MCAK associates with the tips of polymerizing microtubules

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CAK is a member of the kinesin-13 family of microtubule (MT)-depolymerizing kinesins. We show that the potent MT depolymerizer MCAK tracks (treadmills) with the tips of polymerizing MTs in living cells. Tip tracking of MCAK is inhibited by phosphorylation and is dependent on the extreme COOH-terminal tail of MCAK. Tip tracking is not essential for MCAK’s MT-depolymerizing activity. We propose that tip tracking is a mechanism by which MCAK is preferentially localized to regions of the cell that modulate the plus ends of MTs.

Introduction

MCAK is a member of the kinesin-13 (Lawrence, 2004) family of microtubule (MT)-depolymerizing kinesins. Previous studies have implicated MCAK in the regulation of MT dynamics during interphase and mitosis and in ensuring proper chromosome segregation during mitosis (for review see Moore and Wordeman, 2004b). MCAK is a potent depolymerase of stabilized MTs (Desai et al., 1999; Hunter et al., 2003) and increases the catastrophe frequency of dynamic MTs in vitro 10-fold (Newton et al., 2004). MCAK is the most potent cellular MT depolymerizer that has yet been identified (Tournebize et al., 2000; Ogawa et al., 2004). MCAK displays extremely high affinity for both plus and minus MT ends, yet its localization to MT ends in fixed cells is intermittent at best. For these reasons we imaged live cells transfected with levels of GFP or RFP-MCAK that do not appreciably disturb the overall MT polymer level. When imaged in this manner, MCAK is observed to track with polymerizing MT tips.

A number of proteins exhibit the unusual property of targeting and treadmilling (tracking) along polymerizing MT ends. The two most well characterized of these are members of the Clip-170 and EB1 families (for review see Carvalho et al., 2003). Thus far, all known treadmilling tip proteins in mammalian cells exhibit MT-stabilizing activity. For this reason, it is remarkable that the potent MT depolymerase, MCAK, tracks the tips of polymerizing MTs. MCAK’s tip tracking is dependent on phosphorylation, which may regulate the association of MCAK with the MT lattice or with recruiting proteins, thus influencing the ability to localize to MT plus ends. The closely related kinesin-13 family member Kif2A does not tip track in mammalian cells and its activity appears to dominate at MT minus ends (Gaetz and Kapoor, 2004; Ganem and Compton, 2004; Rogers et al., 2004) by virtue of its dynein-dependent preferential localization to centrosomes. Hence, tip tracking may represent a mechanism to preferentially localize MCAK’s activity to regions of the cell, such as the distal face of the centromere, that interact with MT plus ends.

Results and discussion

When fluorescent MCAK is expressed in live cultured cells at levels that do not significantly alter MT polymer levels, MCAK can be detected on MT lattice and in obvious densities on MT tips (Fig. 1 A). Time-lapse imaging shows tips polymerizing toward the cell edge in a manner similar to that which has been previously reported for GFP-EB1 (Fig. 1 B; Video 1, available at http://www.jcb.org/cgi/content/full/jcb.200411089/DC1). In living HeLa cells the MCAK tip densities are coincident with GFP-EB1 (Fig. 1 C). Tracking of MCAK to MT tips is not dependent on a functional motor domain, as it can be clearly seen in cells transfected with GFP-ML-MCAK (Fig. 2 B; Video 3), a construct in which the 50-kD motor domain has been removed and the NH2-terminal domain of MCAK is fused directly to the COOH-terminal domain (Maney et al., 1998). Tracking tips of GFP-ML-MCAK are consistently longer than those seen with GFP-wt-MCAK (Fig. 2, A’ and B’), suggesting that the motor may influence the off-rate from the lattice. Another mutant of MCAK that contains three point mutations in the motor domain rendering the
protein inactive with respect to depolymerizing activity (GFP-hypir-MCAK) also exhibits tip tracking (Fig. 2 C, Table I; Video 4). Tip localization and tracking is seen in both interphase and mitotic cells. This is a striking observation because MCAK has been previously shown to be a potent depolymerizer of MTs (Desai et al., 1999; Hunter et al., 2003). Therefore, the presence of MCAK on the tips of polymerizing MTs in interphase and mitotic cells suggests that MCAK’s depolymerizing activity is transiently inhibited on tips.

To determine whether other known MT depolymerizing proteins are present on MT tips, we assayed two splice variants of another kinesin-13 family member: Kif2Aα and Kif2Aβ. Both proteins exhibited significant depolymerizing activity when overexpressed (Fig. S1 A, available at http://www.jcb.org/cgi/content/full/jcb.200411089/DC1). However, these proteins were not found tracking on MT tips (Table I; Fig. S1 B and Video 8). This is unexpected because the putative Drosophila melanogaster Kif2A orthologue, Klpl0A, does track on polymerizing tips, whereas the putative MCAK orthologue, Klpl59C, does not (Mennella et al., 2005). Furthermore, neither GFP-stathmin (Marklund et al., 1996) nor GFP-ch-TOG (Charrasse et al., 1998) was found to be tracking on polymerizing MT tips. This suggests that MCAK is the major MT depolymerizer in mammalian cells that tracks on the tips of polymerizing MTs. GFP-APC decorated MT tips but did not exhibit the same distribution or dynamics as those decorated with MCAK or EB1. Instead, tips exhibited aperiodic bending previously described in Dayanandan et al. (2003).

We mapped the domains outside of the core motor domain required for tip tracking, and the results are shown in Figure 1.

**Figure 1.** MCAK exhibits tip tracking behavior in living cells. (A) GFP-MCAK can be detected at the ends of MTs in an interphase HeLa cell. (B) Kymograph of tips traveling from the centrosomal region to the edge of the cell. (C) RFP-MCAK roughly colocalizes with GFP-EB1 in live HeLa cells cotransfected with both constructs.

**Figure 2.** Inactive MCAK tip tracks in interphase and mitotic cells. (A) The edge of a cell transfected with GFP-MCAK. (A’) Three-dimensional reconstruction of 30 frames of MCAK tip tracks. Successive moving tips are visible (black arrows). (B) GFP-ML-MCAK tracks at the edge of an interphase cell. (B’) Reconstruction of 30 frames of motorless MCAK show uniform lines (rather than successive tips) because the tips are consistently longer than is seen with GFP-MCAK. (C) Mitotic cell transfected with GFP-hypir-MCAK. Tip tracking is evident in astral MTs (white arrows).
Table II. Neither the core motor domain nor the neck + motor domain was capable of tip tracking. The neck + motor domain is a significantly more potent MT depolymerizer than full-length wtMCAK because it is missing the COOH-terminal negative regulatory domain (Moore and Wordeman, 2004a). Tip tracking is exquisitely sensitive to the presence of the COOH terminus. Loss of 9 amino acids from the COOH terminus eliminated tip tracking (MCAK-Q710), and loss of even 2–5 amino acids significantly impaired tip tracking (Table II).

Because phosphorylation has been invoked as a mechanism for inactivating MCAK’s depolymerizing activity (Andrews et al., 2004; Lan et al., 2004; Ohl et al., 2004), we tested whether tip-associated MCAK was inhibited by phosphorylation. We found that our more active (Fig. S1 A) phospho-mutant of MCAK (GFP-AAAAA-MCAK) exhibited robust tip tracking (Table III, Fig. 3 A; Video 5, available at http://www.jcb.org/cgi/content/full/jcb.200411089/DC1). In this mutant all of the sites known to be phosphorylated by Aurora B kinase have been mutated to alanine (Andrews et al., 2004). In contrast, a less active (Fig. 3 A) phospho-mimic mutant of MCAK (GFP-EEEEEMCAK) showed no tip tracking (Table III, Fig. 3 A; Video 6). The fact that active GFP-AAAAA-MCAK tip tracks suggests that phosphorylation is not the means by which MCAK is inhibited on tips of polymerizing MTs.

Although the neck domain is required for maximally efficient MT depolymerization activity the S196E mutant of MCAK, which mimics phosphorylation within the neck domain, had no effect on tip tracking. This is consistent with our data suggesting that tip tracking is not essential for maximal MT depolymerizing activity. However, a single substitution of S92E, NH2-terminal to the neck domain, significantly reduced the length and intensity of the tracking tips (Table III). This suggests that the Aurora B sites NH2-terminal to the neck are most critical for the regulation of tip tracking. It is significant that although the Aurora B consensus phosphorylation sites appear to be conserved between all the KinI/kinesin-13 family members (Andrews et al., 2004), the protein sequence surrounding the conserved phosphorylation sites are quite divergent between MCAK and Kif2Aα/β. This implicates the NH2 terminus of MCAK, particularly the region surrounding the Aurora B/Ipl1 phosphorylation sites in associating preferentially with MT ends.

Because the tip-tracking behavior of MCAK was negatively regulated by phosphorylation, we observed the behavior of endogenous MCAK in the presence of the kinase inhibitor roscovitine. CHO cells were incubated in either DMSO or 10 μM roscovitine in DMSO for 5 h, and then were fixed and labeled for endogenous MCAK and MTs (Fig. 3 B, top and bottom, respectively). After 5 h of roscovitine treatment endogenous MCAK exhibited significantly greater association with MT tips. The level of roscovitine used is capable of inhibiting a number of kinases (Meijer et al., 1997). Because the assay was performed on interphase cells, the inhibited kinase is not Aurora B. However, we do find that inhibition of Aurora B with small molecule inhibitors or a kinase-dead mutant promotes greater association of MCAK with MT tips in mitotic cells (unpublished data). The redistribution of endogenous MCAK was striking (Fig. 3 B, bottom). We also performed time-lapse imaging of GFP-MCAK transfected cells in order to confirm that MCAK was tracking with tips in the presence of roscovitine (Fig. 3 C; Video 7). To summarize, MCAK’s ability to tip track is negatively dependent on phosphorylation of the NH2-terminal domain and positively mediated by the extreme COOH-terminal tail.

Presently, it is not known how proteins track with MT ends. Both copolymerization with tubulin and preferential affinity for MT end structures have been proposed. MCAK has been previously shown to possess an affinity for both tubulin and MT ends. To distinguish between these two mechanisms, we compared tip-tracking (wt-unphosphorylated) and nontip-tracking (wt-phosphorylated and Q710) MCAK proteins for their ability to be competed off MT lattice by free tubulin. If tubulin copolymerization were the mechanism by which MCAK

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Table II. Tip tracking of MCAK is dependent on the extreme COOH terminus

| Construct                  | Tip tracking | Diagram of deletion |
|----------------------------|-------------|---------------------|
| GFP-MCAK, RFP-MCAK         | Yes         | ![Diagram of deletion](image1) |
| GFP-MCAK-NH2-terminus      | No          | ![Diagram of deletion](image2) |
| GFP-MCAK-Neck+Motor        | No          | ![Diagram of deletion](image3) |
| GFP-MCAK-Motor             | No          | ![Diagram of deletion](image4) |
| GFP-MCAK-M1-A603           | No          | ![Diagram of deletion](image5) |
| GFP-MCAK-M1-Q650           | No          | ![Diagram of deletion](image6) |
| GFP-MCAK-M1-Q710\^[a]\     | No          | ![Diagram of deletion](image7) |
| GFP-MCAK-M1-K715           | Yes\^[b]\   | ![Diagram of deletion](image8) |
| GFP-MCAK-M1-H717           | Yes\^[b]\   | ![Diagram of deletion](image9) |

\^[a]\ Referred to in the text as “MCAK-Q710.”
\^[b]\ Weak.

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preferentially associates with MT ends, then one might expect versions of MCAK that tip track would interact with free tubulin to a greater degree than versions that do not. We have previously shown that tubulin inhibits lattice association of MCAK (Moore and Wordeman, 2004a). We compared the extent to which unphosphorylated and Ipl1/Sli15-phosphorylated wild-type MCAK bind to assembled MTs, and the extent to which that binding is limited by excess tubulin monomer. The former tip tracks, whereas the latter presumably does not. Under conditions in which 100% of nonphosphorylated MCAK is bound to lattice, only about two thirds of Ipl1-phosphorylated MCAK is found in the pellet (Fig. 4 A, lanes 1). However, both unphosphorylated and phosphorylated MCAK can be competed off lattice to a similar extent by addition of excess free tubulin: 19% in the former case, and 13% (9% out of 68%) of the bound fraction in the latter case (Fig. 4 A, lanes 2). Therefore phosphorylation, which is expected to reduce tip tracking, changes the affinity of MCAK for lattice but not for free tubulin.

We performed the same experiment using MCAK-Q710, which does not tip track but exhibits a higher apparent affinity for MT polymer than wt-MCAK (Moore and Wordeman, 2004a). We found that this mutant form can be competed away from MT lattice to a much greater extent than either wt-MCAK or phosphorylated wt-MCAK (Fig. 4 B, lanes 2 and 3). However, phosphorylation of MCAK-Q710 reduced lattice binding by approximately the same level as did phosphorylation of wt-MCAK. Curiously, phosphorylated MCAK-Q710 was only

| Construct         | Tips | Tip tracking |
|-------------------|------|--------------|
| GFP-MCAK, RFP-MCAK | Yes  | Yes          |
| GFP-AAAAA-MCAK    | Yes  | Yes          |
| GFP-EEEEEE-MCAK   | No   | No           |
| GFP-S196E-MCAK    | Yes  | Yes          |
| GFP-S92E-MCAK     | Yes* | Yes*         |

*Tip length is much shorter than wt-MCAK.

Figure 3. Tip tracking of MCAK protein is dependent on phosphorylation. (A) GFP-AAAAA-MCAK binds to MT tips (left), whereas GFP-EEEEEE-MCAK is not found on tips (right). See also Videos 4 and 5 (available at http://www.jcb.org/cgi/content/full/jcb.200411089/DC1). (B) Two fields of CHO cells labeled for endogenous MCAK (B) and MTs (B'). The top field of CHO cells are control cells. The bottom field of CHO cells were cultured for 5 h in 10 μM roscovitine and then fixed. Increased association of endogenous MCAK with distal ends of MTs is evident (B, bottom; arrows). (C) MCAK protein tracks on tips (arrows) in a HeLa cell transfected with RFP-MCAK and cultured subsequently in 10 μM roscovitine (Video 6).

Figure 4. Tip tracking is positively correlated with apparent lattice affinity rather than tubulin affinity. (A) Wild-type MCAK binds MT lattice in the absence of ATP (left, lane 1). Excess free tubulin dimer can compete off 13–15% of lattice-associated wt-MCAK (left, lanes 2 and 3). Phosphorylation of MCAK significantly decreases the apparent affinity of wt-MCAK for stabilized MTs (right, lane 1). Excess free tubulin competes off a further 2–9% of lattice-bound phosphorylated MCAK (right lanes 2 and 3). (B) MCAK-Q710 binds MT lattice in the absence of ATP (left, lane 1). Excess tubulin dimer can compete off 30–45% of lattice-bound MCAK-Q710 (left, lanes 2 and 3). Phosphorylation of MCAK-Q710 decreases the apparent affinity of MCAK-Q710 for MT lattice to a similar extent as is seen with wt-MCAK (right, lane 1). Excess free tubulin competes off a further 9–11% of lattice-bound phosphorylated MCAK-Q710 (right lanes 2 and 3).
competed away from lattice by free tubulin to approximately the same extent as wt-MCAK (8% of 63% = 13% of the bound fraction). These experiments show that tip tracking is correlated positively with lattice affinity but negatively with free tubulin affinity, contrary to the prediction of a simple model for tip tracking by copolymerization.

Instead, tip tracking may simply depend on the ability of MCAK to see a higher affinity-binding site at the end of the MT relative to the lattice. As the MT polymerizes, this higher affinity site becomes a lower affinity site, leading to increased loss of MCAK from the lattice and the characteristic comet tail appearance. Because tip tracking does not depend on the motor domain we propose that this binding site is distinctly different from the very high affinity binding of MCAK to deformable tubulin dimers at both ends of the MT in AMP-PNP described in structural and kinetic studies (Desai et al., 1999; Moores et al., 2002; Hunter et al., 2003; Moore and Wordeman, 2004a; Ogawa et al., 2004). We have previously hypothesized that the COOH-terminal tail domain “sees” a site at the end of the MT that relieves the inhibition of the tail and promotes lattice binding (Moore and Wordeman, 2004a). This site could be the exposed β-tubulin subunit end. In that case, tip tracking would be plus end dependent and might be antagonized by free tubulin (Fig. 4 B). If tip tracking could be reconstituted in vitro this hypothesis could be tested.

A remaining question is why a MT tip that is enriched for bound MCAK would polymerize at all. MCAK’s MT depolymerizing activity must be inhibited. We propose two models, which are not mutually exclusive, to explain the inhibition of MCAK on MT tips (Fig. 5). In the first case, the MT tip clearly associates with both MT stabilizers (such as EB1) and MT destabilizers (MCAK). MT stabilizers, such as EB1, may be capable of successfully competing with MCAK on MT tips by preventing MCAK from reaching a critical concentration at the end of the MT. We coexpressed EB1 and MCAK in live cells and found that excess EB1 can indeed antagonize the MT-depolymerizing activity of MCAK in living cells as long as the excess MCAK levels are relatively low (Fig. 5 A). We call
cDNA plasmids were a gift of N. Santama (University of Cyprus, Nicosia, no. D12644) and Kif2A (INSERM, Montpellier, France). Kif2A H530A, R534A, and K537A mutations in pEGFPC1. EGFP-mmStathmin ries, Inc.). EGFP-ML-cgMCAK (pYOY154) was prepared from GFP-ML-pYOY71 (Ovechkina et al., 2002) into pEGFPC1 (CLONTECH Laboratories). EGFP-cgMCAK (pYOY152) was prepared by pasting a fragment of

Constructs

Materials and methods

Egfp-MCAK phospho-mutants versus wt-MCAK. Kif2A expression and purification were performed as previously described (Maney et al., 1999). lpl1p and Sli15p GST fusion plasmids were gifts of Dr. Sue Biggins (Fred Hutchinson Cancer Research Center, Seattle, WA). Bovine brain tubulin was acquired from Cytoskeleton, Inc. 50 μM tubulin was added to BRB80 [80 mM Pipes, pH 6.8, 1 mM EGTA, and 1 mM MgCl2] with 4 mM MgCl2, 1.8 mM GMP-CP (a nonhydrolyzable form of GTP), and 5% DMSO and incubated for 20 min at 37°C. After the incubation, MTs were diluted 1:12.5 in 37°C BRB80. Phosphorylation of MCAK was performed by incubating 50 nM of active MCAK with 1 μM lpl1p and 0.5 μM Sli15p in kinase buffer (50 mM Tris-HCl, pH 7.5, 0.1 mM EGTA, and 5 mM MgCl2) with 1 mM DTT and 75 mM KCl for 1 h. Supernatants and pellets were assayed for the presence of tubulin and motor on Coomassie-stained SDS-polyacrylamide gels (Invitrogen). Bands were quantified using NIH Image 1.62.

Online supplemental material

Cell transfection and immunofluorescence

Online supplemental material

Fig. S1: relative MT-depolymerizing activity of Kif2A splice variants and MCAK expression and purification were performed as previously described (Maney et al., 1999). lpl1p and Sli15p GST fusion plasmids were gifts of Dr. Sue Biggins (Fred Hutchinson Cancer Research Center, Seattle, WA). Bovine brain tubulin was acquired from Cytoskeleton, Inc. 50 μM tubulin was added to BRB80 [80 mM Pipes, pH 6.8, 1 mM EGTA, and 1 mM MgCl2] with 4 mM MgCl2, 1.8 mM GMP-CP (a nonhydrolyzable form of GTP), and 5% DMSO and incubated for 20 min at 37°C. After the incubation, MTs were diluted 1:12.5 in 37°C BRB80. Phosphorylation of MCAK was performed by incubating 50 nM of active MCAK with 1 μM lpl1p and 0.5 μM Sli15p in kinase buffer (50 mM Tris-HCl, pH 7.5, 0.1 mM EGTA, and 5 mM MgCl2) with 1 mM DTT and 75 mM KCl for 1 h. Supernatants and pellets were assayed for the presence of tubulin and motor on Coomassie-stained SDS-polyacrylamide gels (Invitrogen). Bands were quantified using NIH Image 1.62.
ylation of MCAK that is in the supernatant versus the pellet in these experiments. Video 1: interphase HeLa cell and GFP-MCAK. Collected at 1 frame per 12 s. Video 2: interphase HeLa cell and GFP-MCAK. Collected at 1 frame per 12 s. Video 3: interphase HeLa cell and GFP-MCAK. Collected at 1 frame per 12 s. Video 4: mitotic HeLa cell and GFP-hypir-MCAK. Collected at 1 frame per 20 s. Video 5: interphase HeLa cell and GFP-MCAK. Collected at 1 frame per 5 s. Video 6: interphase HeLa cell and GFP-EEEEE-MCAK. Collected at 1 frame per 5 s. Video 7: interphase HeLa cell, GFP-Kif2Aβ. 1 frame per 5 s. Online supplemental material available at http://www.jcb.org/cgi/content/full/jcb.200411089/DC1.

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