A kinesin-13 mutant catalytically depolymerizes microtubules in ADP

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The kinesin-13 motor protein family members drive the removal of tubulin from microtubules (MTs) to promote MT turnover. A point mutation of the kinesin-13 family member mitotic centromere-associated kinesin/Kif2C (E491A) isolates the tubulin-removal conformation of the motor, and appears distinct from all previously described kinesin-13 conformations derived from nucleotide analogues. The E491A mutant removes tubulin dimers from stabilized MTs stoichiometrically in adenosine triphosphate (ATP) but is unable to efficiently release from detached tubulin dimers to recycle catalytically. Only in adenosine diphosphate (ADP) can the mutant catalytically remove tubulin dimers from stabilized MTs because the affinity of the mutant for detached tubulin dimers in ADP is low relative to lattice-bound tubulin. Thus, the motor can regenerate for further cycles of disassembly. Using the mutant, we show that release of tubulin by kinesin-13 motors occurs at the transition state for ATP hydrolysis, which illustrates a significant divergence in their coupling to ATP turnover relative to motile kinesins.

Introduction

Microtubules (MTs) spontaneously change length by the assembly and disassembly of tubulin dimers (Mitchison and Kirschner, 1984), and members of the kinesin-13 family promote MT turnover or dynamics by driving the removal of tubulin from MTs (Hunter et al., 2003). Highly dynamic MTs are a hallmark of both rapidly dividing and metastatic cells (Jordan and Wilson, 1998; Goncalves et al., 2001). The kinesin-13 family member mitotic centromere-associated kinesin (MCAK) is the most potent MT-depolymerizing ATPase so far identified, and is often up-regulated during tumorigenesis (Nakamura et al., 2007; Shimo et al., 2008). Yet, the role of ATP turnover in the disassembly of MTs by MCAK remains unclear (Ogawa et al., 2004). Previous structural studies have proposed that ATP hydrolysis is coupled to tubulin protofilament bending, and that tubulin release from MCAK would be coupled to phosphate release (Desai et al., 1999; Ogawa et al., 2004; Shipley et al., 2004; Helenius et al., 2006). Instead, using mutational analysis, we define a reaction scheme for MCAK that is related to that of motile kinesins but is distinct in the relationship between the nucleotide states and reaction coordinates.

Kinesin and myosin superfamily motor proteins use conserved structural elements to effect ATP hydrolysis and respond to different nucleotide states (Sablin et al., 1996). Residues in the regions named Switch I (consensus sequence: NxxSSR) and Switch II (DxxGxE) form a network of hydrogen bonds with the nucleotide, Mg2+, and each other. This network senses the presence or absence of the γ-phosphate and triggers both ATP hydrolysis and allosteric changes in the motor structure (Naber et al., 2003; Nitta et al., 2004; Hirose et al., 2006). Switch II is directly connected to the MT-binding site in the kinesin superfamily (Woehlke et al., 1997; Kikkawa et al., 2001). Recent cryoelectron microscopic studies suggest that alterations in these structures and in their contacts with tubulin among different nucleotide states (ADP, no nucleotide, and adenylyl-imidodiphosphate [AMPPNP]) may explain the changes in MT affinity between different nucleotide states (Hirose et al., 2006).

The structural and enzymatic effects of mutating the conserved G and E residues of Switch II to alanine have been well characterized for myosin motors (Sasaki et al., 1998). The G-to-A mutant is presumed to conform to a pre-ATP–like state that precedes the recovery stroke of the myosin head, whereas the E-to-A mutant is interpreted to correspond to a transition state leading to ATP hydrolysis, which allows the recovery stroke (Suzuki et al., 1998). In kinesin-1, the corresponding mutations...
also appear to correspond to analogous states before and after the docking of the neck linker to the MT-bound motor head. Neck linker docking allows the unbound head to move to its next binding site toward the plus end of the MT. This docking is also promoted by AMPPNP, so in kinesin-1 as well, the E-to-A mutant appears to occupy a point on the ATP cycle that immediately precedes or accompanies ATP hydrolysis (Rice et al., 1999; Tomishige et al., 2006). Mutations in the Switch II domain drastically reduce ATP turnover by blocking the hydrolysis step. A detailed reaction scheme linking steps in the ATP hydrolysis cycle and motility for kinesin-1 has been described recently (Guydosh and Block, 2006).

Biochemical and microscopic analysis of a point mutant in the Switch II region of MCAK (E491A) establishes that binding and detachment of tubulin from the MT occurs before ATP hydrolysis. This model is confirmed by our observation that in the presence of ADP, E491A is capable of detaching tubulin dimers from stabilized MTs and then recycling back onto the lattice to release 18 moles of tubulin per mole of motor. This slow catalysis contrasts with the behavior of the E491A in ATP, in which the mutant is capable of detaching a tubulin dimer, but only on a mole-per-mole basis. The mutant, presumably because of its unique conformation, remains strongly bound to the detached tubulin dimer and is unable to recycle back onto lattice. Thus, in the high-ATP environment of living cells, exogenously expressed E491A disassembles MTs via tubulin sequestration without triggering the tubulin autoregulatory response. We have previously demonstrated that disassembly of MTs in living cells via overexpression of wild-type (wt) MCAK leads to dose-dependent inhibition of endogenous cellular tubulin expression (Ovechkina et al., 2002). In summary, our data demonstrate that tubulin detachment by wt MCAK is completed before ATP hydrolysis and is mimicked by the E491A mutant. Release of detached tubulin dimer by the motor is correlated with the attainment of the transition state for ATP hydrolysis. Thus, in wt MCAK, detachment of tubulin may trigger ATP hydrolysis. This feature differs from other kinesins in which the motor remains strongly bound to lattice during the attainment of the transition state for ATP hydrolysis (Asenjo et al., 2006). We speculate that the kinesin-13 family may represent an early branch in the evolution of kinesin subfamilies.

**Results and discussion**

We engineered two mutants that correspond to the conserved G and E residues of Switch II (G489A and E491A; Fig. S1, available at http://www.jcb.org/cgi/content/full/jcb.200805145/DC1). When we assayed them in our in vivo depolymerization assay (Ovechkina et al., 2002), we found that total tubulin levels remained high even in cells with few or no visible MTs (Fig. 1). This contrasts with wt MCAK overexpression, which increases free tubulin dimers and triggers autoregulated tubulin translational suppression that is measurable as a dose-dependent decrease in total tubulin fluorescence (Ovechkina et al., 2002). In contrast, tubulin released by E491 (and to a lesser extent G489A) does not trigger translational repression of tubulin. We hypothesized that these mutant proteins remained bound to tubulin after removing it from MTs, and sequestered it from normal tubulin translational regulation (Cleveland et al., 1981). This suggested that the Switch II mutants are impaired in the final step of catalytic MT disassembly: the release of the detached tubulin dimer.

Consistent with this hypothesis, purified full-length E491A bound tubulin in vitro with much higher affinity ($K_d = 230 \text{ nM}$) than wt MCAK ($K_d = 4.7 \text{ μM}$; Fig. 2). We found that E491A exhibited lower tubulin affinity in ADP-Al-F, an ATP hydrolysis transition state analogue (Fisher et al., 1995; Nitta et al., 2004), which indicates that tubulin release occurs in concert with ATP hydrolysis, not $P_i$ release, as has been previously suggested (Hunter et al., 2003). Although the E491A mutant has a very low basal ATPase rate, the ATPase is still sensitive to stimulation by MTs or unpolymerized tubulin (Fig. S2, available at http://www.jcb.org/cgi/content/full/jcb.200805145/DC1). The $K_{\text{1/2, tubulin}}$ (the tubulin concentration at which the ATPase rate is half-maximal, analogous to the $K_d$ for tubulin) for E491A is 39 nM tubulin, whereas the $K_{\text{1/2, tubulin}}$ for wt MCAK is 1.5 μM, which confirms that the mutant has a much tighter association with tubulin.

Surprisingly, purified recombinant E491A MCAK was able to depolymerize MTs in vitro in the presence of either ATP or ADP (Figs. 3 and S3 A, available at http://www.jcb.org/cgi/content/full/jcb.200805145/DC1). wt EGFP-MCAK depolymerized MTs catalytically in ATP, releasing ~0.05 tubulin dimers per MCAK per second (Fig. 3, A and B). Neither ADP nor the nucleotide analogues AMPPNP or ADP-Al-F were able to promote depolymerization by wt MCAK, even at a stoichiometry of 1:3 MCAK/tubulin (Fig. 3 B), which indicates that completion of the normal nucleotide hydrolysis cycle is required for depolymerization. In contrast, E491A released somewhat more tubulin in ADP or ADP-Al-F than in ATP, though at an apparent rate that was lower than for wt MCAK, releasing 0.002–0.003 dimers per MCAK per second. The pre-ATP state mimic G489A
failed to promote any depolymerization in vitro in ATP-containing buffer (unpublished data), and was not analyzed further. The rate of depolymerization by E491A in ATP appeared to decline as free tubulin reached a concentration approximately equal to that of the mutant MCAK (Fig. 3 B, broken line), which suggests a product-inhibition effect mediated by tighter binding to tubulin in ATP than in ADP.

We directly observed E491A-MCAK depolymerizing MTs in a reaction mix containing excess MTs with a total internal reflection fluorescence (TIRF) microscope (Fig. 3 C, Video 1, available at http://www.jcb.org/cgi/content/full/jcb.200805145/DC1). The assay was performed at different motor concentrations designed to mimic the following stoichiometries: 1:3 (the stoichiometry for E491A in Fig. 3 B), 1:1.7, and supersaturated (1.3:1) stoichiometry, in which all tubulin dimers are expected to be complexed with E491A. Linear depolymerization rates were measured during the first 60–180 s of depolymerization at three E491A concentrations while holding the tubulin (MT) concentration constant. The TIRF rate of depolymerization at three E491A concentrations while holding the tubulin (MT) concentration constant, the percentage of tubulin dimer bound to the motor in each nucleotide and standard deviation from the mean (SD) are indicated. All lanes, respectively, are run on the same gel but the order of the lanes in E491 A (bottom) are arranged to correspond to the nucleotide order of wt MCAK (top).

To verify that E491A depolymerization is catalytic in ADP, we monitored the turbidity of reactions in which MCAK was approximately equimolar with protofilament ends (Fig. 3 D). In the turbidity experiments, the stoichiometry was 1:117 motor/tubulin or 1.4 motors per protofilament end for both wt MCAK and E491A. At this stoichiometry, no depolymerization could be detected by E491A–ATP, although we cannot be sure there is no activity because removal of one equivalent or a small molar excess of tubulin would give too small a change in turbidity to detect reliably. In contrast, after subtraction of background from control reactions, ~18 moles of tubulin were released per mole of E491A in ADP over the course of the experiment, which indicates robust catalysis. At this stoichiometry, wt MCAK, a proven catalytic MT depolymerizer, released ~28 moles of tubulin per mole of motor. Because E491A's catalytic depolymerization occurs in the absence of ATP, we conclude that the energy source for MCAK’s depolymerization activity is the binding energy between MCAK and tubulin, and that ATP hydrolysis gates tubulin release in wt MCAK. This tubulin release step is inhibited by ATP in the E491A mutant.
not caused by cross-linking of depolymerized tubulin by dimeric MCAK. We conclude that full depolymerization requires the attainment of the transition state for ATP hydrolysis that lies beyond the AMPPNP state. Attainment of this state either forces the tubulin dimer off the protofilament end or alters its affinity with neighboring tubulin. Because monomeric deletion constructs can promote full depolymerization in ATP (Maney et al., 2001), we favor the latter mechanism through an allosteric effect between MCAK and tubulin, as the forceful separation of the tubulin dimers would seem to require dimeric MCAK, with its two tubulin binding sites.

Neither E491A nor wt MCAK bind tubulin dimers with high affinity in ADP-Al-F, which shows that this state and tubulin release occur later in the cycle and are distinct from the depolymerization step. ADP-Al-F has been seen to form a bipyramidal transition state in crystal structures of both myosin and kinesin Kif1A, so ADP-Al-F is expected to represent a transition state, not, as sometimes reported, the ADP-Pi state, of MCAK. We conclude that MCAK has two transition states along the reaction coordinate.

Significantly, AMPPNP blocked depolymerization by the mutant protein (Fig. 3 A), thus the AMPPNP-bound state and the E491A state in other nucleotides are not equivalent and represent two successive states within the hydrolysis cycle of MCAK (diagrammed in Fig. 4 D). Kinesin-1 in AMPPNP allows docking of the neck linker to the head domain, though transiently, compared with the more tightly docked ADP-Al-F state (Asenjo et al., 2006), which indicates that the AMPPNP state is early in the reaction scheme. We speculate that for MCAK, AMPPNP stabilizes the predepolymerization configuration even in the E491A mutant, but that ADP or ADP-Al-F enables the mutant to reach a configuration that is farther along the depolymerization reaction coordinate than the AMPPNP state (Fig. 4 D). wt MCAK is known to bind and accumulate at MT ends in AMPPNP, causing shortening of the MTs by fraying of the lattice into protofilament curls, but this does not result in tubulin detachment from the lattice (Desai et al., 1999). A monomeric deletion construct in AMPPNP also frays MTs into oligomeric protofilaments (Moores et al., 2006), which indicates that the effect seen in AMPPNP is not caused by cross-linking of depolymerized tubulin by dimeric MCAK. We conclude that full depolymerization requires the attainment of the transition state for ATP hydrolysis that lies beyond the AMPPNP state. Attainment of this state either forces the tubulin dimer off the protofilament end or alters its affinity with neighboring tubulin. Because monomeric deletion constructs can promote full depolymerization in ATP (Maney et al., 2001), we favor the latter mechanism through an allosteric effect between MCAK and tubulin, as the forceful separation of the tubulin dimers would seem to require dimeric MCAK, with its two tubulin binding sites.

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It is tempting to speculate that detachment of the tubulin dimer from MT lattice may, in fact, promote ATP hydrolysis. This may be the explanation for the increased catalytic efficiency of wt MCAK over E491A, which behaves more like a conventional catalyst that does not actively alter itself to participate in a reaction.

Although kinesin-13 and kinesin-1 share highly homologous motor domains, their very different interactions with MTs are accompanied by remarkable divergence in their coupling to the ATPase reaction cycle. To gain some insight into this evolutionary divergence, we compared the phylogenetic distance between the Switch I, Switch II, and P-loop domains of myosins, kinesins, and ancestral G proteins (Fig. 4F).

**Figure 4. A model of the MCAK depolymerization reaction.** (A) MTs supersaturated with E491A can be rapidly disassembled in ATP. (B) The lower the ratio of E491A to tubulin, the less effective the motor is in disassembling MTs in ATP. (C) E491A is capable of steady catalytic disassembly of MTs in ADP because the motor can release from tubulin and recycle back onto the MT lattice. (D) A model for MCAK-dependent disassembly of MTs. Our data suggests that MCAK can completely remove one tubulin dimer before ATP hydrolysis. M:D:Mt represents ADP–MCAK diffusion on MT, M:T:Mt* represents the first ATP–MCAK depolymerization transition state, M:T*:t represents the ATP hydrolysis transition state/low tubulin affinity transition state, and M:DPi + t represents ADP–Pi–MCAK dissociated from tubulin dimer. A postulated zero ATP state is also shown. This state exhibits high affinity for the MT lattice (Moore and Wordeman, 2004). [E] Phylogenetic distances based on the alignment of the Switch I, Switch II, and the P-loop of kinesins, myosins, and G proteins. Alignments are shown in Fig. S4 (available at http://www.jcb.org/cgi/content/full/jcb.200805145/DC1).

**Comparison of nucleotide cycles between motile motors and kinesin-13.**

<table>
<thead>
<tr>
<th>Nucleotide state</th>
<th>Kinesin-1 (motile)</th>
<th>Kinesin-13 (MCAK)</th>
<th>Myosin (actin)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATP</td>
<td>Strong</td>
<td>Strong</td>
<td>Strong</td>
</tr>
<tr>
<td>ATP*</td>
<td>Strong*</td>
<td>Weak</td>
<td>Weak**</td>
</tr>
<tr>
<td>ADP</td>
<td>Weak</td>
<td>Weak</td>
<td>Strong</td>
</tr>
</tbody>
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**F** Phylogenetic distance between Kinesins, Myosins and small G-proteins.
Materials and methods

In vivo depolymerization assay

MCAK activity was quantified in vivo as described previously (Ovechkin et al., 2002). In brief, EGFP-MCAK–transfected cells were fixed and stained with anti-tubulin antibody. Cells were selected such that the EGFP level was equivalent to the amount of wt EGFP-MCAK observed to be just sufficient to depolymerize essentially all MTs in the cell. Tubulin-staining ratios between transfected cells and untransfected cells were calculated ("tubulin ratio"). wt MCAK-transfected cells with fully depolymerized MTs have a tubulin ratio of ~0.5, whereas cells transfected with GFP alone have an MT ratio of 1. Mutant EGFP-MCAK is expected to give a tubulin ratio greater than in wt cells if depolymerization is impaired.

Protein purification

We purified mutant full-length EGFP-MCAK-6H proteins for in vitro studies from a baculovirus expression system as described previously (Maney et al., 1998), except that the pFastBac1 plasmid (Invitrogen) was used to construct recombinant baculoviruses. All MCAK concentrations are described as active sites, although the protein is believed to be dimeric (Maney et al., 1998).

Tubulin-binding assay

Recombinant EGFP-MCAK was incubated in buffer containing 1.6 mM of the indicated nucleotide for 3.5 h on ice, then incubated at 700 nM in 0.5 ml of the same buffer. Bound proteins were eluted with 500 mM imidazole-HCl and run on SDS-NuPAGE gels (Invitrogen). Coomassie-stained gels were dried, scanned, and measured using ImageJ, and the percent tubulin polymerization and pelleting from GMPCPP solution. The final cycle was pelleting assay, except with 14 nM MCAK and 100 μM ADP or ATP. The turbidity was measured every 6 s in a 1-cm path-length cuvette at 340 nm. After 3,000 s, 10 mM CaCl₂ was added to depolymerize remaining MTs and establish a baseline reading. Loss of MT polymer (Y) was fit to a one-phase exponential decay Y = (Y₀)(1-e⁻ᵏᵗ) using GraphPad Prism (GraphPad Software). This case, Y₀ = initial tubulin polymer concentration of 1.6 M [K = rate constant and X = time]. Each experimental condition was performed identically 3–5 times on different days and averaged to obtain the error.

Online supplemental material

Fig. S1 shows a space-filling model showing the position of key residues in Switch I and Switch II. Fig. S2 shows tubulin-stimulated and MT-stimulated ATPase of recombinant EGFP-MCAK and E491A mutant. Fig. S3 shows in vitro depolymerization in various nucleotide states and MCAK-tubulin stoichiometries. Table S1 shows alignments used to build the distance tree in Fig. 4. Video 1 shows TIRF microscopy of MTs depolymerizing in the presence of two different E491I stoichiometries. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200805145/DC1.

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References


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