The Ndc80 Kinetochore Complex Forms Load-Bearing Attachments to Dynamic Microtubule Tips via Biased Diffusion

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SUMMARY

Kinetochores couple chromosomes to the assembling and disassembling tips of microtubules, a dynamic behavior that is fundamental to mitosis in all eukaryotes but poorly understood. Genetic, biochemical, and structural studies implicate the Ndc80 complex as a direct point of contact between kinetochores and microtubules, but these approaches provide only a static view. Here, using techniques for manipulating and tracking individual molecules in vitro, we demonstrate that the Ndc80 complex is capable of forming the dynamic, load-bearing attachments to assembling and disassembling tips required for coupling in vivo. We also establish that Ndc80-based coupling likely occurs through a biased diffusion mechanism and that this activity is conserved from yeast to humans. Our findings demonstrate how an ensemble of Ndc80 complexes may provide the combination of plasticity and strength that allows kinetochores to maintain load-bearing tip attachments during both microtubule assembly and disassembly.

INTRODUCTION

Chromosomes are organized and separated during cell division by a molecular machine, the mitotic spindle. A central component of this machine is the kinetochore, a specialized site on each chromosome that forms attachments to spindle microtubules. Besides mechanically coupling the chromosomes to microtubules, kinetochores are also regulatory hubs. They control chromosome movement by altering the growth and shortening of attached microtubules, they sense and correct errors in attachment, and they emit diffusible “wait” signals until they are properly attached. These regulatory functions are all vital for accurate mitosis, and they are all related to microtubule attachment. To understand kinetochore function, we must uncover the basis for its most fundamental behavior—microtubule attachment.

As the spindle forms, kinetochores make initial attachments to the sides of spindle microtubules. These lateral (“side-on”) attachments convert quickly into tip (“end-on”) attachments and, thereafter, chromosome movement is coupled to filament assembly and disassembly (Rieder and Salmon, 1998). Kinetochore-microtubule coupling requires a puzzling combination of strength and dynamicity. Molecules at each kinetochore-microtubule interface must maintain a persistent, load-bearing attachment to the filament tip even while thousands of tubulin subunits are added and removed. This strong yet dynamic coupling allows kinetochores to harness microtubule shortening to drive poleward chromosome movement and also to track with growing tips during antipoleward movement (or during microtubule “flux”). While the behavior is conserved from yeast to humans (McAinsh et al., 2003; Rieder and Salmon, 1998), its molecular basis remains uncertain.

Models for kinetochore-microtubule coupling fall into several classes. In one class, coupling is based on ATP-powered motor enzymes. Motors likely contribute to coupling in many organisms (Sharp et al., 2000; Yang et al., 2007), and some motors can mediate attachment to disassembling tips in vitro (Lombillo et al., 1995). However, kinetochore-bound motors are largely dispensable in yeast (Grishchuk and McIntosh, 2006; McAinsh et al., 2003), and their depletion in higher eukaryotes does not completely disrupt tip attachment (Kapoor et al., 2006; Weaver et al., 2003; Yang et al., 2007). In a second class of models, peeling protofilaments at disassembling tips drive kinetochore movement either by pushing against a sliding ring (Koshland et al., 1998) or by tugging on tightly bound kinetochore fibrils (McIntosh et al., 2008). Enthusiasm for the ring model was generated recently by the discovery that the Dam1 kinetochore complex forms rings around microtubules in vitro (Miranda et al., 2005). However, while the Dam1 complex is very likely to contribute to tip coupling, microtubule-based force production, and regulation of kinetochore-attached microtubules in yeast (Asbury et al., 2006; Franck et al., 2007; Tanaka et al., 2007;
Westermann et al., 2006), no clear homologs have been identified outside fungi. Moreover, ring formation is not required for microtubule-driven movement of the Dam1 complex in vitro (Gestaut et al., 2008). The fibrils model offers an attractive, ring-independent way for peeling protofilaments to exert force on the kinetochore (McIntosh et al., 2008). However, progressive disassembly-driven movement by this mechanism requires tight binding of the fibrils to the microtubule, which would prevent assembly-coupled movement. Thus, motor-, ring-, and tight fibril-based mechanisms do not appear to fully account for tip coupling in all eukaryotes.

In a third class of models, kinetochore-microtubule coupling depends on biased thermal diffusion. This idea was first proposed by Hill (1985), who demonstrated theoretically that an array of individually weak microtubule binders will track with disassembling microtubule tips provided that its diffusion along the filament is sufficiently fast and that its total binding energy is large enough. Because this mechanism provides molecular friction that resists translocation of the microtubule through the attachment, it is sometimes called a “slip clutch” (Maddox et al., 2003). Importantly, a biased diffusion-based coupler supports tension continuously while remaining tip attached during both assembly and disassembly. This behavior distinguishes it from models based on ATP-powered motor proteins or peeling protofilaments, which are fundamentally unidirectional. Although Hill imagined a sleeve-like arrangement of binding elements completely encircling the filament, the physical underpinnings of his model can apply generally to any array of microtubule binders that are linked together, even a disordered array.

Extensive genetic, biochemical, and structural studies have established that the Ndc80 complex is crucial for maintaining a robust attachment between kinetochores and microtubules that are linked together, even in disordered arrays (reviewed in Tanaka and Desai, 2008). However, the ability of the complex to support the strong yet dynamic attachment required for kinetochore-microtubule coupling has never been demonstrated. Here, we show that ensembles of Ndc80 complexes can form load-bearing attachments to microtubules for assembly and disassembly tips. The properties of this attachment can be fully described by a biased diffusion mechanism. Based on these findings, we propose that the Ndc80 complex forms a slip clutch at the kinetochore.

RESULTS

The Ndc80 Complex Couples Cargo to Dynamic Microtubules

To study Ndc80-based coupling, we adapted a recently developed bead motility assay (Asbury et al., 2006; Franck et al., 2007). All four subunits of the S. cerevisiae Ndc80 complex—Ndc80, Nuf2, Spc24, and Spc25—were coexpressed in E. coli and purified by affinity chromatography followed by gel filtration (Gestaut et al., 2008; Wei et al., 2005) (Figure S1 available online). Purification and binding of the complex to beads were facilitated by a His6 tag on the C terminus of Spc24, a location near the kinetochore-facing end of the complex (~20 nm away) and farther (~40 nm) from the N-terminal regions of Ndc80 and Nuf2 shown previously to mediate binding to taxol-stabilized microtubules (Cheeseman et al., 2006; Ciferri et al., 2008; Wei et al., 2005, 2006). Beads decorated with the complex were introduced into a chamber containing dynamic microtubules growing from stable, coverslip-anchored seeds. The filaments exhibited dynamic instability typical for microtubules in vitro, switching stochastically between periods of slow growth (at ~10 nm s−1) and rapid shortening (~100 nm s−1). To test for binding, we captured candidate beads diffusing freely in solution with a laser trap and held them near the tip of a growing filament. Binding was evident from a sharp reduction in the Brownian motion of the bead, which was monitored with high spatiotemporal resolution using back focal plane interferometry (Asbury et al., 2006; Franck et al., 2007). Beads coated with full-length Ndc80 complex almost always bound microtubules (82%, 42 of 51 beads tested). Control beads coated with a “headless” version of the complex, 2NCC:2S, lacking the N-terminal regions of Ndc80 and Nuf2 (Wei et al., 2005), failed to bind (116 tested). Thus, binding of Ndc80-coated beads to assembling microtubule tips in our assay requires the same molecular domains that mediate binding of the free complex to taxol-stabilized microtubules.

In our initial experiments, we shut off the laser trap when the bead bound the microtubule tip to record movement in the absence of external load. Without load, the beads exhibited longitudinal diffusion along the filament as the tip continued to grow (65%, 26 of 40 beads tested) (see Figure 1) or remained stationary on the microtubule lattice (35%, 14 of 40). Most attachments persisted until the filament tip underwent a spontaneous “catastrophe,” disassembling rapidly back to the bead and then carrying it in the direction of shortening (73%, 29 of 40), usually all of the way to the coverslip-anchored seed (52%, 21 of 40). Most of the remaining events were terminated artificially (40%, 16 of 40), often because the bead stuck irreversibly to the coverslip. Spontaneous detachment from the microtubules was rare (6%, 3 of 40).

We used automated centroid tracking from video recordings to quantify bead position over time (Figure 1C). The Ndc80-based attachments remained bound to the filaments for a mean duration of 180 ± 30 s (Figure 1F) and underwent disassembly-driven movement over 4800 ± 400 nm on average (Figure 1E). These values underestimate the capacity of the linkages to remain bound and track with disassembly because most events were terminated by the microtubule shortening completely back to the seed, not by detachment. For the subset of beads that diffused along the filaments, mean-squared displacement increased linearly with time and indicated a diffusion constant of 2300 ± 400 nm2 s−1 (Figure 1D). The diffusion was unbiased. Diffusing and nondiffusing beads alike were able to track with disassembly (Figure 1C). These observations demonstrate that the Ndc80 complex forms persistent dynamic attachments to microtubules, allowing filament disassembly to drive movement.

Ndc80-Based Couplers Harness Microtubule Disassembly to Produce Force

Attachments between kinetochores and microtubule tips in vivo withstand tensile forces arising during congression and segregation, but kinetochores lacking the Ndc80 complex fail to
establish such attachments (DeLuca et al., 2005; He et al., 2001). To apply force to Ndc80-based linkages in vitro, we used a laser trapping-based force clamp. Constant tension was applied in the direction of microtubule growth with a computer-controlled piezoelectric stage programmed to maintain a fixed offset between the bead and the trap center (Asbury et al., 2006; Franck et al., 2007). Even with 0.5 to 2.5 pN of continuous tension, the bead-bound Ndc80 complexes maintained an attachment to the filament that persisted through phases of assembly and disassembly and often lasted several minutes (Figure 2 and Table S1). During microtubule growth, the linkages behaved differently under tension than in its absence. Instead of diffusing along the filament or remaining stationary (as in Figure 1), they stayed associated with the growing tip, traveling an average of 670 ± 70 nm before the tip switched spontaneously into rapid shortening (20%, 64 of 314 events) or until movement was interrupted by bead detachment (72%, 226 of 314) or some other event (e.g., the bead sticking to the coverslip or colliding with another bead; 8%, 24 of 314). Many beads remained attached during disassembly (142 events, including 78 that were not preceded by discernable assembly-coupled movement) (see Table S1) and were carried, on average, 1080 ± 140 nm in the direction of filament shortening, against the force of the laser trap (Figure 2E). A few reversed direction when the filament underwent a “rescue,” spontaneously resuming growth (3%, 4 of 142). The remaining beads detached during disassembly (58%, 83 of 142) or were otherwise interrupted (39%, 55 of 142). These results establish that Ndc80 complexes can form load-bearing tip attachments that remain bound during microtubule growth, shortening, catastrophe, and rescue and that the attachments are capable of harnessing energy released during filament disassembly to produce mechanical work.

**Coupling and Force Production Are Conserved Properties of the Ndc80 Complex**

To determine whether the ability to form dynamic, load-bearing tip attachments is conserved in the human Ndc80/Hec1 complex, all four subunits were coexpressed in *E. coli* and purified by affinity chromatography followed by gel filtration (Figure S1). As above, a His6 tag on the C terminus of Spc24 was used both for purification and bead binding. Beads coated

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**Figure 1. The Ndc80 Complex Couples Cargo to Dynamic Microtubules**

(A) An Ndc80-coated bead diffuses back and forth over the microtubule lattice and fails to track with the tip during filament growth (0–600 s). During filament shortening, it tracks with the disassembling tip (600–690 s). Selected frames from Movie S1 are shown. (White arrows) The coverslip-anchored portion of the microtubule seed. (Red crosses) The bead center. (Yellow arrowheads) The microtubule tip. Elapsed times are in seconds.

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(B) Schematic diagram of the Ndc80 bead motility assay. In the absence of tension, Ndc80-based linkages usually exhibit lattice diffusion as the filament grows by addition of tubulin subunits to the tip, and they remain tip bound even as subunits are rapidly lost from the tip during shortening.

(C) Records of bead position versus time, measured without applied force. Increasing position represents movement away from the anchored portion of the microtubule. For clarity, the records are offset vertically by an arbitrary amount.

(D) Mean-squared displacement plotted against time for the subset of beads that underwent lattice diffusion. Data are mean (solid red curve) ± SEM (dotted red curves), computed from n = 26 records. (Black line) Linear fit used to determine diffusion coefficient.

(E and F) Histograms of bead displacement during microtubule shortening (n = 29 records) and total duration (including growth and shortening phases) of microtubule attachment (n = 40) for a population of beads. Dotted vertical lines indicate mean values. The data in (D)–(F) are pooled from experiments using beads prepared with 0.6 to 15 nM Ndc80 complex, corresponding to molar ratios of 110 to 2700 complexes per bead.
with the full-length Ndc80/HeC1 complex behaved similarly in force-clamp experiments to those coated with the yeast complex. After attachment to a growing tip and placement under 0.5 to 2.5 pN of tension, the beads often remained tip bound for many minutes through phases of filament growth and shortening (Table S1 and Figure S2A). The mean displacements during filament assembly, 1900 ± 300 nm (n = 46), and disassembly, 970 ± 270 nm (n = 25), and the average duration of attachment, 280 ± 50 s (n = 50), were comparable to those measured for the yeast complex (see Figures S2B–S2D). These results show that the capacity of the Ndc80 complex to couple force to dynamic microtubule tips is conserved from yeast to humans.

**Coupling and Force Production Persist at Low Surface Densities**

Kinetochores in vivo contain 6 to 30 copies of the Ndc80 complex per attached microtubule (Emanuel et al., 2005; Joglekar et al., 2008). To estimate the number of complexes required in vitro for a load-bearing coupler, we measured coupling performance as a function of the surface density of Ndc80 on the beads. Anti-His beads (at 5.6 pM) were incubated with the Hi6-tagged yeast Ndc80 complex at molar ratios ranging from 110 to 2700 complexes per bead. At each molar ratio, we measured displacements for a population of tip-attached beads subject to 1 pN of continuous tension during microtubule growth and, if possible, during shortening. We also calculated detachment rates for each condition by counting the number of detachment events and dividing by the total observation time. At molar ratios of ~360 complexes per bead and higher (i.e., ≥ 2 nM Ndc80 complex), we routinely observed tip attachments that persisted for minutes and moved 400 to 1000 nm during growth and shortening (Figure 3A). Below ~360 complexes per bead, the detachment rate from growing tips increased (Figure 3B), an effect that reduced the distance moved during growth and made measurements during shortening more difficult (because fewer beads remained bound long enough for the filament to undergo catastrophe). Below ~110 complexes per bead (<0.6 nM Ndc80 complex), microtubule binding still occurred, but the beads detached within a few seconds when placed under tension. These observations suggest a lower bound for the formation of a load-bearing coupler of ~360 complexes per bead.

Simple geometric considerations indicate that Ndc80 complexes on only a small fraction of the bead surface, ≤ 4%, can simultaneously interact with the microtubule (Figure 3C). Assuming a random distribution of complexes on the bead surface, 360 complexes per bead corresponds to 14 ± 5 interacting with the filament. This calculation may overestimate the number of interactions because it assumes (1) that all Ndc80 complexes are functional, (2) that they bind very efficiently to the bead surface, (3) that every bead-bound Ndc80 complex within 40 nm can attach to the filament, and (4) that the bead rests directly against the side of a filament whose tip extends > 120 nm past the point of contact. Relaxing any of these assumptions would lead to a lower estimate. We also note that our estimate includes a relatively large uncertainty (± 33%), arising from the inherent uncertainty in our measurements of bead and protein concentration and from counting statistics (see Supplemental Discussion). Nevertheless, we can conclude that the number of microtubule-interacting Ndc80 complexes required to form a load-bearing coupler in our bead assay is close to the 6 to 30 found at kinetochores in vivo.

**Individual Ndc80 Complexes Exhibit Diffusion along Microtubules**

To study interactions between Ndc80 complexes and microtubules at the level of single molecules, we applied total internal
reflection fluorescence (TIRF) microscopy (Figure 4). A fluorescently tagged version of the yeast complex was created by fusing green fluorescent protein (GFP) to the C terminus of Nuf2, a modification that causes no detectable phenotype in vivo (Shimogawa et al., 2006). At the low concentrations of free Ndc80 necessary for recording single molecules (i.e., <10 nM), we observed very little binding in a standard microtubule growth buffer. However, in buffers with reduced ionic strength and at an Ndc80-GFP concentration of 20 pM, individual fluorescent particles of Ndc80 complex bound transiently to taxol-stabilized microtubules and underwent one-dimensional diffusion along the filaments (Figure 4B). The brightness of the particles suggested that they were composed of one or two complexes. Most disappeared in a single step, which probably represents detachment from the microtubule (Figure 4C). However, a small fraction showed a stepwise loss of half of their intensity while they remained attached to the filament, consistent with photobleaching of one GFP molecule within a particle containing two. We fit the initial brightness distribution for all of the particles by the sum of two Gaussian functions (Figure 4D), corresponding to a large population (89%) with a unitary brightness of 16,200 ± 5500 arbitrary units (a.u.) and a small population (11%) with twice the brightness (32,400 ± 5500 a.u.). The unitary brightness was similar to that of individual GFP molecules, 17,000 ± 11,000 a.u., measured in control experiments in which GFP-MCAK was bound tightly to microtubules under otherwise identical conditions (data not shown).

Sensitivity to ionic strength is a hallmark of protein-protein interactions that rely on electrostatic attraction. To quantify the sensitivity of Ndc80-microtubule interactions to ionic strength, we recorded particle binding and diffusion on taxol-stabilized microtubules across a range of total potassium ion concentrations from 30 to 90 mM (Figure 5). At each potassium ion concentration, we measured association and dissociation rates, $k_\text{on}$ and $k_\text{off}$, and used these values to estimate an equilibrium dissociation constant, $K_\text{d} = k_\text{off}/k_\text{on}$. Diffusion coefficients were estimated for each condition from linear fits to plots of mean-squared displacement versus time. Raising the ionic strength also caused a modest (50%) increase from 0.23 ± 0.02 μM at 30 mM potassium to 0.84 ± 0.12 μM at 90 mM potassium (Figure 5D). These $K_\text{d}$ values fall within the range of previous estimates, which vary from 0.04 μM to 3 μM depending on the Ndc80 construct and the buffer composition (Cheeseman et al., 2006; Ciferri et al., 2008; Wei et al., 2007).

Raising the ionic strength also caused a modest (50%) increase in the apparent diffusion coefficient, which went from 0.117 ± 0.004 μm$^2$ s$^{-1}$ at 30 mM potassium to 0.17 ± 0.01 μm$^2$ s$^{-1}$ at 90 mM potassium (Figure 5E). We note that these changes are unlikely to reflect differences in the oligomeric state of the complex because particle brightness values were indistinguishable at the three ionic strengths (Figure S3). Collectively, these results show that particles consisting of one or two Ndc80 complexes form transient, electrostatically based interactions with the microtubule that allow rapid diffusion along the filament lattice. Lattice diffusion implies that the energetic barrier for complete detachment of the particles from the microtubule is
higher than for movement between neighboring sites on the filament lattice. This confirms a key property required for effective tip coupling via Hill's biased diffusion mechanism.

**Ensembles Are Required for Tracking with Disassembling Tips**

Another critical assumption of Hill's model is that, when the coupler encounters a disassembling tip, its otherwise random diffusion must become biased in favor of movement back toward the filament lattice. To test this, we modified the TIRF assay to study interactions with dynamic rather than stabilized microtubules. Short, GMPCPP-stabilized microtubule seeds were anchored to the coverslip, and dynamic extensions of several micrometers were grown from them by the addition of free tubulin and GTP. Fluorescent Ndc80 complex was then introduced by buffer exchange, which also lowered the level of free tubulin to trigger filament disassembly. We used 1 nM Ndc80 complex, a concentration at which the microtubules were crowded with diffusing particles, so that each tip would encounter many particles as it disassembled. No clear tip tracking was evident.

However, the density of particles sometimes (15%, 20 of 132 tips recorded) became noticeably higher at a disassembling tip as compared to the lattice. When individual particles could be discerned as they encountered a disassembling tip, they sometimes appeared to “bounce” repeatedly off the tip, diffusing back onto the lattice rather than dissociating (Figure 6A, top). These observations confirm that the diffusion of Ndc80 particles becomes biased at disassembling tips, as required for the biased diffusion mechanism.

Given the results from our bead assays, we reasoned that tip tracking might occur in our TIRF assay if larger particles containing more complexes could be generated. To test this, we recorded the behavior of Ndc80 complexes that had been preincubated with antibodies directed against the His6 tag on Spc24 to drive oligomerization. After antibody treatment, the fraction of disassembling tips that accumulated fluorescent particles was higher (46%, 17 of 35 tips), and they were 3-fold brighter than those in control experiments without antibody treatment (Figure 6B). In addition, some brighter particles in the antibody-treated samples exhibited clear episodes of processive tip tracking (Figure 6A, bottom). Together, these observations indicate that tip tracking is not a property exhibited by individual Ndc80 complexes. Assembly of an array of complexes is required.

**Biased Diffusion Fully Describes the Behavior of Ndc80-Based Couplers**

To test whether the biased diffusion mechanism can account for the performance of Ndc80-based couplers, we used computational (Monte Carlo) methods to simulate Hill’s model (Hill, 1985) with parameters chosen to match the properties of
individual Ndc80 complexes. The simulations recapitulate our experimental observations very closely (Figure 7). When bound to growing microtubules in the absence of external load, the simulated couplers have no tendency to follow the tip. Instead, they remain on the microtubule lattice, where they exhibit unbiased diffusion at a rate that slows as the number of binders in the ensemble increases (Figure 7B). At disassembling tips, their random movement becomes biased. Small, highly diffusive couplers (e.g., with \( M = 2 \) binding elements) tend to stay ahead of the disassembling tip and only occasionally bounce off of the tip (Figure 7A). In contrast, couplers with more elements track unidirectionally with disassembly (Figure 7B). Interestingly, couplers with very high numbers of binding elements (e.g., \( M = 100 \)) diffuse at undetectably slow rates, yet they still track with disassembling tips (see Supplemental Discussion). When tensile loads are applied, couplers with very few binding elements detach quickly, but larger couplers can sustain substantial forces indefinitely. Couplers with \( M = 14 \) elements, for example, support 2.5 pN while remaining persistently attached to the filament tip over many micrometers of growth and shortening (Figure 7C). These observations show that the behavior of Ndc80-based couplers is well described by a biased diffusion mechanism.

**DISCUSSION**

The Ndc80 Complex Can Make a Direct and Significant Contribution to Kinetochore-Microtubule Coupling

We measured the mechanical performance of Ndc80-based tip attachments in experiments that recapitulate the in vivo situation far more closely than previous assays. Even in the absence of other kinetochore components, we find that the Ndc80 complex can form dynamic tip attachments that permit assembly- and disassembly-coupled movement over distances comparable to the excursions of tip-attached kinetochores in vivo, which range from 0.5 to 5 \( \mu \text{m} \) depending on the cell type (Pearson et al., 2001; Stumpff et al., 2008). Under continuous tensile load, the linkages often persist for many minutes, similar to the 2 to 10 min during which kinetochores support tension in yeast (Pearson et al., 2001). Also, the number of Ndc80 complexes required to form a load-bearing coupler in our assay is close to the 6 to 30 complexes found per microtubule at kinetochores in vivo (Emanuele et al., 2005; Joglekar et al., 2008).

The loads that we applied here are comparable to the forces supported by kinetochore-microtubule junctions in vivo. Thermodynamic considerations suggest that a disassembling
microtubule tip might be capable of producing as much as 20 to 60 pN of tensile force (Desai and Mitchison, 1997). However, based on the amount of stretching between sister kinetochores and the elasticity of the chromatin that links them, the tension transmitted to bioriented sister kinetochores during normal chromosome movement is probably closer to 0.4 to 8 pN per attached microtubule (Brower-Toland et al., 2002; Nicklas, 1988; Pearson et al., 2001; see Supplemental Discussion).

Ndc80-based linkages in our assay support similar forces of 0.5 to 2.5 pN, so the experiments provide a close approximation to the physiological situation.

**Ndc80-Based Coupling Likely Depends on Biased Diffusion**

At the level of individual complexes, Ndc80 binds microtubules transiently, exhibiting moderate (micromolar) affinity and rapid

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**Figure 6. Small Ensembles of Ndc80 Complexes Track with Disassembling Microtubule Tips**

(A) Kymographs showing movement of particles composed of Ndc80 complexes (green) on disassembling microtubules (red). At 1 nM Ndc80 complex, the microtubules were crowded with diffusing particles, many of which encountered the disassembling tip. No clear tip tracking was evident. However, in some cases, the density of particles became noticeably higher at a disassembling tip relative to the lattice. When individual particles could be discerned as they encountered a disassembling tip, they sometimes appeared to bounce repeatedly off of the tip (upper kymograph). After pretreating the Ndc80 complex with anti-His antibody to drive oligomerization, more disassembling tips accumulated fluorescent particles, and some long episodes of processive tip tracking were observed (lower kymograph). Scale bars, 3 μm.

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**Figure 7. Simulated Behavior of a Diffusion-Based Coupler**

(A–C) Output from Monte Carlo simulations with parameters defined as in Hill (1985) but chosen to match our observations. The number of binding elements in the coupler, M, was varied between 2 and 100, as indicated. In (A) and (B), no external load was applied. The couplers in (A) were initially located at a disassembling tip. In (B), they were initially located on the lattice, where they exhibited unbiased movement for several minutes until being captured and carried by the disassembling tip (denoted by dotted lines). In (C), the couplers were initially located at an assembling tip, which later began to disassemble (>200 s). A load of 2.5 pN was applied in one case (black trace), whereas no load was applied in the other case (red trace). The other parameter values were held constant as follows: $r = e^{-b_{\text{tip}}} = 0.75$, $s = e^{-w_{\text{tip}}} = 10^{-6}$, $l = 0.62$ nm, $x_{\text{tip}} = 0.170$ μm$^2$ s$^{-1}$, $a = 10$ nm s$^{-1}$, $js = 100$ nm s$^{-1}$. Note that the parameters, $s$, $l$, $x_{\text{tip}}$, $a$, and $js$, are constrained by the binding equilibrium for single Ndc80 complexes, the spacing of tubulin subunits in the microtubule lattice, the diffusion speed for single Ndc80 complexes, and the microtubule growth and shortening speeds, respectively. The only free parameter, $r$, was chosen to fall in the same range considered by Hill. For clarity, the records are offset vertically by arbitrary amounts.
lattice diffusion. These properties make it ideally suited for efficient tip coupling through a biased diffusion mechanism (Hill, 1985). A recently proposed alternative view that the complex mediates processive attachment to depolymerizing tips by binding tightly to peeling protofilaments (McIntosh et al., 2008) seems less likely given the mobility and transiency of the Ndc80-microtubule interaction (but the mechanism could apply to other kinetochore components). Instead, a mechanism based purely on biased diffusion is sufficient to describe the ability of Ndc80-based couplers to mediate attachments to both polymerizing and depolymerizing tips. Coupling by this mechanism relies on a gradient in binding energy, which exists at any tip, independent of whether it is assembling or disassembling. Importantly, the mechanism accounts for the ability of Ndc80-based couplers to sustain tensile forces comparable to those found in vivo. It will also provide molecular friction that resists movement of the microtubule through the attachment (Hill, 1985). Together, these observations suggest that the Ndc80 complex forms a slip clutch at the kinetochore.

General Implications for Kinetochore-Microtubule Coupling

The idea that the Ndc80 complex forms a slip clutch at the kinetochore can explain aspects of spindle physiology. For example, when 9G3 antibodies against Ndc80/Hec1 are injected into dividing cells, kinetochore-microtubule attachments are stabilized, the chromosomes stop oscillating, sister kinetochores become hyperstretched, and kinetochore fiber flux slows (DeLuca et al., 2006). These changes are all consistent with an increase in molecular friction at the kinetochore. If an ensemble of Ndc80 complexes provides a kinetochore slip clutch, then antibody treatment could dramatically increase its friction by rafting the complexes together or by otherwise tightening their interaction with the filament (e.g., by blocking phosphorylation). The slip clutch idea can also explain the seemingly incongruous observations that the Ndc80 complex possesses only moderate microtubule affinity, yet its depletion results in severe detachment defects without major loss of other microtubule-binding kinetochore components (DeLuca et al., 2002). Notably, the affinity of the complex for microtubules depends on flexibly tethered positive charges that interact with negative charges on the E hook of tubulin (Ciferri et al., 2008; Wei et al., 2007). This pliant, electrostatic binding is reminiscent of the charge-charge interactions that allow DNA-scanning enzymes to slide freely over the DNA phosphate backbone (Blainey et al., 2006). If flexibly tethered charges on the Ndc80 complex are the basis for a slip clutch activity at the kinetochore, this would explain why their removal severely compromises kinetochore-microtubule coupling (Guimaraes et al., 2008; Miller et al., 2008).

The fact that kinetochores in all eukaryotes maintain load-bearing attachments to growing and shortening microtubule tips suggests that a common molecular mechanism underlies this behavior. We propose that ensembles of Ndc80 complexes acting through a biased diffusion mechanism supply the combination of plasticity and strength that allows kinetochores to hold on loosely to microtubule tips without letting go.

EXPERIMENTAL PROCEDURES

**Protein Expression and Purification**

The S. cerevisiae Ndc80 complex was expressed from two vectors encoding the Ndc80/Nuf2 and Spc24/Spc25 subcomplexes (Wei et al., 2005). For fluorescent studies, the Nuf2 gene was substituted with Nuf2-GFP from yeast strain MY107-5d (Shigemori et al., 2006). For the human Ndc80/Hec1 complex, a polyclastic vector encoding all four subunits was cloned (see Supplemental Experimental Procedures). All complexes included a C-terminal His6 tag on Spc24 for affinity purification and bead binding. Purification was carried out as described previously except buffers that are more amenable to the Ndc80 complex were used (Asbury et al., 2006; Franck et al., 2007; Gestaut et al., 2008). The buffer for purification over talon resin was 50 mM HEPES (pH 7.6) with 300 mM NaCl, 2.5 U ml⁻¹ benzozonase, and protease inhibitors (0.01 mg ml⁻¹ chymostatin, 0.01 mg ml⁻¹ aprotinin, 0.01 mg ml⁻¹ leupeptin, 0.01 mg ml⁻¹ pepstatin, 0.002 mg ml⁻¹ benzamidine, and 1 mM phenylmethylsulphonyl fluoride). For gel filtration, 50 mM HEPES (pH 7.6) with 200 mM NaCl was used.

**Bead Motility Assays**

Recombinant His₆-tagged Ndc80 complex was linked to 0.44 µm diameter streptavidin-coated polystyrene beads (Spherotech, Lake Forest, IL) using biotinylated His₆ antibody (QIAGEN, Valencia, CA) essentially as described in Asbury et al. (2006) and Franck et al. (2007). The amount of complex per bead was adjusted by incubating dilutions of 0.6–15 nM Ndc80 complex, prepared in BRB80 (80 mM PIPES, 1 mM MgCl₂, and 1 mM EGTA [pH 6.9]) plus 8 mg ml⁻¹ bovine serum albumin (BSA) and 1 mM DTT, with a fixed amount of beads (6.5 µM) at 4°C for 90 min. Flow chambers were created by sandwiching two strips of double-sided sticky tape between a KOH-washed coverslip and a standard glass slide and functionalized by coating with 5 mg ml⁻¹ biotinylated BSA (Vector Labs, Burlingame, CA), washing, and then incubating with 0.3 mg ml⁻¹ avidin DN (Vector Labs). Stable, biotinylated microtubule seeds were introduced and then washed with growth buffer (BRB80 plus 8 mg ml⁻¹ BSA with 1 mM GTP) before introduction of Ndc80-coated beads, which were diluted 8-fold into growth buffer supplemented with 1.5–1.9 mg ml⁻¹ tubulin, 1 mM DTT, and an oxygen scavenging system consisting of 250 µg ml⁻¹ glucose oxidase, 30 µg ml⁻¹ catalase, and 4.5 mg ml⁻¹ glucose. Assays were performed at 22°C.

**TIRF Microscopy Assay**

Flow cells for TIRF experiments were prepared as described in Gestaut et al. (2008). After rinsing with ddH₂O, the coverslip was incubated with “rigor” kinase-scented in BRB20 (20 mM PIPES, 1 mM MgCl₂, and 1 mM EGTA [pH 6.9]) plus 200 mM KCl and 8 mg ml⁻¹ BSA for 5 min. Taxol-stabilized Alexa-647-labeled microtubule seeds were introduced and then washed with growth buffer (BRB80 plus 8 mg ml⁻¹ BSA with 1 mM GTP) before introduction of Ndc80-coated beads, which were diluted 8-fold into growth buffer supplemented with 1.5–1.9 mg ml⁻¹ tubulin, 1 mM DTT, and an oxygen scavenging system consisting of 250 µg ml⁻¹ glucose oxidase, 30 µg ml⁻¹ catalase, 25 mM glucose and 5 mM DTT, and the reported amount of K⁺ was then introduced.

For experiments with dynamic microtubules, Alexa-647-labeled GMPCPP-stabilized seeds were bound and washed with growth buffer (BRB80 plus 8 mg ml⁻¹ BSA and 1 mM GTP). Extensions were assembled in growth buffer plus 2 mg ml⁻¹ tubulin (1% Alexa-647 labeled) and oxygen scavenger. After ~5 min, the buffer was exchanged with BRB20 plus 1 mM GTP, 2 mg ml⁻¹ tubulin (1% Alexa-647 labeled), and oxygen scavenger. Then, 1 nM GFP-labeled Ndc80 complex was introduced in a buffer lacking both free tubulin and GTP (BRB20 plus oxygen scavenger) to induce depolymerization. For antibody-induced oligomerization, 525 nM Ndc80 complex was preincubated with 250 nM biotin-conjugated His₆ antibody (QIAGEN, Valencia, CA) for ~60 min. For all experiments, the Ndc80 complex was passed through a 0.22 µm filter after dilution into the final buffer.

**Instrumentation, Data Collection, and Analysis**

Our optical force clamp has been described previously (Asbury et al., 2006; Franck et al., 2007). Bead-trap separation was sampled at 40 kHz, and stage position was updated at 50 Hz to maintain the desired load. Position data were
decimated to 200 Hz before storing to disk. Records of bead position versus time were analyzed using custom software written in Igor Pro (Wavemetrics, Lake Oswego, OR). Periods of microtubule growth and shortening were identified in the records by eye. Catastrophes were scored only when a bead moved in the direction of filament shortening by an amount that clearly exceeded the level of random noise (typically 5–10 nm rms during force clamping). Rescues were scored when an episode of rapid disassembly was clearly followed by slow growth, in assays without load, mean-squared displacements versus time for individual traces were averaged to estimate the diffusion coefficient during microtubule assembly.

TIRF movies were analyzed using software developed in Labview (National Instruments, Austin, TX) for tracking the position and brightness of individual GFP spots. Association rates were estimated by dividing the total number of events by the total microtubule length, the total observation time, and the concentration of free complex in solution. Matlab software (Mathworks, Natick, MA) was developed to calculate the average GFP intensity per pixel as a function of distance from disassembling MT tips. Only intervals when the tips appeared to accumulate more fluorescence than the lattice for > 4 s were included in the averages.

Computer Simulations of the Biased Diffusion Mechanism
Simulations of the biased diffusion model were performed in Igor Pro using standard iterative Monte Carlo methods. Briefly, for each iteration, the rate constants for “inward” and “outward” movement, $k_i$ and $k_o$, were first calculated based on the current state of the system, as described by Hill (1985). A random number between 0 and 1 was then generated and compared with these rates to determine whether the coupler moved during the iteration. The time interval corresponding to each iteration was kept very short (typically < 5 μs) to avoid missing events. Shortening the interval further had no effect on the simulations, confirming that the fraction of missed events was negligible. We also verified that the program recapitulates the steady-state distributions shown in Hill’s paper (i.e., Figure 3) when matching parameters are used.

SUPPLEMENTAL DATA
The Supplemental Data include Supplemental Discussion and Experimental Procedures, one table, five figures, and six movies and can be found with this article online at http://www.cell.com/supplemental/S0092-8674(09)00023-3.

ACKNOWLEDGMENTS
We thank Stephen Harrison and Jennifer DeLuca for plasmids, Anthony Azevedo for conducting early experiments with the 2NCC:2S construct, Ryan Lemke for technical assistance, Jason Stumpff for helpful comments on the manuscript, and all members of the Seattle Mitosis Club for valuable discussion. This work was supported by an NIH traineeship (T32 GM07270 awarded to A.F.P.), an NSF IGERT traineeship (DGE-0504573 to A.D.F.), a Searle Scholar Award (06-L-111 to C.L.A.), a Packard Fellowship for Science and Engineering (2006-30521 to C.L.A.), and by grants R01GM40506, R01GM79373, and R01GM69429 from the National Institute of General Medical Sciences (to T.N.D., C.L.A., and L.W., respectively).

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REFERENCES


SUPPLEMENTAL DATA

The Ndc80 kinetochore complex uses biased diffusion to couple chromosomes to dynamic microtubule tips

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SUPPLEMENTAL DISCUSSION

Uncertainty in the number of complexes required for a load-bearing coupler

We estimated the uncertainty in our calculation of the number of Ndc80 complexes required for load-bearing tip attachment by combining the uncertainties arising from three potential sources of error. First, we used BCA assays (Pierce, Thermo Fisher Scientific, Rockford, IL) to measure the concentrations of our Ndc80 preparations. Based on many replicate measurements, these assays are repeatable to within ± 6% in our hands. The manufacturer additionally reports protein-to-protein variation of ± 15%. Assuming uncorrelated errors, these can be combined (i.e., added in quadrature; Taylor, 1982) to yield a total uncertainty due to protein quantification of ± 16%. The second potential error source is the counting assay used to measure the concentrations of our preparations of anti-His beads. Based on repeated counts, the uncertainty of this measurement is ± 11%. The third source arises from Poisson statistics, which govern the variability in the number of complexes bound to a small area on the bead surface when the binding density is low. In the present case, this source will contribute ± 27% (i.e., $N^{-1/2}$, where $N = 14$ is the expected number of bound complexes based on the
average density). Combining all three sources (by adding them in quadrature) yields a total uncertainty of ± 33% or, equivalently, an estimate of 14 ± 5 complexes.

We considered the possibility that Ndc80 complexes might form clusters on the bead surface large enough to significantly increase the number interacting with the filament above our estimate (i.e. clusters > 20 to 30 complexes). However, several observations indicate that such clustering was unlikely. First, beads incubated with ≤ 2 nM GFP-tagged Ndc80 were uniformly decorated and did not exhibit the large bead-to-bead variability in brightness that would occur if the complexes bound in large clusters (Figure S4A and B). Second, their fluorescence increased linearly with the concentration of the complex, which demonstrates an absence of the binding cooperativity that would be expected if clustering were occurring (Figure S4C and D). Third, gel filtration experiments carried out in the same buffer that was used for bead binding showed no evidence for large clusters. The complex elutes as a clean peak with an apparent Stoke’s radius close to the predicted size for individual heterotetramers (Figure S1C). Thus the complex does not appear to cluster, either in solution or on the bead surface, under the conditions used here for bead binding.

**Estimation of kinetochore-microtubule forces *in vivo***

The most straightforward way to estimate the range of forces transmitted to kinetochores during normal movement of bi-oriented chromosomes is to measure the amount of stretching between sister kinetochores and then multiply by the elasticity of the chromatin that links them together. Nicklas (1988) used this method, after determining chromatin elasticity with a calibrated microneedle, to estimate a force of 7 pN per microtubule in meiotic grasshopper spermatocytes. Essentially the same calculation can be performed for budding yeast, yielding a very similar force estimate. Pearson et al.
(2001) showed that the pericentric chromatin linking sister kinetochores is extended to between 0.08 and 0.15 \( \mu \text{m kb}^{-1} \), depending on whether stretching of the lacO GFP marker contributes. A reasonable estimate for the elasticity of this pericentric chromatin is the elasticity measured for chromatin fibers \textit{in vitro}, which varies between 5 and 50 pN \( \text{kb} \mu \text{m}^{-1} \) depending on histone density (Brower-Toland et al., 2002). Multiplying these values gives the range of forces, 0.4 to 8 pN.

\textbf{Predictions of the biased diffusion mechanism}

Although Hill imagined a sleeve-like coupler completely encircling the microtubule, the physical underpinnings of his model can apply more generally to any array of microtubule binding elements that are linked together. The mechanism he proposed is a form of biased diffusion where the kinetochore undergoes thermally driven movement in a corrugated free energy landscape, as shown schematically in Figure S5A and B. The speed of diffusion depends on the height of the corrugations, which arise because movement along the filament requires breaking some bonds. Moving from one lattice site to another (e.g., from i to ii in Figure S5A and B) results in no net change in energy because the broken bonds can re-form. Thus, lattice-bound couplers in the absence of external load will undergo an unbiased random walk. However, if the coupler begins to move past the tip of the filament (i.e., from ii to iii, where it hangs partially off the end) some binding elements will no longer be able to reach the filament. Thus the free energy increases (in proportion to the number of broken bonds that are unable to re-form), which presents a barrier that biases the diffusion and inhibits detachment. Additionally, the corrugations in the landscape become smaller as the coupler begins to move off the tip because fewer bonds mediate the linkage, so fewer must be broken for further movement. This effect locally increases the rate of diffusion and, as explained in
the following paragraph, allows couplers that are nearly immobile on the lattice to track effectively with shortening tips. Tip tracking requires the diffusion of the coupler to be fast enough and biased enough at the tip to outrun disassembly.

Hill's model predicts several changes in behavior as the number of binding elements changes. The most obvious difference is that couplers with fewer elements are predicted to detach more frequently, owing to their reduced binding energy. Couplers with fewer elements will also diffuse more quickly on the lattice, because fewer bonds must be broken to allow movement. During microtubule shortening, another interesting difference is predicted. Small highly mobile couplers will tend to stay ahead of the disassembling tip by moving randomly on the lattice and only occasionally ‘bouncing’ off the tip (Figure 7A). Conversely, larger less mobile couplers will track closely with the tip, even if they are so large that their diffusion on the lattice is negligibly slow (Figure 7B). More unidirectional tracking occurs in these cases because the mobility is low on the lattice but increases when the tip begins to disassemble out from under the coupler (see above). This in turn promotes lattice-directed movement and formation of new bonds, resulting in a steady state where the rate of new bond formation is balanced by the loss due to disassembly.

SUPPLEMENTAL MATERIALS AND METHODS

Details of expression plasmid construction and protein purification

For expression of the *S. cerevisiae* Ndc80 complex, we used two dicistronic vectors encoding the Ndc80/Nuf2 and Spc24/Spc25 sub-complexes that have been described previously (Wei et al., 2005). To create a fluorescent-tagged version of the complex, the Nuf2 gene was excised from one of these vectors and substituted with Nuf2-GFP, which
was derived through PCR from yeast strain MSY107-5d (Shimogawa et al., 2006) with addition of a Spe1 restriction site at the 5’ end of the 3’ primer. The internal Kpn1 restriction site and Spe1 restriction site were used to ligate the Nuf2-GFP PCR product into the dicistronic vector. For the full-length human Hec1 complex, we created a new polycistronic vector encoding all four subunits. cDNA clones for human Spc25, Spc24 and Hec1 were purchased from ATCC (Manassas, VA), and a clone for human Nuf2 was kindly provided by Jennifer De Luca (Colorado State University). Human Spc25, Spc24, Hec1 and Nuf2 were cloned into cassettes 1, 2, 3, and 4 of the pST39 vector (Tan, 2001) using restriction sites Xba1 & Apa1, EcoR1 & HindIII, EcoRV & Kpn1, and BspE1 & Mlu1, respectively. In addition, a Pac1 site (encoding leucine-isoleucine-asparagine) was added to the 3’ end of Spc25 for future cloning. All versions of the complex included a C-terminal His6-tag on Spc24 to allow affinity purification and bead binding.

Vectors were transformed into E. coli BL21 (Rosetta; Novagen, Madison, WI) and protein expression and purification were carried out as previously described (Asbury et al., 2006; Franck et al., 2007; Gestaut et al., 2008). Cells containing the target plasmid(s) were grown at 37˚C until they reached a density of 30 Klett units, induced overnight (~12 hours) at 23˚C with the addition of 0.2 mM IPTG, then pelleted. Fresh pellets were lysed using a French press in 50 mM HEPES (pH 7.6) containing 300 mM NaCl, 2.5 U mL⁻¹ benzonase, and protease inhibitors (0.01 mg mL⁻¹ chymostatin, 0.01 mg mL⁻¹ aprotinin, 0.01 mg mL⁻¹ leupeptin, 0.01 mg mL⁻¹ pepstatin, 0.002 mg mL⁻¹ benzamidine and 1 mM phenylmethylsulfonyl fluoride) and clarified by centrifugation. Protein was purified by affinity chromatography using TALON resin as described by the manufacturer (BD Biosciences). Peak elutions were concentrated using an Amicon Ultra 50 kD MWCO centrifugal filter (Millipore, Billerica, MA) and then subjected to gel filtration chromatography on an SDX200 16/60 (GE Healthcare, Piscataway, NJ) equilibrated
with 50 mM HEPES (pH 7.6) containing 200 mM NaCl. Glycerol was added to a final concentration of 10% and aliquots were snap frozen and stored at -80˚C.

SUPPLEMENTAL REFERENCES


**Table S1.** Outcomes during force clamp experiments with yeast or human Ndc80 complex, under 0.5 to 2.5 pN of continuous tension.

<table>
<thead>
<tr>
<th></th>
<th>yeast Ndc80 complex</th>
<th>human Ndc80/Hecl complex</th>
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<tbody>
<tr>
<td>A</td>
<td>total events observed</td>
<td>392</td>
</tr>
<tr>
<td>B</td>
<td>events with discernable assembly-coupled movement only</td>
<td>250 (64%)(^1)</td>
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<tr>
<td>C</td>
<td>events with both assembly- and disassembly-coupled movement</td>
<td>64 (16%)(^2)</td>
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<td>D</td>
<td>events with discernable disassembly-coupled movement only</td>
<td>78 (20%)(^2)</td>
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<td>E</td>
<td>total events with discernable assembly-coupled movement (B+C)</td>
<td>314</td>
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<tr>
<td>F</td>
<td>'catastrophes' during assembly-coupled movement</td>
<td>64 (20%)(^2)</td>
</tr>
<tr>
<td>G</td>
<td>detachments during assembly-coupled movement</td>
<td>226 (72%)(^2)</td>
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<tr>
<td>H</td>
<td>other interruptions to assembly-coupled movement</td>
<td>24 (8%)(^2)</td>
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<td>I</td>
<td>total events with discernable disassembly-coupled movement (C+D)</td>
<td>142</td>
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<tr>
<td>J</td>
<td>'rescues' during disassembly-coupled movement</td>
<td>4 (3%)(^3)</td>
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<tr>
<td>K</td>
<td>detachments during disassembly-coupled movement</td>
<td>83 (58%)(^3)</td>
</tr>
<tr>
<td>L</td>
<td>other interruptions to disassembly-coupled movement</td>
<td>55 (39%)(^3)</td>
</tr>
</tbody>
</table>

\(^1\) expressed as a percentage of total events (row A)
\(^2\) expressed as a percentage of total events with discernable assembly-coupled movement (row E)
\(^3\) expressed as a percentage of total events with discernable disassembly-coupled movement (row I)
Figure S1. Purification of the Ndc80 complex

(A) Coomassie stained SDS-PAGE (12% acrylamide) showing the components of the S. cerevisiae Ndc80 complex, the GFP-tagged S. cerevisiae Ndc80 complex, the ‘headless’ 2NCC:2S complex and the human Ndc80/Hec1 complex.

(B) Anti-GFP Western blot showing the GFP-tagged Nuf2 subunit of the S. cerevisiae Ndc80 complex. As negative and positive controls, untagged Ndc80 complex and GFP-tagged Dam1 complex (Gestaut et al., 2008) were run in adjacent lanes.

(C) Gel filtration profile for the yeast Ndc80 complex in our standard purification buffer (50 mM HEPES pH 7.6 plus 300 mM NaCl; red trace) and in the buffer used for binding the complex to anti-His beads (BRB80; solid black trace). For comparison, the elution profile for a mixture of standard proteins in purification buffer is also shown (ferritin, Stoke’s radius of 61 angstroms; aldolase, 48 angstroms; RNAse, 18 angstroms; dotted black trace, offset vertically for clarity).
Figure S2. The human Ndc80/Hec1 complex couples force to dynamic microtubule tips

(A) Beads coated with the human Ndc80/Hec1 complex form persistent load-bearing attachments to growing and shortening microtubule tips, similar to attachments formed by the yeast Ndc80 complex. Records show bead position versus time during continuous application of tensile load, as in Figure 2A.

(B – D) Histograms of distance moved during assembly (N = 46 records), distance moved during disassembly (N = 25), and total attached time (including both assembly and disassembly phases; N = 50) for tip-attached beads moving under 0.5 - 2.5 pN of tension. Dotted vertical lines indicate the average for each distribution. Uncertainties represent s.e.m.
Figure S3. The brightness of diffusing Ndc80-GFP spots is unaffected by ionic strength
Distributions of initial brightness values for Ndc80-GFP particles diffusing on taxol-stabilized microtubules in 30 mM K⁺ (black), 60 mM K⁺ (red) or 90 mM K⁺ (blue) buffer. Each histogram was fit by a Gaussian function (dotted lines) to determine the mean brightness values indicated. Uncertainties represent s.d. For clarity, the 60 mM K⁺ and 90 mM K⁺ distributions are offset vertically by 100 and 200 counts, respectively.
Figure S4. Quantification of GFP-tagged Ndc80 complex bound to anti-His beads
(A) Epifluorescence images of anti-His beads after incubation with 2 nM GFP-tagged Ndc80 complex in our standard binding buffer (BRB80 plus 8 mg mL⁻¹ BSA and 1 mM DTT).
(B) Brightness distributions for beads incubated with 2 nM GFP-tagged Ndc80 complex in our standard binding buffer (black histogram) or in the buffer used for purification of the complex (50 mM HEPES pH 7.6 plus 300 mM NaCl; red histogram).
(C and D) Mean brightness for beads incubated with GFP-tagged Ndc80 in our standard binding buffer (black symbols) or in purification buffer (red symbols) plotted against the concentration of the complex. Uncertainties represent s.d. The bead concentration was 5.6 pM for all incubations.
Figure S5. The biased diffusion model

(A) Schematic picture of a coupler composed of five microtubule binding elements attached to a microtubule at three different locations, two on the lattice (denoted i and ii) and one on the tip in a position where two of the elements are no longer bound (iii).

(B) Free energy landscape for a five-element, biased diffusion-based coupler. The spacing of sites is denoted by \( l \). The gradual increase in free energy on the right arises from the bond energies, \( w \), that must be overcome to move the coupler past the filament tip. The corrugations in the landscape occur because movement along the filament requires breaking and reforming some bonds. Their heights, \( 5b, 4b, ..., b \), decrease as the coupler begins to move past the tip. Red circles (i, ii, and iii) denote the landscape positions corresponding to the couplers shown in (A).
SUPPLEMENTAL FIGURE LEGENDS

Figure S1. Purification of the Ndc80 complex

(A) Coomassie stained SDS-PAGE (12% acrylamide) showing the components of the *S. cerevisiae* Ndc80 complex, the GFP-tagged *S. cerevisiae* Ndc80 complex, the ‘headless’ 2NCC:2S complex and the human Ndc80/Hec1 complex.

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(C and D) Mean brightness for beads incubated with GFP-tagged Ndc80 in our standard binding buffer (black symbols) or in purification buffer (red symbols) plotted against the concentration of the complex. Uncertainties represent s.d. The bead concentration was 5.6 pM for all incubations.
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**SUPPLEMENTAL VIDEO LEGENDS**

**Video 1. An Ndc80-coated bead diffuses on the microtubule lattice then undergoes disassembly-driven movement**

Microtubule disassembly drives movement of an Ndc80-coated bead, shown here in the absence of optical trap-applied force. The bead is initially attached at the growing microtubule tip, but subsequently undergoes one-dimensional diffusion along the lattice. Following catastrophe the filament disassembles back to the diffusing bead, producing sustained bead movement until reaching the coverslip-anchored stabilized seed. The contrast-enhanced movie corresponds to the event in Figure 1A, which was recorded at 30 frames s$^{-1}$ and is displayed at ×20 speed. Beads were incubated with 15 nM S. *cerevisiae* Ndc80 complex. Scale bar, 5 µm.
**Video 2. The Ndc80 complex couples force to a growing and shortening microtubule tip**

Movement of an Ndc80-coated bead is coupled to microtubule growth and shortening while tension is continuously applied with an optical trapping-based force clamp. The tip-attached bead is held stationary in the optical trap and as the microtubule lengthens, the piezo stage moves leftward under feedback control to maintain a constant amount of tension (here, 1.3 pN). When the microtubule switches to shortening, rightward stage movement is used to similarly maintain tension. The approximate location of the coverslip-anchored microtubule seed is slightly below the coverslip-stuck bead. The movie was filmed at 30 frames s⁻¹ and is displayed at ×20 speed. Beads were incubated with approximately 2 nM *S. cerevisiae* Ndc80 complex. Scale bar, 5 µm.

**Video 3. The human Ndc80/Hec1 complex forms load-bearing attachments with a dynamic microtubule tip**

Beads coated with human Hec1 complex remain attached to growing and shortening microtubule tips in the presence of optical trap-applied tension (here, 1.1 pN), demonstrating that the ability to form a dynamic linker is conserved across species. The movie was filmed at 30 frames s⁻¹ and is displayed at ×60 speed. Hec1 complex incubation concentration, 400 pM. Scale bar, 5 µm.

**Video 4. Ndc80-microtubule interactions are affected by ionic strength**

Diffusion of individual Ndc80-GFP complexes on taxol-stabilized microtubules in motility buffer containing 30, 60 or 90 mM K⁺. In each movie, the red fluorescent microtubule is
oriented horizontally, and single Ndc80 complexes (green spots) undergo one-dimensional diffusion along this axis. Fewer particles are found diffusing on microtubules under higher concentrations of buffer K+, and their mean residence time is shorter and speed of diffusion is faster. Each movie was individually collected at 10 frames s⁻¹ using total internal fluorescence microscopy, and is displayed at ×2 speed. *S. cerevisiae* Ndc80 complex-GFP concentration, 20 pM. Scale bar, 3µm.

**Video 5. Lattice diffusion of Ndc80 complex is biased at disassembling microtubule tips**

A dynamic red fluorescent microtubule was grown from a coverslip-anchored seed (on right side of video), then induced to depolymerize by simultaneously flowing out the free tubulin while introducing Ndc80-GFP at single molecule concentration (here, 1 nM). Following introduction of complex, the shortening tip moves rightward, biasing the movement of lattice-bound Ndc80 toward the microtubule seed. The movie corresponds to the upper kymograph in Figure 6A. It was collected at 10 frames s⁻¹ using total internal fluorescence microscopy and is displayed at ×10 speed. Scale bar, 5 µm.

**Video 5. Ensembles of Ndc80 complex track unidirectionally with shortening microtubule tips**

Following antibody-induced oligomerization, particles of Ndc80-GFP move coincident with the depolymerizing tip. The movie corresponds to the lower kymograph in Figure 6A. It was collected at 10 frames s⁻¹ using total internal fluorescence microscopy and is displayed at ×5 speed. Scale bar, 5 µm.