**Direct observation of microtubule dynamics at kinetochores in *Xenopus* extract spindles: implications for spindle mechanics**

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Microtubule plus ends dynamically attach to kinetochores on mitotic chromosomes. We directly imaged this dynamic interface using high resolution fluorescent speckle microscopy and direct labeling of kinetochores in *Xenopus* extract spindles. During metaphase, kinetochores were stationary and under tension while plus end polymerization and poleward microtubule flux (flux) occurred at velocities varying from 1.5–2.5 μm/min. Because kinetochore microtubules polymerize at metaphase kinetochores, the primary source of kinetochore tension must be the spindle forces that produce flux and not a kinetochore-based mechanism. We infer that the kinetochore resists translocation of kinetochore microtubules through their attachment sites, and that the polymerization state of the kinetochore acts a “slip-clutch” mechanism that prevents detachment at high tension. At anaphase onset, kinetochores switched to depolymerization of microtubule plus ends, resulting in chromosome-to-pole rates transiently greater than flux. Kinetochores switched from persistent depolymerization to persistent polymerization and back again during anaphase, bistability exhibited by kinetochores in vertebrate tissue cells. These results provide the most complete description of spindle microtubule poleward flux to date, with important implications for the microtubule–kinetochore interface and for how flux regulates kinetochore function.

**Introduction**

During mitosis, kinetochore fibers produce force that stretches centromere chromatin and pulls chromosomes poleward. In tissue culture cells, poleward movement appears to be generated mainly by kinetochore-based force-generating mechanisms coupled to depolymerization of microtubules at kinetochore plus end attachment sites (the “Pac-Man” mechanism; for review see Inoue and Salmon, 1995; Mitchison and Salmon, 2001). Kinetochore fibers in cultured cells also exhibit anti-poleward movement, coupled to polymerization of microtubule plus ends at kinetochores (“Pac-Man in reverse”). Likewise, kinetochores in cultured cells often oscillate between persistent phases of poleward and anti-poleward movement, behavior termed “directional instability” (Skibbens et al., 1993). Attached kinetochores in budding yeast also exhibit directional instability, suggesting that depolymerization and polymerization states of kinetochores are widely conserved throughout evolution (He et al., 2000; Pearson et al., 2001).

Another source of poleward force at attached kinetochores is poleward movement (flux) of kinetochore microtubules (Mitchison, 1989; Desai et al., 1998). Poleward flux is translocation of tubulin polymer poleward coupled to minus end depolymerization near the spindle poles (Mitchison 1989). Flux may act as a “traction fiber” mechanism for aligning chromosomes to the metaphase plate, generating tension at kinetochores for inactivating the spindle checkpoint and contributing to anaphase segregation of chromosomes (Kapoor and Compton, 2002). Spindle microtubules that exhibit poleward flux also exhibit "treadmilling" in a plus-to-minus direction when their plus ends are polymerizing (Grego et al., 2001).

In tissue culture cells, kinetochore-based mechanisms appear to make the dominant contribution to chromosome movement because kinetochore microtubule flux is slow (~0.5 μm/min) compared with the velocities of kinetochore poleward and anti-poleward movements (1.5–2.5 μm/min; Rieder and...
Salmon, 1998). In *Xenopus* egg extract spindles, flux mechanisms appear dominant because spindle microtubules exhibit average flux rates near the average rate of anaphase A
velocity (~2.0 μm/min; Desai et al., 1998). Similarly, in *Drosophila* embryo mitotic spindles, flux is fast, implying that traction fiber mechanics represent a major component of anaphase A movement (Brust-Mascher and Scholey, 2002; Maddox et al., 2002).

In *Xenopus* and *Drosophila* spindles, where flux is fast, metaphase kinetochores do not exhibit directional instability (Desai et al., 1998; Maddox et al., 2002). Chromosomes in spermatocytes, oocytes, early embryos, and higher plants also do not exhibit kinetochore oscillations. Fig. 1 shows three possible models that could explain both the lack of metaphase chromosome oscillations and the mechanism of anaphase A seen in the systems just mentioned.

In model 1, there is no flux of kinetochore microtubules. This is a possibility because kinetochore microtubules were not absolutely discriminated from the great majority of non-kinetochore microtubules in the previous analyses of flux in *Xenopus* extract spindles and *Drosophila* spindles during metaphase and early anaphase (Desai et al., 1998; Brust-Mascher and Scholey, 2002; Maddox et al., 2002). At metaphase, kinetochores pull on microtubules by Pac-Man mechanisms, but are stalled by tension between sister kinetochores and for unknown reasons do not oscillate. In anaphase, loss of centromere tension from disjunction allows kinetochores to move poleward coupled to plus end depolymerization. Model 2 is of historical significance because it is similar to mechanisms suggested by Inoue and Salmon (1995) and the treadmilling model proposed by Margolis and Wilson (1981) (for review see Mitchison and Salmon, 2001), and conclusions from
got more recent flux measurements by Desai et al. (1998) for *Xenopus* extract spindles. This model is based solely on flux. At metaphase, polymerization at kinetochores occurs at the flux rate, whereas in anaphase, polymerization stops at kinetochores. Kinetochore flux then becomes tightly bound to the microtubule lattice (“park” state), and are pulled poleward by flux. Model 3 predicts that both kinetochore motility and flux contribute to metaphase alignment and anaphase A.

To test these possibilities and other fundamental aspects of the microtubule–kinetochore interface, we used high resolution fluorescent speckle microscopy (FSM) methods (Maddox et al., 2002, 2003) to directly image microtubule polymerization/depolymerization at kinetochores relative to flux during metaphase and anaphase in spindles assembled in *Xenopus* egg extracts. Our results show that model 3 is correct, and we discuss the implications of rapid poleward flux for spindle mechanics at the microtubule–kinetochore interface.

**Results and discussion**

**Direct observation of microtubule dynamics at kinetochores**

Metaphase spindles with replicated chromosomes in *Xenopus* egg extracts (Desai et al., 1998) were labeled with a...
At metaphase, poleward flux creates tension at kinetochores

The stretch of the centromere between sister kinetochores measures tension at metaphase kinetochores (Waters et al., 1998). The analysis showed that fluorescent speckles appeared at kinetochores and moved poleward with a constant velocity of 2.0 ± 0.5 μm/min (n = 72). This result eliminates model 1.

During the analysis period, the distance between chromosomes and spindles poles changed little, allowing comparisons between rates of poleward flux in kinetochore fiber microtubules and in adjacent interpolar microtubule bundles. Fig. 2 C shows a kymograph of a nonkinetochore microtubule bundle (Fig. 2 A, closed arrow; Video 2) adjacent to the kinetochore fiber analyzed (Fig. 2 A, open arrow). Note that flux is bidirectional in the interpolar microtubule bundle near the spindle equator and unidirectional in kinetochore fibers. Fig. 2 D shows a histogram of kinetochore and nonkinetochore microtubule flux velocities relative to stationary kinetochores. Average values measured for the poleward flux of nonkinetochore microtubules was 2.3 ± 0.6 μm/min (n = 113). The 10% lower average rate of poleward flux of kinetochore microtubules relative to nonkinetochore microtubules was statistically different based on a t test (P = 0.004). This suggests that kinetochores impose a small load on flux during metaphase, although it is possible that the CENP-A antibody bound to the kinetochore slightly perturbs kinetochore function.

At metaphase, kinetochore microtubule flux occurs at slightly slower velocities compared with nonkinetochore microtubules

To test if polymerization occurs at kinetochores of metaphase chromosomes, we generated kymographs of speckle movements along aligned kinetochore fibers (Fig. 2 B).
In metaphase spindles, the average separation of sister kinetochores marked by CENP-A antibodies was 1.5 ± 0.4 μm (n = 26; Fig. 2 E). In comparison, when microtubules were completely depolymerized by 10 μM nocodazole, sister separation was 0.6 ± 0.1 μm (n = 20; Fig. 2 F). Thus, metaphase centromeres were stretched between sister kinetochores to 2.5 times their rest length.

Because microtubule plus ends polymerize at metaphase kinetochores, the primary source of kinetochore tension must be the spindle mechanisms that produce microtubule poleward flux and not a kinetochore-based mechanism. We infer that the kinetochore resists translocation of kinetochore microtubules through their attachment sites as a function of translocation velocity and the molecular viscosity of attachment (Howard, 2001). The idea that microtubule binding sites within the kinetochore produce molecular friction to a moving microtubule lattice was conceptualized by Hill (1985). The resistance at polymerizing kinetochores may be generated by transient linkages between dimers in the microtubule lattice and motor or nonmotor linker molecules within the kinetochore outer plate (see diagram in Fig. 4).

In anaphase, kinetochores switch from polymerization to depolymerization, increasing the rate of anaphase A above flux

Anaphase was initiated (Desai et al., 1998) and time-lapse series were recorded (Video 3) to test if Xenopus kinetochores pull their chromosomes poleward along stationary kinetochore microtubules (Fig. 1, model 1), become parked on the lattice of fluxing kinetochore microtubules (Fig. 1, model 2) or switch to depolymerization while flux persists (Fig. 1, model 3). Kymographs generated through aligned kinetochores and kinetochore fibers (Fig. 3, A and B) allowed measurement of polymerization (speckle slopes away from kinetochores) versus depolymerization (speckle slopes toward kinetochores) of microtubule plus ends at kinetochores (Fig. 3 C). Because the position of the right-hand sister kinetochore in Fig. 3 A was fixed in the alignment procedure, it appears as a vertical line in the kymograph, and the sister moved to the left after anaphase onset. In Fig. 3 B, the left-hand kinetochore was fixed, resulting in the sister moving to the right during anaphase. When sister chromosomes separated at anaphase onset, polymerization at the kinetochore slowed. When polymerization was slow enough, fluorescent speckles on kinetochore microtubules abruptly switched to movement toward the kinetochore, indicating microtubule depolymerization at kinetochores (Fig. 3, A and B). The velocity of plus end depolymerization added to the velocity of microtubule poleward flux, resulting in transient anaphase A rates greater than that of poleward flux. Variable switching between persistent polymerization and persistent depolymerization was observed for many kinetochores (Fig. 3 A). This switching likely accounts for the periods of fast and slow velocity seen in plots of kinetochore-to-pole movement, and likely explains why kinetochores move, on average, ∼40% faster than poleward flux of kinetochore microtubules in anaphase (Fig. 3, C and D). Only rarely and for brief periods did we see kinetochores appear to become fixed
in position at the ends of their kinetochore microtubules in anaphase. Thus, model 3 is correct and the hypothetical park state does not play a significant role in spindle mechanics in this system.

A notable observation was the distinctly bimodal distribution of polymerization/depolymerization rates at kinetochores in anaphase. We believe that this observation reflects the nature of the force–velocity relationship for kinetochores, providing fundamental insights into kinetochores as molecular engines. It suggests that the kinetochore–microtubule attachment site has two stable states of force production, and it can switch between either spontaneously or in response to applied force. The "polymerizing" state is not a neutral state (Rieder and Salmon, 1998), but a resistive, or frictional state, coupled to polymerization and plus end-directed movement (Fig. 4, A and B). Our data show that when polymerization equals flux, the kinetochore is stationary, but when sisters initially separate at anaphase onset, polymerization becomes slower than flux and the kinetochore moves poleward at a velocity given by the rate of flux minus the rate of polymerization at the kinetochore (Fig. 3 C and Fig. 4 B). Note that kinetochore resistive tension is high at metaphase when polymerization and the rate of translocation of the lattice through the attachment site equals flux (Fig. 4 A). Resistive tension becomes lower in early anaphase as sisters separate and the rate of polymerization and translocation through the attachment site decreases (Fig. 4 B). The "depolymerizing" state is the actively pulling, motile state, coupled to depolymerization and minus end-directed movement (Fig. 4 C). Fig. 3 C is the most direct and quantitative data to date supporting the idea that kinetochores are fundamentally bistable, exhibiting persistent polymerization and depolymerization states, and that bistability is a property of plus end dynamic instability (Skibbens et al., 1993; Tirnauer et al., 2002; Fig. 4). Our observation of underlying kinetochore bistability in a system where chromosomes do not oscillate in metaphase points to a conserved property that is probably fundamental to kinetochore mechanochemistry.

**Implications for spindle mechanics**

We suggest that kinetochores rarely park on the microtubule lattice in any system; rather, they are inherently bistable, switching between a force-generating depolymerizing state and a more passive, friction-generating polymerization state due to the fundamental mechanochemistry of the kinetochore–microtubule interface. Different spindles appear to vary considerably in the rate of flux, ranging from probably zero in yeast (Mallavarapu et al., 1999; Maddox et al., 2000) to slow in vertebrate somatic cells (for review see Desai et al., 1998) to a large fraction of anaphase A in meiotic and embryonic systems (Desai et al., 1998; Brust-Mascher and Scholey, 2002; Maddox et al., 2002). We believe that fundamentally conserved kinetochores behave differently in the three types of system as a response to this differing flux rate. At metaphase, we propose that chromosome oscillations occur for lower (but not higher) microtubule flux rates, because only high flux rates produce sufficient kinetochore tension to prevent kinetochores from switching to depolymerization (Skibbens et al., 1993). We also suggest that the polymerization state of the kinetochore represents a "slip-clutch" safety mechanism that prevents strong force from pulling plus ends out of their kinetochore attachment sites during chromosome congression or segregation. The slip-
ping clutch at the kinetochore allows the system to reduce the force on chromosomes if there is a mechanical problem.

Materials and methods

Labeled tubulins, labeled antibodies, and preparation of *Xenopus* extract spindles

Labeling of tubulin using X-rhodamine was performed as described previously (Waterman-Storer et al., 1998). For assaying flux, labeled tubulin was initially diluted into CSF extract 1:50, and that extract was diluted into experimental extract by 1:50 for confocal FSM. Antibodies to *Xenopus* Cenp-A were prepared and fluorescently labeled as described in the supplemental materials.

FSC

FSC was performed using diffraction-limited spinning-disk confocal fluorescence microscopy as described in detail in Maddox et al. (2002, 2003), and with a 100x/1.4 NA Plan Apochromat objective (Nikon) with 2 x 2 binning in the cooled CCD camera (model ER; Hamamatsu Corporation). MetaMorph® software (Universal Imaging Corp.) was used to control shutters, wavelength selection, image acquisition, and storage. Sequential images of different fluorophores were acquired at 1–5 s intervals, depending on the experiment. Paired images from different color channels were within 0.5 s of each other.

Data analysis

Measurements of the movement of fluorescent speckles and the leading edge of kinetochores relative to the spindle poles was mainly performed by hand tracking their positions or from kymographs using MetaMorph® software as described previously (Maddox et al., 2002). Kinetochore and speckle-to-pole movements were measured from the margin of the structure being analyzed to the end of the spindle fibers. Speckles were chosen for analysis based on the speckle remaining visible for at least five consecutive time points. “Custom Align” and “Rotation” algorithms in MetaMorph® were used to align spindles throughout the time-series stack of images to a given point (e.g., a single kinetochore). Color overlays were used for both kymograph methods to compare movements of microtubule fluorescent speckles to kinetochores. Velocities were obtained from the slopes of the speckle or leading-edge trajectories in the kymograph images. Statistical analysis and graphs were done in Excel (Microsoft).

Online supplemental material

Online supplemental materials consist of methods and materials for EM and antibody production. Also included is a figure comparing wide-field imaging to confocal imaging and a figure showing thin-section EM of *Xenopus* extract kinetochores. Three movies corresponding to Fig. 2 and Fig. 3 are also included. Online supplemental material available at http://www.jcb.org/cgi/content/full/jcb.200310088/DC1.

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