of kinetochore microtubules under low tension and release under high tension. This increase of the release rate with tension is opposite to the catch-bond behavior of unphosphorylated kinetochores, in which force stabilizes attachments (Akiyoshi et al., 2010). These seemingly contradictory effects of tension can be explained by a two-state model, in which the detachment rate is lower for polymerizing compared with depolymerizing microtubule plus-ends, as shown in vitro (Sarangapani and Asbury, 2014; Akiyoshi et al., 2010). At unphosphorylated kinetochores, the catch-bond is due to tension promoting the polymerizing state, which is more strongly attached (Akiyoshi et al., 2010). Kinetochore substrate phosphorylation promotes microtubule depolymerization (Umbreit et al., 2012), preventing the transition to the more strongly attached state, which converts the catch-bond into a more conventional slip-bond that breaks under increasing force. Thus, phosphorylated kinetochores maintain attachment while depolymerizing under low tension, but release under higher tension (Huis in’t Veld et al., 2019). Phosphorylation also increases the detachment rate for polymerizing microtubules (Sarangapani et al., 2013), and there may be additional complexity in the cell if phosphorylation of distinct Aurora B kinase substrates (e.g., the Ndc80 or Ska complexes or MCAK) depends on tension, promoting either microtubule depolymerization or release (Carmena et al., 2012; Krenn and Musacchio, 2015). For example, partial release may be an intermediate state leading to depolymerization while maintaining attachment of the remaining kinetochore microtubules under low tension. Furthermore, centromeric microtubule depolymerases might confer distinct functions under the control of Aurora B (Cimini et al., 2006; Bakhoum et al., 2009)—driving poleward chromosome movement or generating tension at syntelic or bi-oriented attachments, respectively.

By integrating spindle mechanics and kinetochore biochemistry, we propose that the tension-dependent response facilitates correction of distinct attachment errors (Fig. 4E). Lower tension is associated with increased Aurora B activity at syntelic attachments (Welburn et al., 2010; Liu et al., 2009), which typically scatter around spindle poles. Microtubule release would leave chromosomes to be pushed away from the spindle by polar ejection forces, whereas microtubule depolymerization pulls chromosomes toward the spindle, where they subsequently detach near the pole and congress by gliding along K-fibers of other chromosomes (Ye et al., 2015; Chmátal et al., 2015; Kapoor et al., 2006). Aurora B kinase is also activated at merotelic attachments, in which one kinetochore attaches to both spindle poles, either by recruitment or by interactions with microtubules (Knowlton et al., 2006; Salimian et al., 2011; Trivedi et al., 2019). Because tension is higher with microtubules pulling in opposite directions, phosphorylation would promote microtubule release and turnover to achieve bi-orientation (Zaytsev and Grishchuk, 2015; Cimini et al., 2006; Bakhoum et al., 2009). This mechanism can explain the failure to correct merotelic errors associated with defective spindle mechanics, such as disruptions of poleward microtubule flux, tissue architecture, or tubulin homeostasis (Ganem and Compton, 2006; Knouse et al., 2018; Lin et al., 2020).
Materials and methods

Dimerizers

Synthesis and characterization of the photo-caged (CTH) and uncaged (TNH) dimerizers followed published protocols (Zhang et al., 2017, 2018). Both stocks were dissolved in DMSO at 20 mM and distributed in amber-colored 1.5-ml microcentrifuge tubes at −80°C for long-term storage. For CTH, a stock aliquot was diluted in growth medium to a 10-µM working concentration with 1 ml vol and then kept at −80°C until use. For TNH, a stock aliquot was diluted to 10 µM in the L-15 medium without phonol red (Invitrogen) containing 10% FBS and 1% penicillin/streptomycin.

Plasmids

All constructs for anchors and effectors were integrated into the pEM7/05 backbone containing a CAG promoter for constitutive expression (Khandelia et al., 2011; Zhang et al., 2017). The kinetochore protein, SPC25, was chosen as an anchor because of its outer kinetochore localization and slow cytosolic exchange. The N terminus of SPC25 was fused to three tandem copies of HaloTag and to GFP to make 3×Halo-GFP-SPC25. The full-length inner centromere protein, CENP-B, was fused to GFP and a HaloTag at its C terminus to make CENP-B-GFP-Halo. To recruit and activate Aurora B kinase, the INBox domain (human INCENP819-918) was fused to mCherry at its N terminus and to eDHFR at its C terminus to make mCherry-INBox-eDHFR. To prevent Aurora B activation upon recruitment, the TSS motif of INBox—corresponding to residues 892–894 of INCENP—was replaced with AAA to make mcherry-INBoxAAA-eDHFR. To maximize Aurora B kinase activity, a second INBox domain was fused to the N terminus of mcherry-INBox-eDHFR to make INBox-mcherry-INBox-eDHFR (2×INBox). As a negative control, mCherry was fused to the N terminus of eDHFR to make mCherry-eDHFR (Void). As a positive control, human MCAK583 (Maney et al., 1998; Talapatra et al., 2015) was fused to the N terminus of mcherry-eDHFR to make mcherry-MCAK-eDHFR. To mark the spindle poles, the PACT (Pericentrin and AKAP Centrosome Targeting) domain was fused to GFP (PACT-GFP) in the pcDNA3.1 backbone containing cytomegalovirus and T7 promoters (Gillingham and Munro, 2000).

Cell cultures and transfection

For kinetochore recruitment, assays were performed with HeLa RMCE (recombination-mediated cassette exchange) acceptor cells (with LoxP and Lox2272 recombination sites; obtained from E.V. Makayev, Nanyang Technological University, Singapore) stably expressing 3×Halo-GFP-SPC25 (Khandelia et al., 2011; Ballister et al., 2014; Zhang et al., 2017). In brief, acceptor cells were cotransfected with a plasmid expressing Cre recombinase (10 ng) and another donor plasmid containing 3×Halo-GFP-SPC25 and puromycin selection marker inserted in a transgenic cassette flanked by LoxP and Lox2272 sites (1 µg). Cells were cultured at 37°C in growth medium containing DMEM plus 10% FBS (Clontech) and 1% penicillin/streptomycin (Life Technologies), with 5% CO2 and a humidified atmosphere. For selection, FBS (Clontech) and 1% penicillin/streptomycin (Life Technologies) were included in growth medium containing DMEM plus 10% FBS and 1% penicillin/streptomycin. For monopolar spindle assays, 250 ng of PACT-GFP and/or CENP-B-GFP-Halo plasmid was cotransfected with 1-2 µg of the eDHFR-tagged constructs—Void, INBox, or 2×INBox—for another 24 h. For photoactivation experiments, 3×Halo-GFP-SPC25 with transiently expressed INBox-mcherry-INBox-eDHFR were incubated with 20 µM STLC plus 500 nM TNH for 24 h, followed by 1 h TNH washout to remove unbound dimerizers. After quickly rinsing with PHEM buffer (60 mM Pipes, 25 mM Hepes, 10 mM EGTA, and 4 mM MgSO4 [pH 6.9]), cells were permeabilized for 1 min in freshly prepared lysis buffer (PHEM + 0.5% Triton X-100) containing 100 nM microcystin (Sigma-Aldrich), fixed by 4% formaldehyde (Thermo Fisher Scientific) in PHEM, washed 5 × 5 min in PHEM-T (0.1% Triton X-100 in PHEM), and then blocked in 20% boiled donkey serum (BDS; Sigma-Aldrich) with 2% BSA (Fisher BioReagent) in PHEM for 1 h. The primary rabbit antibody against phosphorylated Hec1-Ser44 (DeLuca et al., 2011) was incubated for 1 h after 2,000× dilution with 10% BDS plus 1% BSA in PHEM. Next, cells were washed out with 5 × 5 min in PHEM-T and then incubated with the secondary donkey anti-rabbit antibody conjugated to Alexa Fluor 647 (1:1,000 dilution in 10% BDS + 1% BSA; A-31573; Life Technologies) for another hour. Before infiltrating the coverslips with DAPI-embedded mounting medium (Vectashield; Vector Labs), cells were washed 5 × 5 min in PHEM-T and three times in PHEM to reduce nonspecific binding. All processes before the end of fixation were performed in the dark with care at 37°C and at RT for the rest of the procedure.

Photoactivation and dimerization

For photoactivation experiments, cells were incubated with CTH for 1 h, followed by a 30-min washout with growth medium to remove unbound molecules, as previously described (Zhang et al., 2017). For dimerization experiments without photo-activation (Fig. S1, D–J), 500 nM TNH was added after t = 0. For imaging, coverslips were mounted in a magnetic chamber (Chamlide CM-S22-1; LCI) with L-15 medium without phenol red (Invitrogen) containing 10% FBS and 1% penicillin/streptomycin, and then placed on a heated stage in a 37°C environmental chamber (Incubator BL; PeCon). Targeted uncaging was performed using a 405-nm laser (model #DL405-050-O, 27-mW output after fiber coupling; CrystalLaser LC) under the control of an iLas2 software module (Roper Scientific).
For monopolar spindle assays or metaphase-arrested bipolar spindle assays, 20 µM STLC (Skoulfias et al., 2006) or 20 µM proTAME (Zeng et al., 2010) was added during CTH incubation and imaging. Kinesin rigor inhibitors were added by media exchange directly on the stage, followed by a 30-min incubation to equilibrate the system. To avoid unwanted CTH uncaging during preparation stage, care was taken to minimize light and heat exposure, using low-luminescence or red light in the room and a long-pass filter to find cells under differential interference contrast microscopy.

Image acquisition
Live imaging at 37°C was performed with a confocal microscope (DM4000; Leica), equipped with a 100× 1.4 NA oil immersion objective (Leica), an XY Piezo-Z stage (Applied Scientific Instrumentation), a CSU10 spinning disk (Yokogawa), an electron multiplier charge-coupled device camera (ImageEM; Hamamatsu Photonics), and a laser merge module (LMMS; Spectral Applied Research) equipped with 488- and 593-nm lasers, as previously described (Zhang et al., 2017). To minimize photobleaching in monopolar spindle assays, images were acquired with a 1-min time interval, with 1-µm spacing for GFP and mCherry z-stacks covering 15 µm total. To improve precision for the bipolar spindle assays, the time interval was 30 s with 0.5-µm z-spacing covering 3 µm total, so not all kinetochore pairs are visualized. For fixed cell imaging, the same microscope was used with a four-line laser module (405 nm, 488 nm, 561 nm, and 639 nm; Vortran Stradus VersaLase 4).

Image processing and data analyses
Images are shown as maximum-intensity z-projections. The Fiji plug-in, TrackMate, was used to define kinetochore coordinates globally. For monopolar spindle assays, the identified objects in the targeted cell were separated as kinetochores and poles based on their intensity and quality. Kinetochore and pole coordinates were imported into MATLAB (MathWorks). The distance from each kinetochore to the center of the monopolar spindle, defined by averaging the pole coordinates, was calculated to determine the average distance at each time point. The reported displacements are average distances at each time point relative to the average distance at t = 0, with poleward movement defined as the positive direction.

For bipolar spindle assays, the position of the metaphase plate was determined by fitting the interior kinetochore ensemble using linear regression and minimizing the mean squares of x- and y-deviation, and the kinetochores of unaligned chromosomes were omitted. The Fiji plug-in, MtrackJ, was used to manually track the activated single kinetochores and identify the sister kinetochores. The relative distances between the activated kinetochore and the fitted metaphase plate were calculated in MATLAB. Movement toward the pole attached to the sister kinetochores was defined as the negative direction. Hence, the depolymerization or release models predict positive or negative directional movements, respectively, for both monopolar and bipolar spindle assays. Classification of each event in the bipolar spindle assay is based on the earliest time point distinguishable from the negative control so that the two models are mutually exclusive and in a kinetic race. For example, if an event shows depolymerization followed by release, then it is classified as a depolymerization event. The steady-state period of a released kinetochore was manually defined by the best linear fit of each trajectory moving beyond the negative threshold (see example in Fig. 2F). Experiments were repeated at least three times on different days, and data points across repeats were pooled for statistical analysis. To compare INBox recruitment in the monopolar and bipolar spindle assays (Fig. S2G), mCherry intensity at kinetochores at 2 min after activation was quantified after background subtraction.

Correlative EM
Cells were fixed for 30 min in PBS containing 2.5% glutaraldehyde (Sigma-Aldrich) immediately after the last frame of live imaging. Complete Z-series at 0.2-µm steps were then recorded to map positions of chromosomes in differential interference contrast and Hoechst 33342 fluorescence (0.1 µg/ml; Life Technologies). The images were acquired on a Nikon eclipse Ti2E microscope with a Plan Apochromat 100× 1.45 NA objective lens and Photometrics9550Prime camera at 43-nm XY pixels. EM embedding, relocation of cells, and serial sectioning were done as previously described (Rieder and Cassels, 1999). 80-nm sections were imaged on a JEM 1400 microscope (JEOL) operated at 80 kV using a side-mounted 4.0 megapixel XR401 sCMOS AMT camera (AMT). Complete image series recorded at 10K magnification were used to reconstruct partial volumes containing activated chromosomes. These volumes were aligned with the light microscopy images by matching positions of prominent landmarks, such as chromosome arms. Serial higher-magnification images (40K) were then collected to detail the distribution of microtubules in the vicinity of activated kinetochores. In two cells, we successfully activated a single kinetochore, identified the activated kinetochore based on 2×INBox fluorescence signal in fixed cells, and analyzed it by correlative EM. In both cases, the activated kinetochore lacked end-on microtubule attachments, as shown in the representative sample in Fig. 3.

Statistical tests
A nonparametric Kolmogorov-Smirnov test was used to compare the cumulative waiting time distributions, and a two-sample Z-test was used to compare the fractions of release vs. depolymerization events. Titration curves were fit by a rectangular hyperbola to measure the IC50 of interkinetochore distance for KIF15-IN-1 and BRD9876. The two-sample comparisons of velocity, interkinetochore distance, kinetochore displacement, kinetochore phosphorylation level, or INBox recruitment level were performed using Student’s t test. Unless otherwise specified in figure legends, two-tailed tests were used to compare samples.
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Author contributions: G.-Y. Chen designed and conducted cell biology experiments and wrote the manuscript. H. Zhang and A. Golden tested the feasibility of this project in the initial stage. F. Renda and A. Khodjakov acquired and analyzed EM images. D.Z. Wu and D.M. Chenoweth synthesized and characterized the CTH dimerizer. M.A. Lampson designed experiments and edited the manuscript.

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References


Figure S1. **Related to Fig. 1.** (A) Two proposed models of error correction mediated by Aurora B kinase. State 1 is the initial incorrect state. In the release model, microtubules detach from kinetochores (state 2) to allow new attachments to form. The process is iterative until bi-oriented attachments (state 3) are stabilized. In the depolymerization model, microtubules depolymerize while maintaining kinetochore attachment, generating poleward chromosome movement. Microtubules detach near the pole (state 2), followed by translocation along another K-fiber to the spindle equator (state 3), and binding of a microtubule from the opposite pole to achieve bi-orientation (state 4). (B) The photocaged and uncaged chemical dimerizers used in this study. CTH consists of the following components: A coumarin photocage; trimethoprim (TMP), which binds eDHFR; and a Halo ligand, which binds covalently to the HaloTag protein. TNH, consisting of TMP linked to NVOC (6-nitroveratryl oxycarbonyl) and a Halo ligand, was used for dimerization without requiring uncaging. (C) Representative images of the negative control (Video 2; yellow triangles label spindle poles). (D and E) Hec1-Ser44 phosphorylation after 2xINBox recruitment with TNH. Images (D) show immunostaining with a phospho-specific antibody in the presence or absence of 2xINBox recruitment to kinetochores. Quantification of Hec1-S44p signals shows higher phosphorylation with 2xINBox recruitment (n = 10 and 11 cells for nonrecruited and recruited kinetochores, respectively). Black lines: Mean ± SEM. (F–J) Depolymerization depends on Aurora B kinase activity and the location of the recruitment site. Representative images show INBox (F; n = 27) or INBoxAAA (G; n = 15) recruitment to the outer kinetochore anchor SPC25, or INBox recruitment to the inner centromere anchor CENP-B (H; n = 24), using TNH. INBoxAAA denotes alanine substitutions at the INBox TSS motif to prevent kinase activation. Images are maximum-intensity projection across z-slices that cover spindle poles for analyses. Displacements (H and J) quantified as in Fig. 1 G. Black lines: Mean ± SEM. (K–M) Aurora B kinases recruitment to kinetochores on monopolar spindles, using CTH, partially triggers microtubule release (same data as Fig. 1, E and F, Fig. S1 C, and Videos 2, 3, and 4). (K) Radial distance from the outermost kinetochores to the population median in the 10-min time window (same dataset as Fig. 1 G). Most kinetochores move toward the pole after activation, but kinetochores that release microtubules should be pushed away from the pole, leading to a large deviation (M, magenta triangles). Magenta bars indicate two of 41 control (Void-recruited) cells (4.9%), with at least one kinetochore pushed away. Cells with INBox recruitment contain fewer such kinetochores (L; 11.9%) than with 2xINBox recruitment (M; 23.1%). *, P < 0.05. Scale bars: 5 µm or 1 µm in insets.
Figure S2. Related to Figs. 2 and 3. (A and B) Representative images (A) and displacements (B) for Void recruitment to a single kinetochore (yellow triangles) of a bi-oriented pair, as a negative control. Displacements, plotted as in Fig. 2, D and E, were used to define the range of naturally occurring chromosome oscillations (dashed lines cover 45 of 47 traces, 96%; Video 5). (C) Representative images showing kinetochore movement indicating microtubule depolymerization after recruiting INBox to a single kinetochore (white triangles; Video 7). (A and C) Inset: The targeted kinetochore pair at higher magnification. Scale bars in A and C: 5 µm or 1 µm in insets. (D) The fraction of release vs. depolymerization events from Fig. 2, D and E (2×INBox: 87 ± 5% mean ± SEM, n = 47; INBox: 67 ± 9% mean ± SEM, n = 30; *, P < 0.05, one tailed). (E and F) Steady-state velocities (E) and interkinetochore distances (F) for released kinetochores. Distances are plotted before activation and at steady-state after 2×INBox recruitment. Each data point represents a single kinetochore or pair of sisters. Black lines: Mean ± SEM (n = 41 for 2×INBox, n = 20 for INBox; *, P < 0.05). (G) INBox recruitment on monopolar and bipolar spindles, measured as mCherry intensity (mean ± SEM). (H) Higher magnification of the image shown in Fig. 3 B. Full series of sections through the centromeres of chromosomes within yellow boxes are shown in H’ and H”. (H’) The activated kinetochore Ka and its non-activated sister kinetochore Kb. Microtubules have been released from Ka but are present at Kb. Arrowheads denote kinetochore plates. (H”) Non-irradiated sister kinetochores of a properly aligned chromosome within the metaphase plate. Similar number of microtubules are attached to both kinetochores.
Video 1. Related to Fig. 1 D. MCAK recruitment triggers poleward chromosome movement on a monopolar spindle. Left: 3×Halo-GFP-SPC25. Right: MCAK-mCherry-eDHFR.

Video 2. Related to Fig. S1 C. Void recruitment as a negative control for optogenetic recruitment on a monopolar spindle. Left: 3×Halo-GFP-SPC25. Right: mCherry-eDHFR.

Video 3. Related to Fig. 1 E. INBox recruitment triggers poleward chromosome movement on a monopolar spindle. Left: 3×Halo-GFP-SPC25. Right: mCherry-INBox-eDHFR.
Video 4. **Related to Fig. 1 F.** 2×INBox recruitment triggers poleward chromosome movement on a monopolar spindle. Left: 3×Halo-GFP-SPC25. Right: INBox-mCherry-INBox-eDHFR.

Video 5. **Related to Fig. S2 A.** Void-recruitment defines naturally occurring chromosome oscillations on a bipolar spindle. Left: 3×Halo-GFP-SPC25. Right: mCherry-eDHFR.

Video 6. **Related to Fig. 2 B.** INBox recruitment triggers microtubule release on a bipolar spindle. Left: 3×Halo-GFP-SPC25. Right: mCherry-INBox-eDHFR.

Video 7. **Related to Fig. S2 C.** Example of INBox recruitment triggering microtubule depolymerization on a bipolar spindle. Left: 3×Halo-GFP-SPC25. Right: mCherry-INBox-eDHFR.

Video 8. **Related to Fig. 2 C.** 2×INBox recruitment triggers microtubule release on a bipolar spindle. Left: 3×Halo-GFP-SPC25. Right: INBox-mCherry-INBox-eDHFR.

Two supplemental datasets are available online. Data S1 and Data S2 provide MATLAB codes to calculate the average distance of all kinetochores to monopolar spindle poles in a single cell and to calculate the kinetochore displacement relative to the metaphase plate of a bipolar spindle.