Single-molecule analysis of the microtubule cross-linking protein MAP65-1 reveals a molecular mechanism for contact-angle-dependent microtubule bundling

Amanda Tulin
Sheri McClerklin
Yue Huang
Ram Dixit

Washington University in St Louis, ramdixit@WUSTLEdu

Follow this and additional works at: https://openscholarship.wustl.edu/bio_facpubs

Part of the Biology Commons, Biophysics Commons, Cell Biology Commons, and the Plant Biology Commons

Recommended Citation
Tulin, Amanda; McClerklin, Sheri; Huang, Yue; and Dixit, Ram, "Single-molecule analysis of the microtubule cross-linking protein MAP65-1 reveals a molecular mechanism for contact-angle-dependent microtubule bundling" (2012). Biology Faculty Publications & Presentations, 76.
https://openscholarship.wustl.edu/bio_facpubs/76

This Article is brought to you for free and open access by the Biology at Washington University Open Scholarship. It has been accepted for inclusion in Biology Faculty Publications & Presentations by an authorized administrator of Washington University Open Scholarship. For more information, please contact digital@wumail.wustl.edu.
Single-molecule analysis of the microtubule crosslinking protein MAP65-1 reveals a molecular mechanism for contact-angle-dependent microtubule bundling
Amanda Tulin, Sheri McClerklin, Yue Huang and Ram Dixit

Biology Department
Washington University in St. Louis
1 Brookings Dr, CB 1137
St. Louis, MO 63130.

Running title: Microtubule bundling by MAP65-1

Keywords: MAP65, microtubule bundle, TIRF microscopy, Arabidopsis

Corresponding Author:
Ram Dixit
Biology Department
Washington University in St. Louis
1 Brookings Drive, CB 1137
St. Louis, MO 63130.
T: (314) 935-8823
F: (314) 935-4432
Email: ramdixit@wustl.edu
Abstract

Bundling of microtubules (MTs) is critical for the formation of complex MT arrays. In land plants, the interphase cortical MTs form bundles specifically following shallow-angle encounters between them. To investigate how cells select particular MT contact angles for bundling, we used an in vitro reconstitution approach consisting of dynamic MTs and the MT-crosslinking protein MAP65-1. We found that MAP65-1 binds to MTs as monomers and inherently targets antiparallel MTs for bundling. Dwell-time analysis showed that the affinity of MAP65-1 for antiparallel overlapping MTs is about three-times higher than the affinity of MAP65-1 for single MTs and parallel overlapping MTs. We also found that purified MAP65-1 exclusively selects shallow-angle MT encounters for bundling, indicating that this activity is an intrinsic property of MAP65-1. Reconstitution experiments with mutant MAP65-1 proteins with different numbers of spectrin repeats within the N-terminal rod domain showed that the length of the rod domain is a major determinant of the range of MT bundling angles. The length of the rod domain also determined the distance between MTs within a bundle. Together, our data show that the rod domain of MAP65-1 acts both as a spacer and as a structural element that specifies the MT encounter angles that are conducive for bundling.
Introduction

Microtubule (MT) bundles play a crucial role in the formation and maintenance of organized MT arrays. In plant cells, the acentrosomal interphase MTs at the cell cortex are highly bundled and their spatial organization dictates the direction of cell expansion (1). These so-called cortical MTs are nucleated from dispersed sites at the cell cortex (2-4) and are attached to the plasma membrane along their lengths (5, 6). The cortical MTs are highly dynamic and treadmill along the plasma membrane surface, leading to frequent interactions between them (2). A subset of these interactions leads to the formation of cortical MT bundles. Importantly, the encounter angle between interacting cortical MTs is a key determinant of the bundling probability (7). Specifically, bundling is observed to occur only after shallow-angle interactions (< 40°) between cortical MTs (7). Steep-angle cortical MT interactions are followed by either MT crossover or depolymerization (7). The dependency on the encounter angle for cortical MT bundling appears to be important for the proper organization of the cortical MT array because large shifts in the distribution of bundling angles were found to hinder cortical MT organization in computer simulations (8). However, the molecular basis for why only shallow-angle encounters lead to cortical MT bundling is unknown.

MT bundles are generated by the activity of MT-crosslinking proteins. The conserved MAP65/Ase1/PRC1 family of MT-crosslinking proteins plays a major role in the formation of both interphase and mitotic MT arrays. The Arabidopsis genome encodes nine MAP65 proteins of which MAP65-1, MAP65-2, MAP65-5 and MAP65-8 localize to cortical MTs in vivo (9-11). Recently, genetic analyses have revealed that MAP65-1 and MAP65-2 together regulate cell growth during interphase (12) and play a role in cytokinesis (13). The Arabidopsis MAP65-1 is the most extensively studied isoform and is the focus of this study. Purified MAP65-1 bundles taxol-stabilized MTs in vitro and appears as filamentous cross-bridges that separate adjacent MTs by a distance of about 25 nm (14, 15). Electron microscopic observation of bundled cortical MTs in vivo shows that the spacing between adjacent MTs is also about 25 nm (6, 16, 17), indicating that the MAP65 proteins are the major MT bundling proteins in this system. Similar to other MAP65/Ase1/PRC1 members, MAP65-1 is able to discriminate between parallel and antiparallel MTs in vitro and localizes to regions of antiparallel MT overlap with high specificity (15). Consistent with these results, MAP65-1 has been recently shown to preferentially label bundled cortical MTs in vivo, a significant subset of which contain antiparallel MTs (12).

Structural modeling of MAP65-1 based on fold-recognition predicts the presence of four spectrin repeats that are thought to form an extended rod-like structure about 25-nm in length (18). This N-terminal “rod” domain of MAP65-1 is thought to be flexible when bound to a single MT, based on the presence of several disordered domains in its sequence and on its hydrodynamic properties (15). Recent structural analysis of PRC1, the human MAP65 homolog, also suggests that the rod domain is likely to be flexible when PRC1 is bound to a single MT (19). Monomers of MAP65-1 are proposed to homodimerize through their rod domains, thus creating a cross-bridge between adjacent MTs (15). We hypothesize that a long and flexible rod domain might allow MAP65-1 to homodimerize within a particular range of angular orientations, thus specifying the range of bundling angles.

To investigate whether the rod domain of MAP65-1 is responsible for specifically selecting shallow-angle MT encounters for bundle formation, we developed a cell-free in vitro reconstitution assay consisting of dynamic MTs and purified MAP65-1. We found that MAP65-1 inherently selects shallow-angle encounters between antiparallel MTs for bundling. Time-lapse imaging of GFP-tagged MAP65-1 showed that MAP65-1 preferentially accumulates at and
dynamically tracks with regions of antiparallel MT overlap. This property is associated with an increase in the dwell-time of MAP65-1 within regions of antiparallel MT overlap. Reconstitution experiments with mutant versions of MAP65-1 that either lack a spectrin repeat or have additional spectrin repeats showed that the length of the rod domain determines both the spacing between crosslinked MTs and the range of encounter angles that lead to MT bundling. Together, these data provide a molecular mechanism for why only certain encounter angles lead to cortical MT bundling in plant cells.
Materials and Methods

Protein expression and purification
Constructs for protein expression were prepared using PCR and verified by sequencing. See supplemental Table S1 for the list of primers used to generate the constructs. Verified PCR products were introduced into the pET-28a(+) vector (Novagen) which encodes for a 6x histidine tag at the N-terminus of proteins. The assembled plasmids were introduced into Rosetta (DE3) cells (Novagen) for protein expression. His-tagged recombinant proteins were affinity-purified using a nickel column and subsequently desalted using a PD-10 column (GE Biosciences) and exchanged into BRB80 buffer (80 mM piperazine-1,4-bis(2-ethanesulfonic acid), 1mM MgCl₂, 1 mM EGTA, pH 6.8). Protein aliquots were flash frozen in liquid nitrogen and stored at -80 °C until use.

MT-binding assays
All MTs in this study were assembled in BRB80 buffer using purified bovine tubulin (Cytoskeleton, Inc.). The MT-binding assays were conducted by co-incubating increasing concentrations of MTs with 1.5 μM of the specified recombinant protein along with 20 μM paclitaxel (Cytoskeleton, Inc.) at 25 °C for 30 min. The samples were then centrifuged at 39,000g for 20 min at 25 °C to sediment the MTs. The resultant supernatant and pellet fractions were analyzed by SDS/PAGE and densitometry to calculate the bound fraction. The ΔR1 protein co-migrates with tubulin and therefore we used western blot analysis with a monoclonal Tetra-His antibody (Qiagen) to detect ΔR1 in the supernatant and pellet fractions. Densitometry was carried out using ImageJ. For analysis of MT bundling using taxol-stabilized MTs, 1 μM rhodamine-labeled and taxol-stabilized MTs were co-incubated with 1 μM of the specified recombinant protein at 25 °C for 30 min and then visualized using fluorescence microscopy.

Reconstitution experiments with dynamic MTs
The in vitro reconstitution assay was developed based on our previously described method (20). Briefly, flow chambers of about 20 μl volume were prepared using silanized coverslips attached to slides with double-sided sticky tape. The flow cell was coated with 20% monoclonal anti-biotin antibody (clone BN-34, Sigma) and then blocked with 5% pluronic F-127 (Sigma). About 150 nM rhodamine-labeled and biotinylated guanosine 5‘-(α,β-methylene)triphosphate (GMPCPP) MT seeds were then introduced into the flow cell. MT growth and bundling was initiated by introducing 20 μM 1:40 rhodamine-labeled bovine tubulin in BRB80 buffer and the specified MAP65-1 protein along with 0.15% methylcellulose, 100 mM DTT, an oxygen scavenging system consisting of 250 μg/ml glucose oxidase, 35 μg/ml catalase and 4.5 mg/ml glucose, and 2 mM GTP. The samples were excited with 488-nm (at 10 mW output) and 561-nm (at 4 mW output) diode-pumped solid-state lasers (Melles Griot) to visualize MAP65-1-GFP and rhodamine-labeled MTs respectively. Time-lapse images were captured with a back illuminated electron-multiplying CCD camera (Hamamatsu, ImageEM) and GFP (500-550 nm emission) and rhodamine (582-636 nm emission) filter sets. The polarity of growing MTs was assigned based on the difference in growth velocity between the plus- and minus-end. Kymograph analysis was conducted using Slidebook 5.0 (Intelligent Imaging Innovations). Curve fitting and statistical analysis was conducted using KaleidaGraph (Synergy Software).

Single molecule imaging
For photobleaching assays, 1 nM MAP65-1-GFP bound to rhodamine-labeled and taxol-stabilized MTs were imaged at higher laser power (20 mW output from the 488-nm laser) and the fluorescence intensities of individual spots were measured over time to determine the number of bleaching steps. For comparison, 10 nM human kinesin1-GFP bound to rhodamine-labeled and taxol-stabilized MTs in the presence of AMPPNP were analyzed using identical image acquisition conditions. For dwell-time analysis, reconstitution assays were conducted using 400 nM unlabeled MAP65-1 containing 8 nM MAP65-1-GFP. Kymographs of single and bundled MTs were generated using Slidebook 5.0 and used to measure the dwell-times of individual molecules.

**Electron microscopy**

For negative-stain electron microscopy of MT bundles, 1 μM of taxol-stabilized MTs and 1 μM recombinant protein were co-incubated at 25 °C for 30 min. The MT suspension was then applied to formvar-coated grids and stained with a 7% (aqueous) solution of uranyl acetate for two minutes. The grids were then blotted dry and examined in a LEO 912 AB energy filter TEM operated at 120 kV.
Results

MAP65-1 inherently selects shallow-angle MT encounters for bundling

To study the MT bundling activity of MAP65-1, we purified full-length MAP65-1 expressed in bacteria (Fig. 1A). In vitro MT binding experiments showed that the equilibrium $K_d$ of MAP65-1 for MTs is $1.03 \pm 0.75 \mu M$ (Fig. 1B), which is similar to that of Ase1, PRC1 and tobacco MAP65-1b (19, 21, 22). Since MAP65-1 is thought to dimerize within MT bundles (15), the measured $K_d$ of MAP65-1 is likely to be a composite of MT binding and MAP65-1 dimerization.

Next, we developed an in vitro reconstitution assay consisting of dynamic MTs and purified MAP65-1 to observe MT bundling by MAP65-1 using time-lapse total internal reflection fluorescence microscopy. In these experiments, growing MTs encountered each other along the cover glass surface and we noticed that only a subset of these MT encounters led to bundling. In control experiments lacking MAP65-1, we never observed MT bundling (Fig. S2). Analysis of the polarity of MAP65-1-induced MT bundles demonstrated that about 90% of the MT bundles consisted of antiparallel MTs ($N = 148$; Fig. 1C; Movie S1). Only about 10% of MT bundles were between parallel MTs (Movie S2). Therefore, MAP65-1 inherently discriminates between parallel and antiparallel MTs.

In addition to the strong preference of MAP65-1 for crosslinking antiparallel MTs, we found that only a narrow range of MT encounter angles yielded MT bundles. Specifically, shallow-angle MT encounters invariably led to MT bundling, whereas steep-angle MT encounters led to MT crossover. This was true for both antiparallel (Figs. 1D and 1E) and parallel MT bundling (Figs. S1A and S1B). To determine if the MT bundling angle is a function of the MAP65-1 concentration, we conducted reconstitution experiments at increasing MAP65-1 concentrations. Increasing the MAP65-1 concentration from 100 nM to 400 nM shifted the bundling probability to larger encounter angles (Fig. 1D). A further increase in MAP65-1 concentration to 800 nM did not significantly affect the probability of MT bundling compared to 400 nM MAP65-1 (Fig. 1D). Therefore, 400 nM MAP65-1 is sufficient to result in maximal MT bundling under our experimental conditions.

The distribution of bundling angles at 400 nM MAP65-1 (Fig. 1E) is strikingly similar to the distribution of bundling angles for cortical MTs in living Arabidopsis plants (5). The different types of cortical MT bundling events that have been seen in cells were also observed in our reconstitution experiments: i) in 64% of the cases, the growing plus-end of a MT encountered the sidewall of another MT followed by reorientation of its growth trajectory and continued polymerization alongside the impeding MT (Movie S3). This scenario has been called plus-end entrainment (1); ii) in 30% of the cases, MTs are observed to progressively coalign along their lengths (Movie S4), which has been called zippering (1, 7); and iii) in 6% of the cases, individual MTs instantly snapped together to form a bundle (Movie S5), as seen in both wild-type Arabidopsis plants and clasp-1 mutants (1, 5). Together, these data suggest that our reconstitution experiments with 400 nM MAP65-1 mimic the physiological conditions in plant cells.

The dependency on the encounter angle for MT bundling was most convincingly demonstrated in cases when a MT initially crossed over another MT (at a steep encounter angle) but later become bundled as the crossover angle decreased to a shallow angle (Fig. 1F and Movie S6). These examples highlight the inherent ability of MAP65-1 to discriminate between MT encounter angles and to selectively target shallow-angle MT encounters for bundling.
MAP65-1 dynamically tracks regions of MT overlap

To understand how MAP65-1 selectively bundles particular MT configurations, we generated a construct to express full-length MAP65-1 with GFP fused to its C-terminus. In vitro MT binding experiments showed that the $K_d$ of MAP65-1-GFP for MTs is $1.27 \pm 0.68 \mu M$, which is statistically indistinguishable from the $K_d$ of untagged MAP65-1 (Fig. 1B). Initial tests also showed that MAP65-1-GFP is able to bundle taxol-stabilized MTs (Fig. S2). Therefore, the GFP tag does not interfere with MAP65-1’s ability to bind and crosslink MTs.

Ase1 and PRC1 bind to MTs as dimers (19, 23, 24). Attempts to determine whether MAP65-1 binds to MTs as a dimer or as a monomer have yielded mixed results (14, 15). To directly determine if our purified MAP65-1 binds to MTs as a monomer or dimer, we performed photobleaching experiments of 1 nM MAP65-1-GFP bound to taxol-stabilized MTs. Analysis of the intensity traces of individual spots revealed that the fluorescence intensity of a majority of MAP65-1-GFP spots decreased to background levels in a single step, indicating the presence of a single GFP molecule that photobleached during the observation period (Fig. 2A). In contrast, photobleaching analysis of human Kinesin1-GFP under identical imaging conditions showed mostly two bleaching steps, consistent with the presence of two GFP molecules in the kinesin-1 dimer (Fig. 2A). Therefore, our data indicate that MAP65-1-GFP binds to MTs predominantly as a monomer.

We next carried out reconstitution experiments using 400 nM MAP65-1-GFP. We found that MAP65-1-GFP specifically accumulated at regions of antiparallel MT bundling following shallow-angle MT encounters (Movie S7). Kymograph analysis of MT bundles showed that MAP65-1-GFP dynamically tracks the regions of MT overlap (Fig. 2B). In the same experiments, we detected little MAP65-1-GFP accumulation along single MTs and parallel MT bundles. Analysis of the dwell time of individual MAP65-1-GFP spots revealed that the dwell time of MAP65-1-GFP increased by about 3-fold on antiparallel MT bundles as compared to the dwell time on single MTs (Fig. 2C) and on parallel MT bundles (Fig. S1C). These results indicate that a decrease in the MT unbinding rate underlies the ability of MAP65-1 to selectively accumulate at regions of antiparallel MT overlap, similar to that described for Ase1 (23).

The length of the rod domain of MAP65-1 specifies the range of MT bundling angles

To test if the rod domain of MAP65-1 is involved in specifying the MT bundling angles, we generated constructs to express and purify several mutant versions of MAP65-1 with either shorter or longer rod domains compared to wild-type MAP65-1 (Fig. 3A and 3B). The mutant proteins were designated ΔR1 (first spectrin repeat deleted), ΔR2 (second spectrin repeat deleted) and R1R4 (entire spectrin repeat domain duplicated). In vitro MT binding experiments showed that the $K_d$ of the various mutant proteins for MTs is similar to that of wild-type MAP65-1 (Fig. 3C). Once again, we note that these values represent a convolution of both MT binding and MAP65 protein dimerization, whose individual contributions cannot be discriminated in these binding experiments. The mutant proteins are also able to bundle taxol-stabilized MTs (Fig. S2). Therefore, all of the mutant proteins are able to bind and bundle MTs. We found that 400 nM ΔR1 bundles MTs more weakly as compared to the other proteins. Therefore, for our subsequent experiments we increased the protein concentration to 800 nM ΔR1. This increase in ΔR1 protein concentration does not hinder interpretation of data because the MT bundling angle distributions are similar using either 400 nM or 800 nM of wild-type MAP65-1 protein (Fig. 1D).
To confirm that the ΔR1, ΔR2 and R1R4 mutants altered the spacing between bundled MTs as expected from the predicted lengths of their rod domains, we performed negative-stain electron microscopy of MTs incubated with these proteins. Electron micrographs of MTs bundled by wild-type MAP65-1 showed coaligned MTs separated by an average distance of about 24 nm (Fig. 4B). In contrast, the spacing between MTs in bundles induced by ΔR1 and ΔR2 is about 9 nm and 10 nm, respectively (Fig. 4C and D). This is consistent with previous measurements of distances separating MTs bundled by ΔR1 and ΔR2 (15). Electron micrographs of MTs bundled by R1R4 showed that the average distance between MTs is increased to about 37 nm (Fig. 4E). These results indicate that the ΔR1, ΔR2 and R1R4 mutants indeed produce the expected decrease or increase in inter-MT spacing as predicted by the number of spectrin repeats in their rod domain.

To determine if the ΔR1, ΔR2 and R1R4 mutants altered the distribution of the MT bundling angles, we conducted reconstitution experiments with 800 nM ΔR1, 400 nM ΔR2 and 400 nM R1R4. We found that both ΔR1 and ΔR2 target only very shallow-angle MT encounters for bundling and generally took several attempts to initiate MT bundling as compared to MAP65-1 (Movies S8 and S9). Analysis of the MT bundling angles showed that both ΔR1 and ΔR2 shift the distribution of MT bundling angles to smaller angles compared to MAP65-1 (Fig. 5A, B and C). The mean bundling angles are 16° and 18° with ΔR1 and ΔR2 respectively, which are significantly lower than the mean bundling angle of 28° with MAP65-1 (p < 0.0001 using the t-test). In contrast, R1R4 frequently resulted in MT bundling even after steep-angle MT encounters (Movie S10). Analysis of the MT bundling angles showed that R1R4 dramatically expands the distribution of MT bundling angles to include steep angles (Fig. 5D). The mean bundling angle with R1R4 is 36°, which is significantly higher than the mean bundling angle with MAP65-1 (p < 0.0001 using the t-test). Analysis of the bundling probability as a function of the contact angle shows a striking leftward shift for ΔR1 and ΔR2, while R1R4 shows a striking rightward shift as compared to MAP65-1 (Fig. 5E). Based on these results, we conclude that the length of the rod domain of MAP65-1 is a major determinant of the MT bundling angle.
Discussion

In this study, we sought to understand the molecular basis for the observation that only shallow-angle encounters between cortical MTs result in bundle formation in plant cells. This feature is an important aspect of cortical MT array organization because only similarly oriented cortical MTs are allowed to productively interact and form bundles, thus promoting the formation of linearly ordered arrays. Using a minimal system consisting of dynamic MTs and purified MAP65-1, we found that the ability to selectively bundle MTs that interact at a shallow angle is an intrinsic property of MAP65-1 and does not require additional factors. Furthermore, we found that the length of the rod domain of MAP65-1 determines the range of MT bundling angles, thus providing insight into the structural feature of MAP65-1 that is responsible for bundling angle selection.

We found that increasing the MAP65-1 concentration increases the range of MT bundling angles up to a certain limit. This observation is consistent with the prediction from a theoretical model of cortical MT interactions which posits that an increase in the concentration of a MT crosslinking protein will increase the probability of MT bundling by increasing the torque necessary to bend an incoming MT along the impeding MT (25). Notably, once the torque exerted by the crosslinking protein exceeds the bending rigidity of the incoming MT, any further increase in the concentration of the crosslinking protein would have little effect, in agreement with our finding. Based on our data, regulation of the intracellular concentration of MAP65 proteins offers cells a mechanism to specify which MT encounters will lead to bundling. This ability may be important during MT array formation, remodeling and disassembly.

Like other members of the MAP65/Ase1/PRC1 family, we found that MAP65-1 can inherently distinguish between parallel and antiparallel MTs. In our in vitro experiments, about 90% of the MAP65-1-induced MT bundles consisted of antiparallel MTs. This is comparable to Ase1p and PRC1, which yield antiparallel MT bundles about 70% (23) and 90% (19) of the time, respectively. Our results are also consistent with previous work which showed that MAP65-1 localizes to antiparallel MT bundles both in vitro (15) and in vivo (12). Dwell-time analysis of individual MAP65-1 molecules showed that the off rate on antiparallel MT overlaps was about 3-fold lower than on single MTs and parallel MT overlaps. The increased affinity for antiparallel MTs provides a possible explanation for the selective cross-linking of antiparallel MTs by MAP65-1.

In our assays, the constituent MTs within a bundle remain dynamic and MAP65-1 is observed to dynamically track the regions of antiparallel MT overlap, strikingly illustrating the differential binding of MAP65-1 to antiparallel MT overlaps versus single MTs. Fluorescently tagged MAP65-1 is similarly observed to track along bundled segments of cortical MTs in Arabidopsis plants (12). This property of MAP65-1 is similar to that of the mitotic MAP65-4 (26), MAP65-3 (27), Ase1 (28) and PRC1 (19, 24) and thus appears to be a conserved feature of the MAP65/Ase1/PRC1 family.

Our measured dwell-time of about 2 sec for individual MAP65-1 molecules on antiparallel MTs in vitro is in good agreement with the reported bulk turnover rate of about 5 sec for MAP65-1 on cortical MT bundles in vivo (11, 12). Photobleaching analysis of individual MAP65-1 molecules showed that MAP65-1 binds to MTs as a monomer, which is consistent with Gaillard et al. (15) who concluded that MAP65-1 is monomeric in solution based on analytical ultracentrifugation and size exclusion chromatography experiments. Therefore, it is not necessary for MAP65/Ase1/PRC1 homologs to assemble into pre-formed dimers to be able to bundle MTs.
MT bundling requires the formation of antiparallel dimers from monomeric MAP65-1 subunits bound to separate MTs in order to form a stable crosslink between encountering MTs. Biochemical evidence indicates that the spectrin repeats in the rod domain of MAP65-1 mediate the formation of an antiparallel dimer as described for muscle α-actinin (15, 18). The rod domain of MAP65-1 is also likely to be a flexible structure when bound to a single MT (15, 19). The conformational flexibility of the rod domain may allow MAP65-1 monomers at multiple orientations to dimerize, thus increasing the chances for MT bundling. A possible mechanism for why only certain MT encounter angles lead to MT bundling is that these MT orientations position the MAP65-1 monomers in a way that allows them to productively interact and dimerize. Thus, MT orientations that allow MAP65-1 monomers to dimerize will lead to bundling while other MT orientations which are not conducive for MAP65-1 dimerization will fail to produce MT bundles (Fig. 6A-6C).

In our in vitro reconstitution experiments, the length of the rod domain had a strong effect on the MT bundling angle. Shortening the rod domain by deleting a spectrin repeat constrained the bundling angles to smaller values, while lengthening the rod domain by including additional spectrin repeats greatly expanded the range of bundling angles to include larger values, as compared to wild-type MAP65-1. Deletion of either the first or the second spectrin repeat resulted in a similar shift in the distribution of bundling angles, indicating that the length of the rod domain and not a particular sequence is the key determinant of the MT bundling angle. The length of the rod domain of MAP65-1 may impact the MT bundling angle in at least two ways that are not mutually exclusive: i) it might affect the efficiency and/or strength of dimer formation based on the extent of overlap that would be possible between the rod domains of MAP65-1 monomers; and ii) it might affect the range of the angular sector that the rod domain explores given its conformational flexibility. In particular, the shorter rod domains of the ΔR1 and ΔR2 mutants might be stiffer, thus allowing their dimerization and consequently MT bundling only at very shallow encounter angles (Fig. 6D). In contrast, the longer rod domain of the R1R4 mutant is envisioned to be more flexible than the rod domain of wild-type MAP65-1, which would allow the R1R4 mutant to dimerize and bundle MTs that are even higher encounter angles (Fig. 6E). Besides affecting the MT bundling angle, we found that the length of the rod domain of MAP65-1 also acts as a spacer that determines the distance between MTs within a bundle.

A similar mechanism for selectively bundling shallow-angle MT interactions is probably applicable to MAP65 homologs that bind to MTs as dimers. Ase1 and PRC1 bind to MTs as dimers and both specifically bundle MTs that interact at shallow angles (19, 28). The flexibility of the rod domain of PRC1 dimers has been proposed to allow contact with a second MT within a certain range of MT orientations, thus determining the acceptable MT bundling angles (19). Interestingly, the distribution of MT bundling angles for Ase1 is very similar to that of the ΔR1 and ΔR2 mutants (28) and the distance between Ase1-induced MT bundles is about 6 nm (21), which is in the range of the MT spacing by the ΔR1 and ΔR2 mutants. Therefore, the length of the rod domain is likely to be an important determinant of the MT bundling angle even for dimeric MAP65 homologs.
Acknowledgements
The authors thank Dr. Howard Berg at the Donald Danforth Plant Science Center for the electron microscopy of microtubule bundles. This work was partially supported by funds from I-CARES, Washington University in St. Louis.
References
Figure Legends

Figure 1: MAP65-1 preferentially bundles antiparallel MTs after shallow-angle encounters. (A) Coomassie-stained gel of purified MAP65-1 and MAP65-1-GFP proteins. The expected protein sizes are marked by asterisks. (B) Binding curves with 1.5 μM MAP65-1 and MAP65-1-GFP proteins at increasing MT concentrations. Each data point represents the mean ± SD from at least three independent experiments. The data were fit to the Michaelis-Menten equation yielding Kd’s of 1.03 ± 0.75 μM and 1.27 ± 0.68 μM for MAP65-1 and MAP65-1-GFP respectively. (C) Montage showing antiparallel MT bundling by 400 nM MAP65-1. The plus-ends of the MTs of interest are indicated in the first frame. Arrowheads mark the position of the plus-end within the MT bundle. (D) Plots showing the probability for MT bundling as a function of the encounter angle at various MAP65-1 concentrations. The bundling probability was calculated as a percentage of the number of MT encounters that resulted in MT bundling at a particular angle. The total number of MT encounters observed for 100 nM, 200 nM, 400 nM and 800 nM of MAP65-1 are 245, 243, 323 and 311 respectively. (E) Distribution of the frequency of MT bundling at various encounter angles in the presence of 400 nM MAP65-1 (N = 199 events). The mean MT bundling angle is 28 ± 13°. (F) MT bundling following a decrease in the crossover angle from 60° to 35°. The arrow indicates the direction of the growing plus-end of the MT of interest. Numbers in (C) and (F) indicate time in seconds. Scale bars = 2 μm.

Figure 2: MAP65-1 binds to MTs as a monomer and preferentially localizes to regions of MT overlap. (A) Bar graph of the number of bleaching steps for MAP65-1-GFP and Kinesin1-GFP molecules bound to taxol-stabilized MTs (N = 177 and 171 for MAP65-1-GFP and Kinesin1-GFP respectively). Examples of fluorescence intensity traces showing one and two bleaching steps are shown to the right. (B) Kymograph showing the localization of 400 nM MAP65-1-GFP in an antiparallel MT bundle. MAP65-1-GFP specifically tracks the region of MT overlap and is barely detectable along stretches with a single MT. (C) To the left are kymographs showing the binding of 8 nM MAP65-1-GFP to a single MT and an antiparallel MT bundle. To the right are the distributions of dwell-times of single binding events of MAP65-1-GFP on single MTs (N = 257) and bundled MTs (N = 384). Exponential fits to the data yielded halftimes of 0.62 ± 0.07 s and 1.82 ± 0.01 s respectively.

Figure 3: Purification and MT-binding of MAP65-1 mutants. (A) Schematic of the domain architecture of MAP65-1 and the various mutants used in this study. The four predicted spectrin repeats are labeled R1 to R4. Tail refers to the unstructured domain at the C-terminus of MAP65-1. (B) Coomassie-stained gel of purified ΔR1, ΔR2 and R1R4 proteins. The expected protein sizes are marked by asterisks. (C) Binding curves with 1.5 μM ΔR1, ΔR2 and R1R4 proteins at increasing MT concentrations. Each data point represents the mean ± SD from at least three independent experiments. The data were fit to the Michaelis-Menten equation yielding Kd’s of 1.17 ± 0.73 μM, 1.04 ± 0.61 μM and 1.04 ± 0.63 μM for ΔR1, ΔR2 and R1R4 respectively. The binding curve for MAP65-1 is reproduced from Figure 1B for comparison to the mutant proteins.

Figure 4: The length of the rod domain of MAP65-1 determines the distance between MTs in a bundle. Negative-stain electron microscopy of 1 μM MTs alone (A) or 1 μM MTs co-incubated with 1 μM of MAP65-1 (B), ΔR1 (C), ΔR2 (D) and R1R4 (E) respectively. The mean distance ± SD in nm are shown in the figure. The number of independent measurements between separate MTs is shown within parentheses. Scale bars = 50 nm.
Figure 5: The length of the rod domain of MAP65-1 determines the MT bundling angle. (A-D) Distribution of the frequency of MT bundling at various encounter angles in the presence of 400 nM MAP65-1 (A), 800 nM ΔR1 (B), 400 nM ΔR2 (C) and 400 nM R1R4 (D). The data for MAP65-1 is reproduced from Figure 1E for comparison to the mutant proteins. N = 199, 189, 231 and 295 for MAP65-1, ΔR1, ΔR2 and R1R4 respectively. The mean bundling angle ± SD are shown in the figure. (E) Plots showing the probability for MT bundling as a function of the encounter angle in the presence of 400 nM MAP65-1, 800 nM ΔR1, 400 nM ΔR2 and 400 nM R1R4.

Figure 6: Model for encounter-angle-dependent MT bundling by MAP65-1. MAP65-1 monomers are shown bound to MTs (a single MT protofilament is shown for simplicity). The blue portion of MAP65-1 represents the fourth spectrin repeat which contains the MT-binding domain. The unstructured tail domain of MAP65-1 is shown in red. The N-terminal rod domain is shown in green and the conformational flexibility of this domain is represented by its multiple positions. If two MTs encounter each other nearly parallel (A) or at a shallow-angle (B), the MAP65-1 monomers are able to dimerize and form a stable crosslink, thus resulting in MT bundling. In contrast, if two MTs encounter each other at a steep angle, the MAP65-1 monomers are unable to dimerize because their rod domains cannot interact productively at these angles (C). Consequently, these MTs do not bundle. In the case of the ΔR1 and ΔR2 mutants, their shorter rod domains are probably stiffer, thus requiring even shallower encounter angles for dimer formation (D). In contrast, the R1R4 mutant has a longer rod domain that is likely to be more flexible than the rod domain of wild-type MAP65-1, which allows dimer formation and MT bundling even at steep encounter angles (E).
Figure 2

A

B

C
Figure 3

A

B

C

Bound fraction (%) vs. MT concentration (µM)
Figure 4

<table>
<thead>
<tr>
<th></th>
<th>Distance ± SD (N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAP65-1</td>
<td>23.7 ± 3.8 (26)</td>
</tr>
<tr>
<td>△R1</td>
<td>8.7 ± 0.6 (31)</td>
</tr>
<tr>
<td>△R2</td>
<td>9.8 ± 0.9 (25)</td>
</tr>
<tr>
<td>R1R4</td>
<td>36.8 ± 6.3 (28)</td>
</tr>
</tbody>
</table>
Figure 5

(A) MAP65-1
26° ± 13

(B) ∆R1
16° ± 13

(C) ∆R2
18° ± 11

(D) R1R4
36° ± 17

(E) Bundling probability (%)

Legend:
- MAP65-1
- ∆R1
- ∆R2
- R1R4
Figure 6

A

B

C

D

E