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The spindle assembly function of Caenorhabditis elegans katanin does not require microtubule-severing activity

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ABSTRACT Katanin is a heterodimeric microtubule-severing protein that is conserved among eukaryotes. Loss-of-function mutations in the Caenorhabditis elegans katanin catalytic subunit, MEI-1, cause specific defects in female meiotic spindles. To determine the relationship between katanin’s microtubule-severing activity and its role in meiotic spindle formation, we analyzed the MEI-1(A338S) mutant. Unlike wild-type MEI-1, which mediated disassembly of microtubule arrays in Xenopus fibroblasts, MEI-1(A338S) had no effect on fibroblast microtubules, indicating a lack of microtubule-severing activity. In C. elegans, MEI-1(A338S) mediated assembly of extremely long bipolar meiotic spindles. In contrast, a nonsense mutation in MEI-1 caused assembly of meiotic spindles without any poles as assayed by localization of the spindle-pole protein, ASPM-1. These results indicated that katanin protein, but not katanin’s microtubule-severing activity, is required for assembly of acentriolar meiotic spindle poles. To understand the nonsevering activities of katanin, we characterized the N-terminal domain of the katanin catalytic subunit. The N-terminal domain was necessary and sufficient for binding to the katanin regulatory subunit. The katanin regulatory subunit in turn caused a dramatic change in the microtubule-binding properties of the N-terminal domain of the catalytic subunit. This unique bipartite microtubule-binding structure may mediate the spindle-pole assembly activity of katanin during female meiosis.

INTRODUCTION Katanin was first purified from sea urchin eggs based on its ATP-dependent, microtubule-severing activity (McNally and Vale, 1993). Katanin is a heterodimer consisting of a catalytic subunit and a regulatory subunit (McNally and Vale, 1993; Hartman et al., 1998). The katanin catalytic subunit is a member of a family of closely related AAA ATPases which also includes the microtubule-disassembly proteins spastin and fidgetin. These proteins play roles in a number of diverse cellular processes, including cell division, cilia biogenesis, deflagellation, and neurogenesis (Roll-Mecak and McNally, 2010).

In Caenorhabditis elegans, the katanin catalytic subunit, MEI-1, is essential for meiotic spindle function (Mains et al., 1990) but is destroyed by a ubiquitin-dependent pathway before assembly of the first mitotic spindle (Pintard et al., 2003). The MEI-1(P99L) mutant protein does not bind its cognate ubiquitin ligase (Pintard et al., 2003), and therefore persists on the first mitotic spindle (Clark-Maguire and Mains, 1994a), resulting in embryonic lethality due to defective mitosis (Mains et al., 1990). Genetic screens for suppressors of this embryonic lethality have identified a large number of intragenic suppressor mutations in MEI-1 and extragenic suppressor mutations in MEI-2 (Mains et al., 1990; Clandinin and Mains, 1993). These suppressor mutations all reduce the ectopic activity of MEI-1(P99L) during mitosis but otherwise fall into two classes. Mutants in the first class, including the null allele, mei-1(p99l;ct101), are 100% maternal-effect embryonic lethal as homozygotes. mei-1(null) or mei-2(null) meiotic spindles consist of a disorganized mass of microtubules that lack the discrete parallel bundles and focused spindle poles characteristic of wild-type meiotic spindles (Mains et al., 1990; Yang et al., 2003). Because of
these spindle defects, the chromosomes in a mei-1(null) or mei-2(null) spindle never congress or exhibit organized anaphase chromosome separation (Yang et al., 2003). Deep invaginations of the cortex result in expulsion into polar bodies of random subsets of chromosomes, and none of these embryos develop to hatching (Mains et al., 1990; Yang et al., 2003).

A second class of mutation includes alleles like mei-1(P99L;P235S) that, when homozygous, result in low rates of embryonic lethality, suggesting that meiotic spindles are functional (Mains et al., 1990). Of these homozygous viable suppressors of mei-1(P99L), the mei-2(A237T) mutant has been characterized in the greatest detail. In contrast with mei-1(null) spindles, mei-2(A237T) meiotic spindles are bipolar and mediate anaphase chromosome segregation (McNally et al., 2006). mei-2(A237T) spindles are longer than wild-type spindles during metaphase, suggesting that reduced microtubule-severing activity leads to longer spindles. After activation of the APC ubiquitin ligase, wild-type meiotic spindles first shorten in the pole–pole axis as microtubule density increases. This early shortening occurs normally in mei-2(A237T) spindles. Wild-type spindles then rotate to a perpendicular orientation at the cortex and continue shortening as microtubule density decreases. mei-2(A237T) spindles do not rotate or shorten during this phase, but microtubule density still decreases (McNally et al., 2006). These results revealed a role for katanin in regulating spindle length during metaphase and in the second phase of spindle shortening. The effect on postrotation spindle shortening suggested that MEI-1/MEI-2 might sever microtubules at spindle poles, thus causing the spindle to transform into a cylindrical shape during late anaphase.

There are two distinct models that could explain the phenotypic differences between mei-1(null) or mei-2(null) spindles and mei-2(A237T) spindles. In the first model, katanin’s only in vivo function is to sever microtubules, and different mutations cause quantitative reductions in microtubule-severing activity. Homozygous viable mutations like mei-2(A237T) reduce severing activity to a level that does not interfere with mitotic spindle function but allows assembly of a long, bipolar, meiotic spindle that does not exhibit postrotation spindle shortening. Further reduction in severing activity leads to a failure in the assembly of meiotic spindle poles and parallel microtubule bundles. It is difficult to explain why microtubule-severing activity would be absolutely required for the assembly of spindle poles and parallel microtubule bundles. In the second model, some in vivo functions of katanin, like regulation of metaphase spindle length, are mediated by microtubule-severing activity, whereas other functions, such as spindle pole assembly, are mediated by different biochemical activities of katanin.

We reasoned that a separation-of-function allele of mei-1 that eliminated microtubule-severing activity but retained spindle pole–forming activity would mediate assembly of the longest bipolar spindles. In search of such a separation-of-function allele, we measured meiotic spindle lengths in a collection of homozygous viable mei-1 mutants. We found that the mutant with the longest meiotic spindles also lacked microtubule-severing activity. This is the first demonstration of an essential nonsevering activity for katanin.

RESULTS

In search of a separation-of-function allele, we first assayed the MEI-1 protein expression levels of a series of homozygous-viable mei-1 point mutants (Mains et al., 1990; Clandinin and Mains, 1993) by immunoblotting synchronized young adult worms. If MEI-1/MEI-2 had multiple activities, then any mutation causing a decrease in protein concentration, like mei-2(A237T) (Srayko et al., 2000), would reduce all of these activities and thus could not be used to separate microtubule-severing activity from other functions. Of 11 homozygous viable mei-1 mutants, only 1, P99L;sb10, had reduced MEI-1 protein levels (Supplemental Figure S1). Of the 10 mutants with wild-type protein levels, 6 were crossed with green fluorescent protein (GFP):tubulin-expressing worms to allow an in vivo analysis of meiotic spindle lengths. Live imaging revealed that all meiotic spindles were bipolar in embryos of worms homozygous for these six viable alleles. Wild-type meiotic spindles maintain a steady-state metaphase spindle length for 6 min before initiating APC-dependdent shortening (Yang et al., 2003, 2005). Therefore, pole-to-pole spindle length measurements were made of metaphase I and metaphase II spindles from short time-lapse sequences that exhibited a constant spindle length for at least 1 min to ensure that measurements of shortening spindles were not included (Figure 1 and Table 1). A subset of spindles was monitored until they initiated spindle shortening (Figure 1A) to ensure that embryos were viable. As has been reported previously, metaphase spindle length was decreased in mei-1(P99L) relative to mei-1(+). This observation is consistent with the interpretation that MEI-1(P99L) protein levels are increased during meiosis and that the resulting increase in microtubule-severing activity leads to a reduction in the length of spindle microtubules (Johnson et al., 2009). For each of the viable missense mutants that were characterized, both MI and MII spindle lengths were increased relative to those of mei-1(P99L) (Figure 1B and Table 1). For the alleles P99L;P235S and P99L;A338S, however, spindle lengths were increased to the greatest extent and were longer than those of the wild type. These results suggest that microtubule-severing activity might be decreased to some extent in each of these mutants and reduced to the greatest extent in P99L;P235S and P99L;A338S.

Ala-338, which is changed to serine in mei-1(P99L;A338S) is conserved not only among AAA ATPases but also in prokaryotic AAA+ enzymes like the Escherichia coli helicase, RuvB (Supplemental Figure S2A). Mutation of this alanine in RuvB results in a complete loss of function (Iwasaki et al., 2000). Ala-338 is within a β strand that has been referred to as “sensor I,” and modeling work in E. coli FtsH suggests that it positions the downstream asparagine that interacts with the γ-phosphate of ATP (Karata et al., 1999, 2001). Pro-235, which is changed to serine in the mei-1(P99L;P235S) mutant (Clark-Maguire and Mains, 1994b), is also conserved in both AAA and AAA+ ATPases (Supplemental Figure S2B) and lies within the Walker A motif or P loop. The adjacent prolines Pro-234 and Pro-235 form a sharp turn that positions the downstream lysine which is required for ATP binding (Babst et al., 1998; Scott et al., 2005a; Roll-Mecak and Vale, 2008). Thus either of these mutations has the potential to completely eliminate ATP-dependent, microtubule-severing activity.

To test MEI-1(P99L;P235S) and MEI-1(P99L;A338S) for microtubule-disassembly activity, GFP-tagged wild-type and mutant MEI-1 proteins were coexpressed with GST:MEI-2 in Xenopus fibroblasts and in Xenopus A6 cells, and the effect on interphase microtubule arrays was assayed by anti-tubulin immunofluorescence (Figure 2A). Expression of wild-type MEI-1 sometimes resulted in microtubules with two discrete ends visible (Figure 2A, left), and sometimes resulted in microtubules still emanating from the centrosome but reduced in number (Figure 2A, right). The average fluorescence intensity of anti-tubulin staining relative to that of adjacent untransfected cells was 50% (Figure 2B), indicating a 50% reduction in polymer mass. Similar results were obtained in a control experiment using cells expressing the human MEI-1 orthologue KATNA1 and the MEI-2 orthologue KATNB1 (previously referred to as p60 and p80 katanin). GFP fluorescence intensity was used to choose cells expressing the mutant forms of MEI-1 at the same or higher expression levels as wild

| mei-1 | mei-2 | mei-3 | mei-4 | mei-5 | mei-6 | mei-7 | mei-8 | mei-9 | mei-10 |
|------|------|------|------|------|------|------|------|------|-------|
| mei-1 | mei-2 | mei-3 | mei-4 | mei-5 | mei-6 | mei-7 | mei-8 | mei-9 | mei-10 |
| 1551 | 1551 | 1551 | 1551 | 1551 | 1551 | 1551 | 1551 | 1551 | 1551 |
Allele | Amino acid change | Phenotype | MI metaphase length (μm) | MII metaphase length (μm) \\
--- | --- | --- | --- | --- \\
Wild-type, mei-1+ | None | Wild-type | 8.31 ± 0.10 (n = 10) | 6.93 ± 0.09 (n = 10) \\
mei-1(ct46) | P99L | Dominant maternal effect lethal | 7.52 ± 0.06 (n = 12) | 6.37 ± 0.06 (n = 13) \\
mei-1(ct46, sb9) | P99L, G126S | Homozygous viable | 8.45 ± 0.09 (n = 19) | 7.27 ± 0.17 (n = 11) \\
mei-1(ct46, sb16) | P99L, A338S | Homozygous viable | 10.39 ± 0.16 (n = 16) | 9.00 ± 0.08 (n = 16) \\
mei-1(ct46, sb21) | N.D. | Homozygous viable | 8.66 ± 0.35 (n = 14) | 7.54 ± 0.20 (n = 13) \\
mei-1(ct46, sb22) | P99L, R128C | Homozygous viable | 8.96 ± 0.13 (n = 9) | 7.31 ± 0.10 (n = 14) \\
mei-1(ct46, sb23) | P99L, P360S | Homozygous viable | 8.47 ± 0.11 (n = 14) | 7.13 ± 0.10 (n = 10) \\
mei-1(ct46, ct103) | P99L, P235S | Homozygous viable | 9.69 ± 0.13 (n = 15) | 8.44 ± 0.17 (n = 12) \\
mei-1(ct46, ct101) | P99L, nonsense | Homozygous maternal effect lethal | No spindle poles (n = 50) \\
mei-1(b284) | P225L | Homozygous maternal effect lethal | No spindle poles (n = 18) \\
mei-1(ct81) | L231F | Homozygous maternal effect lethal | No spindle poles (n = 20) \\
mei-1(ct46, ct82) | P99L, G473D | Homozygous maternal effect lethal | No spindle poles (n = 9) \\
mei-1(ct46, ct84) | P99L, E308K | Homozygous maternal effect lethal | No spindle poles (n = 18) \\
mei-1(ct46, ct89) | P99L, R414K | Homozygous maternal effect lethal | No spindle poles (n = 20) \\

All other amino acid changes are from Clark-Maguire and Mains (1994b). The schematic drawing in Figure 1 shows how metaphase length was measured.

**TABLE 1:** Amino acid change for sb16 was determined by sequencing two independent PCR-amplifications of the mei-1 gene.
FIGURE 2: Expression of wild-type and mutant katanins in Xenopus tissue culture cells. Xenopus fibroblasts and A6 cells were cotransfected with GFP-tagged wild-type or mutant MEI-1 and with GST:MEI-2. Control cells were cotransfected with GFP-tagged human KATNA1 and with GST:KATNB1. Cells were fixed and immunostained with antibodies directed against α-tubulin, and images of high GFP expressers were captured. (A) Representative images of transfected fibroblasts. Expression of wild-type MEI-1 sometimes resulted in fragmented microtubules with both ends clearly visible (top left) but more frequently caused a reduction in the density of microtubules still emanating from the centrosome (top right) as did expression of the human homologue, KATNA1 (bottom right). Microtubule density was somewhat reduced by MEI-1(P99L,P235S). No reduction in microtubule density was seen with the homozygous viable allele, P99L;A338S, or with any of the maternal-effect lethal alleles L231F, P99L;G473D, or P225L. Bar = 25 μm.

(B) Graph of relative microtubule density in transfected fibroblasts and A6 cells. For every frame captured, the average α-tubulin pixel intensity in transfected cells was expressed as a percentage of the average α-tubulin pixel intensity in adjacent nontransfected cells. Columns represent the average of the percentages for wild-type and mutant MEI-1 and KATNA1 in Xenopus fibroblasts (light shading) and A6 cells (dark shading). N = the number of transfected cells analyzed; bars = SEM. Relative GFP fluorescence is the average GFP pixel intensity of transfected cells for each experiment divided by the average GFP pixel intensity of wild-type MEI-1–expressing cells. A number greater than one indicates that the GFP construct was expressed at a greater level than GFP-MEI-1(wild-type).
MEI-1's microtubule-disassembly activity is not required for bipolar spindle assembly and that microtubule-disassembly activity in transfected worms homozygous for each of five homoyzous maternal-effect lethal alleles of MEI-1 (L231F, P99L;G473D, P99L;E308K, P99L;R414K, ct101) exhibited no staining of ASPM-1 on the disorganized array of microtubules associated with the chromosomes (Figures 3 and 4). The ct101 allele is a nonsense mutant that produces no MEI-1 protein (Clark-Maguire and Mains, 1994b). L231F and G473D are mutations in highly conserved amino acids (Supplemental Figure S2, B and C; Clark-Maguire and Mains, 1994b) and have no detectable microtubule disassembly activity (Figure 2). These results show that MEI-1 protein, but not MEI-1's microtubule-dissevering activity, is required to assemble spindle poles.

We previously found that depletion of KLP-18, a potential microtubule cross-linker, suppressed the spindle translocation defect of mei-1(null) spindles (McNally et al., 2006), suggesting that a spindle composed of long, cross-linked microtubules cannot translocate to the cortex. Improved confocal imaging indeed suggested that spindle microtubules are cross-linked to cytoplasmic microtubules in mei-1(null) embryos (Figures 3 and 4). To test whether KLP-18 depletion can rescue the spindle pole defect of mei-1(null) spindles, we subjected mei-1(ct46;ct101); klp-18(RNAi) double-mutant embryos to anti-ASPM-1 staining. As previously reported (Wignall and Villeneuve, 2009), klp-18(RNAi) single-mutant embryos had a monopolar spindle with a single focus of ASPM-1 staining. Both mei-1(null) single-mutant embryos and mei-1(null); klp-18(RNAi) double-mutant embryos had no ASPM-1 staining (Figure 4). Thus MEI-1 protein, but not MEI-1's microtubule-dissevering activity, is directly required for spindle pole assembly.

Because mei-1(P99L;A338S) appeared to be a clean separation-of-function allele, we analyzed its effect on spindle rotation and on microtubule-severing activity in transfected worms. MEI-1 with the P99L;P235S mutations had reduced microtubule-disassembly activity (Supplemental Figure S3). Anti-GFP immunoblots revealed that all of the GFP:MEI-1 constructs were expressed as full-length fusion proteins (Supplemental Figure S3). The result that MEI-1(P99L;A338S) had no microtubule-disassembly activity in transfected Xenopus cells and that this mutant allowed assembly of bipolar spindles (Figure 1) suggested that microtubule-disassembly activity is not required for bipolar spindle assembly and that loss of some other activity of MEI-1 must be responsible for the strong defect in spindle assembly previously reported for a mei-1 null mutant and for mei-1(RNAi) (Mains et al., 1990; Yang et al., 2003).
postrotation spindle shortening. Rotation and postrotation spindle shortening were defective in the mei-2(A2377) mutant (McNally et al., 2006), but this mutant has a reduced amount of MEI-2 protein (Srayko et al., 2000). Time-lapse imaging of GFP:tubulin revealed that both rotation and postrotation spindle shortening occurred as in wild type in the mei-1–1(P99L;ct101) embryos with the exception that spindle length was somewhat increased (Figure 5A). This result indicated that microtubule-severing activity is not required during these spindle stages. On the basis of the mei-2(A2737) phenotype, we had previously postulated that microtubule severing at spindle poles during anaphase might mediate the apparent disappearance of microtubules from outside the chromosomes. This disappearance of microtubules is significant because it has been proposed that attachments between spindle poles and chromosomes are not present during anaphase in C. elegans female meiotic spindles (Dumont et al., 2010). To address this issue more carefully, we stained wild-type meiotic spindles with anti-ASPM-1 and found that ASPM-1–containing spindle poles are present outside the chromosomes throughout anaphase chromosome separation (Figure 5B). By both fixed immunofluorescence and improved live imaging, it is also clear that some microtubules are still present outside the chromosomes during anaphase (Figure 5, A and B). Thus the failure of mei-2(A2737) spindles to shorten and change in shape during anaphase is not due to a failure in severing spindle pole microtubules to place chromosomes on the end of the spindle but may reflect an altered spindle pole structure when the amount of MEI-2 protein is reduced.

What nonsevering activity of MEI-1 might mediate spindle pole assembly and ASPM-1 recruitment? Work on katanins and spastins from other species indicates that the monomeric AAA domain can form a transient hexameric ring on ATP binding and that the ATP-bound hexamer has a higher binding affinity for microtubules than do monomers (Hartman and Vale, 1999; Roll-Mecak and Vale, 2008). X-ray structures of monomers modeled into a hexamer (Scott et al., 2005; Roll-Mecak and Vale, 2008) suggest that the N-terminal non-AAA domains extend outward from the hexameric ring. The N-terminal domains of katanin catalytic subunits from sea urchin (Hartman and Vale, 1999) and Arabidopsis (Stoppin-Mellet et al., 2007) exhibit microtubule-binding activity suggesting that the hexameric ring might be able to cross-link six microtubules into a bundle that might lead to spindle pole formation. Furthermore, microtubule binding and bundling might be enhanced by interaction of hexameric rings with the regulatory subunits of katanin. Possible enhancement of microtubule binding by the regulatory subunit is suggested by the observations that the first 29 amino acids of the N-terminal domain of a human katanin catalytic subunit, KATNA1, are also required for binding to the regulatory subunit, KATNB1, and that KATNB1 enhances microtubule binding by KATNA1 (McNally et al., 2000).

In search of nonsevering activities of MEI-1, we first tested whether the N-terminal domain is necessary and sufficient for interaction with its regulatory subunit, MEI-2. Lynates from insect cells expressing full-length MEI-1, a deletion missing the N-terminal 29 amino acids or the N-terminal domain alone were mixed with an E. coli–expressed maltose binding protein (MBP) fusion to MEI-2 or with MBP alone. Complexes were purified by amylose affinity chromatography, and bound complexes were analyzed by SDS–PAGE and immunoblotting. As shown in Figure 6A, the N-terminal domain was both necessary and sufficient for binding to MEI-2. Because katanin N-terminal domains had not previously been tested for binding to their corresponding regulatory subunits, we also assayed for binding of the N-terminal domain of the human katanin catalytic subunit, KATNA1, to KATNB1, or to the previously uncharacterized homologue of KATNB1, c15orf29 (Roll-Mecak and McNally, 2010). Like the N-terminal domain of MEI-1, the N-terminal domain of KATNA1 was sufficient for binding both KATNB1 and c15orf29 (Figure 6B). Deletion of the N-terminal 29 amino acids of KATNA1 had only a minor effect on binding to KATNB1 but had a stronger effect on binding to c15orf29 (Figure 6B). These results indicate that a complex between the N-terminal domain of MEI-1 and MEI-2 might be the physiologically relevant microtubule-binding structure conferring MEI-1’s essential spindle pole assembly activity. Because KATB is required for KATA function in Tetrahymena ciliogenesis (Sharma et al., 2007) just as MEI-2 is absolutely required for MEI-1’s in vivo function (Mains et al., 1990; Srayko et al., 2000), the complex between the N-terminal domains of katanin catalytic subunits with their corresponding regulatory subunits might be generally significant.

| Genotype | n=  | Tubulin | ASPM | DAPI | Merge |
|----------|-----|---------|------|------|-------|
| wild-type | 14  | ![Tubulin](image1) | ![ASPM](image2) | ![DAPI](image3) | ![Merge](image4) |
| klp-18 (RNAi) | 12  | ![Tubulin](image5) | ![ASPM](image6) | ![DAPI](image7) | ![Merge](image8) |
| mei-1 (P99L; ct101) | 10  | ![Tubulin](image9) | ![ASPM](image10) | ![DAPI](image11) | ![Merge](image12) |
| mei-1 (P99L; ct101), klp-18 (RNAi) | 14  | ![Tubulin](image13) | ![ASPM](image14) | ![DAPI](image15) | ![Merge](image16) |

FIGURE 4: ASPM-1 is not present in meiotic spindles of klp-18(RNAi); mei-1(null) double mutants. Worms were grown overnight at 25°C, and embryos were fixed and stained with antibodies directed against ASPM-1 and α-tubulin. A bright ring of ASPM-1 is present at the single spindle pole in klp-18(RNAi) embryos, but no significant ASPM-1 staining occurs in mei-1(P99L;ct101) embryos or klp-18(RNAi), mei-1(P99L;ct101) double-mutant embryos. ct101 is a nonsense allele of mei-1 and behaves as a complete null (Mains et al., 1990). Each phenotype was observed in 100% of embryos of that genotype, and the number of spindles examined for each genotype is indicated. Bar = 5 μm.
Because complexes between katanin N-terminal domains with their regulatory subunit had not previously been characterized, we tested the microtubule-binding activity of these complexes in vitro and in vivo. The N-terminal domain of human KATNAL1 showed saturable binding to microtubules assembled from highly purified, microtubule-associated protein (MAP)-free porcine brain tubulin with an apparent $K_d$ of 4 μM (Figure 7). The MEI-2-like human c15orf29 also exhibited saturable binding to microtubules with an apparent $K_d$ of 2 μM. When these proteins were combined, the apparent $K_d$ for the N-terminal domain was modestly decreased to 0.8 μM, and the $K_d$ for c15orf29 decreased to 0.9 μM. The more dramatic effect was seen in the stoichiometry of binding at saturation. The N-terminal domain of KATNAL1 saturated with only 14% bound. The presence of c15orf29 increased the fraction of the N-terminal domain bound at saturation to 50%. These results indicate that the complex between the N-terminal domain and the regulatory subunit binds microtubules in a qualitatively different manner than does the N-terminal domain alone.

To investigate whether this microtubule binding is significant in vivo, a GFP fusion to the N-terminal domain of MEI-1 was co-expressed with MEI-2 in Xenopus fibroblasts. In the presence of MEI-2, the N-terminal domain of MEI-1 decorated a rare subset of microtubules (Figure 8). None of the full-length MEI-1 constructs assayed for microtubule-disassembly activity (Figure 2) showed any colocalization with perinuclear microtubules. These results indicate that the structure generated by dimerization of the MEI-1 N-terminal domain with MEI-2 can bind microtubules in vivo and that this binding is somehow inhibited in intact MEI-1/MEI-2 complexes. Exposure of the N-terminal MEI-1/MEI-2 complex may be regulated by the ATP hydrolysis cycle of the AAA domains or interactions with additional proteins in meiotic embryos.

**FIGURE 5:** Spindle rotation and postrotation spindle shortening in mei-1(P99L;A338S). (A) Time-lapse images of embryos expressing GFP:tubulin indicate that meiotic spindle rotation and postrotation shortening occurred in 7/7 mei-1(P99L, A338S) embryos and in 6/6 wild-type embryos. Bar = 3 μm. (B) Wild-type and mei-1(P99L, P235S) worms were grown overnight at 25°C, and embryos were fixed and stained with antibodies directed against ASPM-1 and α-tubulin. Wild-type embryos (22/22) and mutant embryos with anaphase meiotic spindles (10/10) had bright foci of anti-ASPM-1 staining at the spindle poles. A small amount of anti-tubulin staining was also observed at the poles in embryos with the brightest staining. Bar = 3 μm.

**DISCUSSION**

Complexes of wild-type MEI-1 and MEI-2 sever microtubules in vitro (McNally et al., 2006) and cause disassembly of interphase microtubule arrays in HeLa cells (Sranyo et al., 2000) and Xenopus fibroblasts. MEI-1(P99L; A338S) appears to lack microtubule-severing activity based on the following criteria: It has no detectable microtubule disassembly activity when expressed in Xenopus fibroblasts, and the mutation is in a residue conserved even in the distantly related bacterial ATPase, RuvB, where that residue is essential for function. Because our preparations of purified MEI-1(P99L;A338S) had lower protein concentration than the preparations of MEI-1(wild-type) that have robust in vitro microtubule-severing activity (McNally et al., 2006), we could not use in vitro severing assays to demonstrate a lack of activity for MEI-1(P99L;A338S). In contrast, we could readily find individual Xenopus cells expressing GFP-MEI-1(P99L;A338S) with higher GFP fluorescence intensity than that of cells expressing GFP-MEI-1(wild-type). No individual cells expressing GFP-MEI-1(P99L;A338S) showed any discernable signs of microtubule disassembly.

Meiotic spindles assembled by MEI-1(P99L; A338S) have metaphase lengths that are significantly longer than those assembled by MEI-1(P99L; P235S), a protein with greatly reduced microtubule-disassembly activity in fibroblasts. Because the expression level of MEI-1(P99L; A338S) protein and that of MEI-1(P99L; P235S) protein are similar to wild type, these results strongly suggest that microtubule-severing activity causes a reduction in metaphase spindle length. Because the nonsevering mutant, MEI-1(P99L; A338S), mediates assembly of spindles with two ASPM-1–positive poles but a lack of MEI-1 or MEI-2 protein causes assembly of spindles containing no ASPM-1 foci, our results indicate that a complex of MEI-1 and MEI-2 mediates spindle pole assembly through a nonsevering mechanism.
In other cell types, it has been proposed that spindle pole assembly requires the microtubule cross-linking activities of kinesin-14 (Goshima et al., 2005) or cytoplasmic dynein (Merdes et al., 2010). Because katanin is proposed to form a transient radial hexamer with N-terminal microtubule binding domains extending outward (Hartman and Vale, 1999; Roll-Mecak and McNally, 2010), it is possible that katanin promotes spindle pole formation by a microtubule cross-linking mechanism. Because mei-1(null) or mei-2(null) spindles lack poles, it is likely that a structure composed of both subunits mediates the essential function of katanin.

In this study, we showed for the first time that the N-terminal domain of MEI-1 is sufficient to bind MEI-2 and that the N-terminal domain of human KATNAL1 (one of three human MEI-1 orthologues) is sufficient to bind either of two human MEI-2 orthologues, KATNB1 and c15orf29. Both KATNB1 and c15orf29 copurify with KATNAL1 from human cells (Torres et al., 2009), but the relative concentrations and importance of different combinations of subunits are not known. The nuclear magnetic resonance structure of the N-terminal domain of mouse KATNAL1 revealed that it is a three-helix bundle called a microtubule interacting and transport (MIT) domain (Iwaya et al., 2010), a protein fold conserved in the N-terminal domains of VPS4 (Scott et al., 2005b) and spastin. Whereas the N-terminal domains of sea urchin (Hartman and Vale, 1999) and Arabidopsis (Stoppin-Mellet et al., 2007) katanin catalytic subunits exhibit saturable, low-affinity microtubule binding, qualitative binding assays suggested that the N-terminal domain of mouse KATNAL1 bound tubulin dimers or aggregates rather than microtubules (Iwaya et al., 2010). In this study, we found that the N-terminal domain of human KATNAL1 has saturable, low-affinity microtubule-binding activity like the equivalent domains of sea urchin and Arabidopsis katanins. Binding of the human KATNAL1 MIT domain to microtubules saturated with only 14% of the protein bound at saturation, however, possibly explaining the lack of binding reported for the mouse KATNAL1 MIT domain. The putative katanin regulatory subunit, c15orf29, bound microtubules by itself and saturated with 90% bound. Remarkably, an equimolar mixture of the KATNAL1 MIT domain and c15orf29 bound microtubules with a higher affinity than either protein alone and saturated at 50% bound. These results support a model in which binding of the katanin regulatory subunit to the catalytic subunit generates a completely new microtubule-binding structure. We suggest that this novel bipartite structure is the physiologically relevant microtubule-binding structure that may cross-link microtubules to allow spindle pole formation.
**MATERIALS AND METHODS**

**C. elegans strains**

Strains were cultured according to standard procedures (Brenner, 1974). Wild type refers to AZ244 (unc-119(ed3);rul57[pAZ147:pie-1/β-tubulin::GFP;unc-119(+)], which is derived from N2 (Prattis et al., 2001). Worms carrying the following temperature-sensitive alleles were maintained at 16°C as homozygous stocks: mei-1(ct46, sb9, sb10, sb16, sb17, sb21, sb22, sb23, ct103). Worms carrying the following nonconditional alleles were maintained as balanced heterozygous stocks: mei-1(ct81, ct82, ct84, ct89, b284). All mei-1-bearing chromosomes included an unc-13 or unc-29 mutation in cis to recognize the appropriate homozygous segregant. For live imaging of worms carrying temperature-sensitive mei-1 alleles, the mei-1 strains were crossed with AZ244, and progeny homozygous for mei-1, unc-13, and GFP-tubulin were isolated.

**Live imaging**

Adult hermaphrodite worms grown for 24 h at 25°C were anesthetized with tricaine/tetramisole (Kirby et al., 1990; McCarter et al., 1999) and gently mounted between a coverslip and a thin 3% agarose pad on a glass slide. Live imaging was primarily done on a Microphot SA (Nikon, Tokyo, Japan) microscope equipped with a 60× PlanApo 1.4 objective and a charge-coupled device (CCD; Qimagming Retiga Exi Fast 1394 camera). Excitation light from an HBO100 light source was attenuated with a heat- and UV-reflecting dichroic mirror (Chroma Technology, Brattleboro, VT) and a 25% transmission neutral density filter. A GFP long pass filter set (Omega Optical, Brattleboro, VT) was used. Excitation light from a U-RFL-T 100W Mercury burner power supply (Olympus) was controlled by a Sutter Lambda 10–3 controller and MetaMorph software. Spindle length was measured as the distance along the pole-to-pole axis.

**Immunostaining of C. elegans embryos**

Adult hermaphrodite worms were washed in a watch glass in 0.8% egg buffer and then transferred to a poly-L-lysine–coated Super-frost/Plus glass slide (Fisher Scientific, Pittsburgh, PA). Excess liquid was wicked away, a coverslip was gently applied, and the slides were submerged into liquid nitrogen for 10 min. Immunostaining was performed using standard freeze-fracture methods followed by –20°C methanol fixation as described previously (Tsou et al., 2002; DeBella et al., 2006). Primary antibodies were diluted in phosphate-buffered saline containing 0.05% Tween 20 (PBST) in the following ratios: anti-ASPM-1 (van der Voet et al., 2009a), 1:100; anti-tubulin (DM1alpha), 1:200. All secondary antibodies were diluted 1:200 in PBST. The secondary antibodies used were Alexa594 anti-rabbit immunoglobulin (Ig)G (Molecular Probes, Eugene, OR) and Alexa488 anti–mouse IgG. DAPI (4',6-diamidino-2-phenylindole) staining was used to visualize DNA. Stained embryos were imaged using an Olympus (Center Valley, PA) IX71 inverted microscope equipped with a 60× PlanApo N.A. 1.42 objective, an Olympus DSU spinning disk attachment, and a Hamamatsu (Bridgewater, NJ) Orca R2 deep cooled C10600–10B digital monochrome CCD camera. Excitation light from a U-RFL-T 100W Mercury burner power supply (Olympus) was controlled by a Sutter Lambda 10–3 controller and MetaMorph software.

**Plasmid constructions**

The coding sequence of each protein expressed was PCR-amplified to add restriction enzyme cleavage sites and cloned into one of the following vectors: pFastBac Dual (for baculovirus expression), pMAL-CR1 (for E. coli expression), pEGFP-C1 (Clontech; for expression in Xenopus cells). Point mutations in MEI-1 were generated by site-directed mutagenesis.

**Transfection of Xenopus cells**

Xenopus fibroblasts (provided by Vladmir Rodionov, University of Connecticut Health Center, Farmington, CT) and Xenopus A6 cells were grown on coverslips in 70% Leibovitz’s L15 Medium supplemented with 10% fetal bovine serum and penicillin/streptomycin at 27°C. Cells were transfected with Lipofectamine 2000 using equal concentrations of GFP-MEI-1 plasmids and a GST-MEI-2 plasmid constructed in pCDNA3. Thirty-six hours after transfection, coverslips were fixed in –20°C methanol and stained with anti-tubulin antibody (DM1alpha) and Alexa594 anti–mouse secondary antibody.

**FIGURE 8:** The N-terminal domain of MEI-1 binds microtubules in vivo. Xenopus fibroblast cells were cotransfected with a GFP fusion to the 198 N-terminal amino acids of MEI-1 and GST-MEI-2. Images of two transfected cells show tracks of GFP:MEI-1(1–198) that overlap or are interspersed with anti-tubulin staining. Arrows point to sites where individual microtubules are dotted with GFP:MEI-1(1–198). Bar = 20 μm.

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Protein expression and purification

E. coli BL21DE3 carrying pMAL-CR1 plasmids were grown at 18°C and induced with IPTG (isopropyl-β-D-thiogalactopyranoside) for 12 h before harvesting and lysis with a microfluidizer. Lysates were centrifuged, and supernatants were subjected to amyllose affinity chromatography and eluted with 20 mM maltose. SF9 cells were grown in SF900 serum-free medium (Invitrogen, Carlsbad, CA) in shaking flasks and infected with each virus for 48 h. Cells were harvested and lysed, and proteins were purified by Ni-chelate chromatography as previously described (McNally et al., 2000).

Protein interaction assay

Amylose beads (New England Biolabs, Ipswich, MA) were incubated with excess lysate from E. coli expressing MBP, MBP-MEI-2, MBP-c15orf29, or MBP-KATNB1(412–655), then washed in binding buffer before addition of Ni-chelate purified baculovirus-expressed proteins. Beads were then washed in binding buffer before elution of bound proteins with SDS Laemml buffer. Binding buffer consisted of 20 mM K-HEPES, pH 7.4, 0.4 mM EGTA, 4 mM MgCl₂, 0.1% Triton X-100, 1 mg/ml lysozyme, and either 400 mM KCl (for MEI-1/MEI-2) or 150 mM KCl (for human katanin). Bound proteins were detected by immunoblotting with either an anti-MEI-1 antibody or an anti-KATNA1 antibody.

Microtubule-binding assays

Ni-chelate–purified 6His-KATNAL1(1–213) (0.6 μM), 0.6 μM amyllose-affinity-purified MBP-c15orf29, or 0.6 μM both proteins was incubated with increasing concentrations (0–19 μM) of paclitaxel (Taxol; Sigma-Aldrich, St. Louis, MO)-stabilized microtubules assembled from phosphocellulose-purified, MAP-free porcine brain tubulin in: 100 mM KCl, 20 mM K-HEPES, pH 7.4, 0.4 mM EGTA, 4 mM MgCl₂, 0.1% Triton X-100, 20 μM paclitaxel, 4 mg/ml bovine serum albumin (BSA) in a volume of 100 μl. Reactions were layered on a 50 μl cushion of 33% glycerol in binding buffer without BSA. Reactions were centrifuged in a TL100 (Beckman Coulter, Brea, CA) fixed-angle tabletop ultracentrifuge rotor at 60,000 rpm for 12 h before harvesting and lysis with a microfluidizer. Supernatants were removed, and pellets were resuspended in SDS-Laemml buffer and analyzed by immunoblotting with an anti-KATNAL1 antibody (McNally and Thomas, 1998) or an anti-MBP antibody. Signals were detected with a Li-COR (Lincoln, NE) Odyssey laser scanner, and the relative fluorescence signals in pellet and supernatant fractions were determined using iVision software. Data were fitted to hyperbolic curves using DeltaGraph software. Data were fitted to hyperbolic curves using DeltaGraph software. Data were fitted to hyperbolic curves using DeltaGraph software. Data were fitted to hyperbolic curves using DeltaGraph software.

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