Novel septin 9 repeat motifs altered in neuralgic amyotrophy bind and bundle microtubules

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Septin 9 (SEPT9) interacts with microtubules (MTs) and is mutated in hereditary neuralgic amyotrophy (HNA), an autosomal-dominant neuropathy. The mechanism of SEPT9 interaction with MTs and the molecular basis of HNA are unknown. Here, we show that the N-terminal domain of SEPT9 contains the novel repeat motifs K/R-x-x-E/D and R/K-R-x-E, which bind and bundle MTs by interacting with the acidic C-terminal tails of β-tubulin. Alanine scanning mutagenesis revealed that the K/R-R/x-x-E/D motifs pair electrostatically with one another and the tails of β-tubulin, enabling septin–septin interactions that link MTs together. SEPT9 isoforms lacking repeat motifs or containing the HNA-linked mutation R88W, which maps to the R/K-R-x-E motif, diminished intracellular MT bundling and impaired asymmetric neurite growth in PC-12 cells. Thus, the SEPT9 repeat motifs bind and bundle MTs, and thereby promote asymmetric neurite growth. These results provide the first insight into the mechanism of septin interaction with MTs and the molecular and cellular basis of HNA.

Introduction

Septins are hetero-oligomeric GTP-binding proteins that assemble into nonpolar filamentous structures, which are essential for microtubule (MT)-dependent cell processes such as mitosis and vesicle transport (Spiliotis, 2010; Saarikangas and Barral, 2011; Mostowy and Cossart, 2012). In mammalian cells, septin depletion affects MT organization, dynamics, and post-translational modifications (Spiliotis et al., 2008; Bowen et al., 2011), but how septins interact with MTs is unknown. Septin 9 (SEPT9) is a ubiquitously expressed septin that caps the ends of septin heteromers (Füchtbauer et al., 2011; Kim et al., 2011; Sellin et al., 2011). Alternative splicing and translation start sites give rise to SEPT9 isoforms, which differ in the length and sequence of their N terminus, which is critical for the association of septin heteromers with MTs (Sellin et al., 2012). Missense mutations in the N terminus of the SEPT9 are genetically linked to hereditary neuralgic amyotrophy (HNA), a hereditary neuralgic disorder with symptoms (shoulder/arm pain and atrophy) similar to those of the idiopathic Parsonage-Turner syndrome (Kuhlenbäumer et al., 2005; Hannibal et al., 2009; van Alfen, 2011). Abnormal expression of SEPT9 isoforms has also been reported in many cancers (Connolly et al., 2011a). Overexpression of SEPT9 isoforms induces oncogenic phenotypes including resistance to the anti-cancer drug paclitaxel, which stabilizes and bundles MTs (Orr et al., 2003; Gonzalez et al., 2007; Chacko et al., 2012). Unraveling the mechanism of SEPT9 interaction with MTs could point to the molecular roles of SEPT9 in these disorders and lead to therapeutic strategies.

Results and discussion

The basic N-terminal domain of SEPT9 binds and bundles MTs by interacting with the acidic C-terminal tails of β-tubulin

To determine how SEPT9 interacts with MTs, we first sought to identify the SEPT9 domain that binds MTs using in vitro co-sedimentation assays. Recombinant His-tagged fragments were made from the longest SEPT9_i1 isoform (SEPT9-FL; Fig. 1A), which consists of the GTP-binding domain (aa 283–586; SEPT9-G) and a structurally disordered N-terminal tail (aa 1–283; SEPT9-N; Fig. S1A) with a basic region (aa 1–142; SEPT9-B). Alternative splicing and translation start sites give rise to SEPT9 isoforms, which differ in the length and sequence of their N terminus, which is critical for the association of septin heteromers with MTs (Sellin et al., 2012). Missense mutations in the N terminus of the SEPT9 are genetically linked to hereditary neuralgic amyotrophy (HNA), a hereditary neuralgic disorder with symptoms (shoulder/arm pain and atrophy) similar to those of the idiopathic Parsonage-Turner syndrome (Kuhlenbäumer et al., 2005; Hannibal et al., 2009; van Alfen, 2011). Abnormal expression of SEPT9 isoforms has also been reported in many cancers (Connolly et al., 2011a). Overexpression of SEPT9 isoforms induces oncogenic phenotypes including resistance to the anti-cancer drug paclitaxel, which stabilizes and bundles MTs (Orr et al., 2003; Gonzalez et al., 2007; Chacko et al., 2012). Unraveling the mechanism of SEPT9 interaction with MTs could point to the molecular roles of SEPT9 in these disorders and lead to therapeutic strategies.
proline-rich acidic domain (aa 143–283; SEPT9-A). SEPT9-FL co-sedimented with MTs (Fig. 1, B and J) and the apparent $K_d$ value was 3.2 µM (Fig. S1 B). In contrast to SEPT9-G and SEPT9-A, which pelleted with MTs weakly (Fig. 1, D–F and J), SEPT9-N and SEPT9-B bound MTs similarly to SEPT9-FL (Fig. 1, C, E, and J).

Next, low speed MT pelleting and visual MT assays were used to examine whether SEPT9 can bundle MTs. At low speed (8,000 g), MT sedimentation was increased by SEPT9-FL, SEPT9-N, and SEPT9-B (Fig. 1, G–I and K), but SEPT9-G and SEPT9-A had no effect (Fig. 1, H, I, and K). Incubation of fluorescent MTs with SEPT9-FL showed a marked increase in the length, thickness, and brightness of MT bundles (Fig. 1, L, N, and O). This bundling effect was independently confirmed by negative stain EM. MTs were arranged in bundles and doublets of parallel tubules that made tight contacts (Fig. 1 M).

Elongated MT bundles were also observed with fluorescent MTs that were stabilized with the nonhydrolyzable GTP analogue GMP-CPP (Fig. S1 C). Decoration of paclitaxel- and GMP-CPP-stabilized MT bundles with recombinant GFP-tagged SEPT9-FL, but not SEPT2, demonstrated that SEPT9 binds and bundles MTs strongly and specifically (Fig. S1, D–I). In agreement with the MT sedimentation assays, significant bundling of fluorescent MTs was observed for only SEPT9-N and SEPT9-B (Fig. 1, L, N, and O). These data indicate that SEPT9 binds and bundles MTs via its basic N-terminal domain.

Because SEPT9 forms hetero-oligomeric complexes with other septins (e.g., SEPT7, SEPT6), we purified SEPT9 in the presence of SEPT6/7 and tested how SEPT6/7/9 complexes affect MT bundling compared with SEPT9 alone or SEPT2/6/7. SEPT6/7/9 induced low speed sedimentation of MTs at a concentration (0.2 µM) that neither SEPT9 nor SEPT2/6/7 had an effect (Fig. S1, J and K). Thus, SEPT9 appears to be a key subunit for the bundling of MTs by septin complexes.

Electrostatic interactions between positively charged domains and the negatively charged C-terminal tails (CTTs) of tubulin underlie the mechanism by which some microtubule-associated proteins (MAPs) and kinesin motors associate with MTs (Amos and Schlieper, 2005; Marx et al., 2006; Akhmanova and Steinmetz, 2008). Although we were not able to identify any known motifs, the N terminus of SEPT9_i1 contains multiple repeats of the tetrapeptide sequences K/R-x-x-E/D and R/K-R-x-E (Fig. 3, A and B), which are often flanked by proline or serine residues. By truncating the basic N-terminal domain of SEPT9_i1, we determined that aa 61–113 comprise the most minimal region of SEPT9_i1 that binds and bundles MTs (Fig. S2, A–C) and contains multiple repeats of the identified motifs.

We set out to elucidate the role of these novel motifs in MT binding and bundling by alanine scanning mutagenesis. The first half of the SEPT9_i1(61–113) peptide contains a region (R1) of three K/R-x-x-E/D motifs and the second half (R2) consists of a K-x-x-E motif flanked by two R-R-x-E motifs (Fig. 3 C); neither R1 nor R2 alone bound or bundled MTs. Alanine substitution of the basic residues in two or all three of the R1 motifs resulted in an ~50% decrease in MT binding (Fig. 3 D and Fig. S2 D). A similar decrease was observed by mutating the basic residues of the R2 motifs (Fig. 3 D and Fig. S2 E). We assessed the effects of the same mutations in MT bundling using a low speed sedimentation assay. Strikingly, alanine substitutions of the basic residues of the R-R-x-E motifs of R2 decreased MT bundling, but mutating the basic residues of the K/R-x-x-E/D motifs of R1 did not decrease MT bundling (Fig. 3 E; Fig. S2, F and G). These data indicate that although the basic residues of both K/R-x-x-E/D and R-R-x-E contribute to MT binding, the arginine residues of the R-R-x-E motifs are uniquely critical for MT bundling.

Next, we assessed the role of the acidic residues. Alanine substitutions of the acidic residues in one or two of the motifs of either R1 or R2 resulted in a moderate (~30%) increase of MT-SEPT9_i1(61–113) binding (Fig. 3 F and Fig. S2 H). MT bundling, however, decreased by ~50% when the acidic residues in two of the K/R-x-x-E/D motifs of the R1 region were mutated (Fig. 3 G and Fig. S2 I). In contrast, alanine substitutions of glutamate in the R-R-x-E motifs of the R2 region resulted in only a marginal decrease in MT bundling (Fig. 3 G and Fig. S2 I). These results show that although the acidic residues dampen somewhat the electrostatic affinity of all motifs for the CTTs of β-tubulin, the acidic residues of the K/R-x-x-E/D motifs (R1)
Figure 1. SEPT9 binds and bundles MTs via a basic N-terminal domain. (A) Sequence and domains of SEPT9_i1. (B–F) Coomassie-stained SDS-PAGE gels of the supernatant (S) and pellet (P) fractions after high speed (39,000 g) sedimentation of pre-polymerized paclitaxel-stabilized MTs with domains of SEPT9_i1. (G–I) Low speed (8,000 g) sedimentation of MTs in the presence of SEPT9_i1 domains. (J and K) Graphs show percentages of total protein pelleted with MTs at 39,000 g (J) and percentage of total MTs pelleted at 8,000 g (K) in three independent experiments. (L) Images show X-rhodamine–labeled MTs after mixing with recombinant SEPT9 fragments. Bars, 10 µm. (M) Negative stain EM images of MTs before and after mixing with SEPT9_i1. Arrows and arrowheads point to MT bundles and doublets, respectively. (N and O) Plots show the intensity of X-rhodamine fluorescence per micron of MT (O; n = 100) and the length of MTs (P; n = 100) per condition.
Figure 2. SEPT9 interacts with the acidic C-terminal tails of β-tubulin. (A) Gel shows supernatant (S) and pellet (P) fractions after sedimentation (39,000 g) of SEPT9-FL with untreated and subtilisin-treated MTs (S-MTs). Graph shows percentage of total SEPT9-FL in the S and P fractions. (B) Images show untreated and subtilisin-treated X-rhodamine–labeled MTs after mixing with SEPT9-FL. Bars, 10 µm. (C and D) Plots show the fluorescence intensity per micron of MT (C; n = 50) and the length of MTs (D; n = 50). (E and F) Graph shows the fluorescence intensity per micron of MT (C; n = 50) and the length of MTs (D; n = 50). (E and F) Graph shows fraction of SEPT9-FL pelleted with MTs in the presence of increasing concentrations of α-tubulin, βIII-tubulin, and scrambled CTT peptide relative to no peptide in three independent experiments. Gels show the pellet fractions after MT pelleting with SEPT9-FL in the presence of increasing concentrations of peptides. (G) Gels show the pellet fractions after sedimentation of MTs with SEPT9-FL in the presence of βI, βII, and βIII-tubulin CTT peptides. Graph shows the fraction of SEPT9-FL pelleted in the presence of peptides relative to no peptide in three independent experiments. (H) Increasing amounts of bovine brain tubulin was separated by 7.5% SDS-PAGE and transferred to nitrocellulose membranes, which were stained with Ponceau S red. Membranes were blotted with DM1A and TUB2.1 antibodies against α- and β-tubulin, respectively, and overlaid with His-tagged SEPT9-FL and SEPT2, which were detected with anti-His antibodies.
Novel SEPT9 repeat motifs bundle microtubules

We hypothesized that SEPT9 isoforms that lack K/R-R/x-x-E/D motifs from their N terminus could affect MT bundling and asymmetric neurite growth. Among the known SEPT9 protein isoforms, SEPT9_i4 lacks the N-terminal 164 aa and therefore most of the MT-binding and -bundling repeat motifs of SEPT9_i1. To test if the absence of these motifs affects intracellular MT bundling, we expressed GFP-tagged chimeras of these isoforms in MDCK cells, in which septins were previously shown to colocalize with MTs (Spiliotis et al., 2008; Bowen et al., 2011). We found that SEPT9_i1 colocalized strongly with perinuclear MT bundles, whereas SEPT9_i4 was distributed at peripheral lamellae showing little to no colocalization with MTs (Fig. 4, A and C).

Upon treatment with paclitaxel, an MT-stabilizing drug that induces MT bundling, SEPT9_i1 colocalized with MT bundles extensively, whereas SEPT9_i4 showed little colocalization with MTs (Fig. 4, B and C). Interestingly, paclitaxel-treated cells that expressed SEPT9_i4 cells appeared to have less number of long and straight MT bundles (Fig. 4 B). This effect was quantified by measuring the relative percentage of MTs with fluorescence intensities 5x or 10x greater than the mean intensity of the are uniquely critical for MT bundling. Because the basic residues of the R-R-x-E motif are also uniquely involved in MT bundling, we posit that the extra arginine residues of the R-R-x-E motifs allow for additional interactions with the acidic residues of the K/R-x-x-E/D motifs, enabling homophilic trans interactions between SEPT9_i1(61–113) peptides that cross-link MTs into bundles (Fig. 3 H). Consistent with this model, a GST-tagged version of the N terminus of SEPT9 bound His-tagged SEPT9-N and SEPT9-FL (Fig. S2, J and K). Thus, homophilic trans interactions between the N termini of SEPT9 are likely to facilitate septin and MT cross-linking.

SEPT9 repeat motifs are required for intracellular MT bundling and the generation of neurite asymmetry

Given the mechanistic role of the K/R-R-x-x-E/D motifs in MT binding and bundling in vitro, we examined the functional significance of these motifs for intracellular MT bundling and neuronal morphogenesis, which requires MT bundling by MAPs (e.g., tau, MAP1B) that support the asymmetric growth of neuronal protrusions termed neurites (Caceres and Kosik, 1990; Teng et al., 2001; Feltrin et al., 2012). We hypothesized that SEPT9 isoforms that lack K/R-R-x-x-E/D motifs from their N terminus could affect MT bundling and asymmetric neurite growth. Among the known SEPT9 protein isoforms, SEPT9_i4 lacks the N-terminal 164 aa and therefore most of the MT-binding and -bundling repeat motifs of SEPT9_i1. To test if the absence of these motifs affects intracellular MT bundling, we expressed GFP-tagged chimeras of these isoforms in MDCK cells, in which septins were previously shown to colocalize with MTs (Spiliotis et al., 2008; Bowen et al., 2011). We found that SEPT9_i1 colocalized strongly with perinuclear MT bundles, whereas SEPT9_i4 was distributed at peripheral lamellae showing little to no colocalization with MTs (Fig. 4, A and C). Upon treatment with paclitaxel, an MT-stabilizing drug that induces MT bundling, SEPT9_i1 colocalized with MT bundles extensively, whereas SEPT9_i4 showed little colocalization with MTs (Fig. 4, B and C). Interestingly, paclitaxel-treated cells that expressed SEPT9_i4 cells appeared to have less number of long and straight MT bundles (Fig. 4 B). This effect was quantified by measuring the relative percentage of MTs with fluorescence intensities 5x or 10x greater than the mean intensity of the

Figure 3. MT bundling by novel SEPT9 repeat motifs K/R-x-x-E/D and R/K-R-x-E. (A and B) WebLogo alignments of eleven hexapeptide sequences containing the K/R-x-x-E/D motif (A) and six sequences containing the R/K-R-x-E motif (B) within the N terminus [aa 1–286] of SEPT9_i1. The height of each residue indicates the frequency of its presence at the indicated positions. (C) Sequence of aa 61–113 of SEPT9_i1. The blue and red stars highlight the basic and acidic residues mutated to alanine. (D and E) Graphs show percentage of total SEPT9_i1(61–113) [wild-type and basic residue mutants] pelleted with MTs at 39,000 g (D), and percentage of total tubulin pelleted at 8,000 g (E) in three independent experiments. Representative gels are shown in Fig. S2. (F and G) Graphs show percentage of total 61–113 peptide [wild-type and acidic residue mutants] pelleted with MTs at 39,000 g (F), and percentage of total tubulin pelleted at 8,000 g (G) in three independent experiments. Representative gels are shown in Fig. S2. (H) Schematic shows a model of electrostatic interactions between the acidic (red) CTTs of tubulin and the basic (blue) residues of the SEPT9 repeat motifs. MT cross-linking is achieved by interactions between the acidic and basic residues of the K/R-x-x-E/D and R/K-R-x-E motifs, respectively.
Figure 4. **SEPT9 repeat motifs are required for MT bundling and asymmetric neurite growth.** (A and B) Maximal projections of 3D confocal microscopy images of MDCK cells expressing GFP-tagged SEPT9_i1 and SEPT9_i4 before (A) and after (B) treatment with 5 µM paclitaxel for 1.5 h. (C) Manders coefficients for the colocalization of GFP-tagged SEPT9_i1 and SEPT9_i4 with MTs in MDCK cells (n = 15). High and low colocalization are indicated by coefficients >0.5 and <0.5, respectively. (D and E) Plots show the fluorescence intensity of putative MT bundles with 5× and 10× the mean intensity of single MTs as percentage of total MT intensity in MDCK cells (n = 15) before (D) or after (E) treatment with paclitaxel. (F) Phase-contrast images show PC12 cells transfected with GFP and GFP-tagged SEPT9_i1 and SEPT9_i4 after a 2- and 3-d NGF treatment. Insets show GFP fluorescence in inverted monochrome. Bars, 10 µm. (G) Graph shows percentage of PC12 cells (n = 90) with one or more neurites. Pooled data from three independent experiments are shown.
dimmest single peripheral MTs, using a segmentation analysis of 3D confocal image stacks as previously published (Sammak and Borisy, 1988; Bowen et al., 2011). MDCK cells that expressed SEPT9_i4-GFP had consistently lower percentage of 5× or 10× MTs compared with control and SEPT9_i1-GFP in the presence or absence of paclitaxel (Fig. 4, D and E). A lower percentage of MT bundles was also observed in fibroblasts from Sept9 knockout mice at steady-state and after treatment with paclitaxel (Fig. S3). Thus, SEPT9 is critical for the intracellular organization of MTs into bundles.

During neuronal morphogenesis, MT bundling supports the asymmetric growth of a neurite, which extends faster and longer than other neurites to become the presumptive axon. To test whether SEPT9_i1 and the K/R-R-x-x-E/D motifs affect neurite growth, we assayed for NGF-induced neurite formation in PC12 cells. After 48 and 72 h of NGF treatment, the percentage of GFP-SEPT9_i1-expressing cells with single neurites was respectively fivefold and twofold higher than control cells (Fig. 4, F and G). In contrast to this amplification of asymmetric neurite growth by SEPT9_i1, expression of SEPT9_i4 had the opposite effect, increasing the percentage of cells with more than three neurites by twofold after 48 h of NGF treatment (Fig. 4, F and G). Notably, after 72 h of NGF exposure, the percentage of SEPT9_i4-expressing cells with single neurites was twofold lower than control cells (Fig. 4 G) and the average neurite length was shorter than the length of single neurites in SEPT9_i1-expressing cells (2.3 vs. 3.3 times the cell width). These results indicate that the MT-bundling properties of SEPT9 promote asymmetric neurite growth, which depends on the presence of K/R-R-x-x-E/D motifs within the N-terminal sequence of SEPT9.

HNA-linked mutation R88W impairs MT bundling and asymmetric neurite growth

Alterations in the N-terminal sequence of SEPT9 have been genetically linked to HNA, a neuralgic disorder whose molecular etiology remains unknown (Kuhlenbäumer et al., 2005; Hannibal et al., 2009; van Alfen, 2011). Several HNA patients carry the mutation R88W in isoform 3 of SEPT9 (Hannibal et al., 2009), which has one K/R-x-x-E/D motif less than SEPT9_i1 and contains seven unique residues in place of aa 1–25 of SEPT9_i1. Because the R88W mutation of SEPT9_i3 maps to the first arginine of an R-R-x-E motif, we asked whether this mutation affects the MT-binding and -bundling properties of SEPT9_i3. The missense mutation did not affect high speed pelleting of SEPT9_i3 with MTs (Fig. 5, A and C), but low speed sedimentation of MTs decreased by ∼30% (Fig. 5, B and D). In fluorescent MT-bundling assays, the R88W mutation weakened the MT-bundling effects of SEPT9_i3 (Fig. 5, E–G). To determine if the R88W mutation alters MT bundling in vivo, we obtained dermal fibroblasts from healthy control individuals and HNA patients with the R88W genotype. We found that the fraction of MTs with 5× the mean intensity of single MTs was lower in the HNA patient fibroblasts compared with cells from a healthy individual (Fig. 5 H). Given the role of SEPT9_i1 and its repeat motifs in asymmetric neurite growth, we tested how the wild-type and R88W mutant versions of SEPT9_i3 affect neurite outgrowth in PC12 cells. Expression of GFP-SEPT9_i3 increased the percentage of cells with a single neurite by 50% relative to GFP-expressing cells (Fig. 5 I). The R88W mutant, however, increased the overall percentage of cells with more than three neurites (Fig. 5 I). Taken together, these data show that the HNA-linked R88W mutation has adverse effects on MT bundling and asymmetric neurite growth.

Our results provide the first insight into the mechanism of septin interaction with MTs. We have shown that the N terminus of SEPT9 binds and bundles MTs via repeat motifs similar to those of conventional MAPs (Noble et al., 1989; Butner and Kirschner, 1991; Cravchik et al., 1994; Ferralli et al., 1994). Formation of uniform linear arrays of bundled MTs is likely to involve highly ordered cis and trans interactions between SEPT9 containing hetero-oligomers. Cis interactions between the GTP-binding domains of septins mediate their assembly into hetero-oligomeric filaments (Sirajuddin et al., 2007). Conversely, trans interactions between the C-terminal coiled-coil domains of septins mediate the pairing of septin filaments (John et al., 2007; Bertin et al., 2008; DeMay et al., 2011; de Almeida Marques et al., 2012). SEPT9 molecules lack C-terminal tails, but our data indicate that their N-terminal extensions could mediate trans interactions by electrostatic pairing between the acidic and basic residues of the K/R-x-x-E/D and R/K-R-x-E motifs, respectively. SEPT9-mediated MT cross-linking could be further reinforced by the trans interactions between the C-terminal coiled-coil domains of SEPT2, SEPT6, and SEPT7, which hetero-oligomerize with SEPT9 (Kim et al., 2011; Sellin et al., 2011). Although SEPT2 does not bind MTs (Fig. S1), previous work indicates that SEPT7 and possibly SEPT6 interact with MTs (Hu et al., 2012; Moon et al., 2013). These subunits, however, lack the repeat motifs of SEPT9.

Septins colocalize with MTs in various cell types, but the physiological significance of this interaction is poorly understood. Here, we have found that SEPT9 and its N-terminal repeat motifs affect intracellular MT bundling and asymmetric neurite growth. Importantly, expression of SEPT9 isoforms with deletions (SEPT9_i4) and mutations (HNA-linked R88W) in the repeated K/R-R-x-x-E/D sequence motifs impair MT bundling and asymmetric neurite growth. Interestingly, cancer cells that overexpress SEPT9_i4 have increased resistance to the drug paclitaxel (Chacko et al., 2012). It is unknown if paclitaxel-mediated MT bundling contributes to the cytotoxicity of the drug, but our data indicate that SEPT9 and its repeat motifs are partly required for paclitaxel-induced MT bundling. This role of SEPT9, however, could vary between different cell types. Given the preferential interaction of SEPT9 with β-tubulin, variations in the expression of β-tubulin isoforms could affect septin association with MTs. In summary, our results suggest that alterations in SEPT9 isoform expression trigger changes in the intracellular organization and function of MTs, and thereby could contribute to the pathology of HNA and cancer.

Materials and methods

Cells, peptides, and plasmids

MDCK/II/G cells were maintained in low glucose DME media (Sigma-Aldrich) supplemented with 10% FBS (Cell Generation) and 1 g/liter NaHCO₃. MDCK cells were transfected with the plasmids pEGFP-SEPT9_v1.
Industries) was used to take a skin sample from the extensor surface of the upper arm. After removal of the subcutaneous fatty tissue, skin explants were cultured in high glucose DME media with 20% FBS for 5–7 d. Skin fibroblasts that grew out of the explants were isolated and passaged. Experiments with R88W and control 1 fibroblasts were performed at passages 7–11 and 9–13, respectively. The peptides NH$_2$-CEVGVDSVEGEGEEE-GEEY-COOH ($\alpha$-tubulin CTT), NH$_2$-CQETAEEYQDEEQGEADAEDFG-COOH (control, scrambled $\beta$II-tubulin CTT), and NH$_2$-CQYQDATAEDEQGEFEEEEGEDEA-COOH ($\beta$I tubulin CTT) were purchased at >95% purity from GenScript. The peptides NH$_2$-CQYQDATADEQGEFEEEEGEDEA-COOH ($\beta$II-tubulin CTT) and NH$_2$-CQYQDATAEEEEGEMYEDDEEESEAAQPK-COOH ($\beta$III-tubulin CTT) were purchased at >95% purity from LifeTein, LLC.

His-SEPT9$_{i1}$-expressing plasmid was constructed by PCR amplifying human SEPT9$_{i1}$ (NP_001106963) using the primers 5’-CGTAAGCTTG-CATGAAGAAGTCTTACTC-3’ and 5’-GTACTCGAGCTACATCTCTGGGGCC-3’ and cloning the amplified fragment into the HindIII and XhoI sites of pEGFP-SEPT9$_{v4}$, which encode respectively for GFP-tagged SEPT9$_{i1}$ and SEPT9$_{i4}$ and were constructed by PCR amplification of SEPT9$_{v1}$ (AF189713) and SEPT9$_{v4}$ (AJ312322) from normal breast tissue cDNA and insertion into pEGFP-C2 (Connolly et al., 2011b). PC-12 cells were maintained in high glucose DME supplemented with 6% donor-defined equine serum (Hyclone), 6% defined bovine calf serum (Hyclone), and 1 g/liter NaHCO$_3$. PC-12 differentiation and neurite growth were induced after 24 h of transfection by incubation in low serum (1% horse serum, 1% bovine calf serum) DME containing 100 ng/ml 2.5S NGF (Harlan Biosciences). Embryonic fibroblasts derived from Sept9$^{cond/del}$ mice were provided by E.-M. Füchtbauer (University of Aarhus, Aarhus, Denmark) and maintained in DME with 10% FBS as described previously (Füchtbauer et al., 2011). Dermal cells were derived from skin biopsies taken from a healthy individual (control 1) and HNA patient with the R88W genotype stained for $\alpha$-tubulin. Bars, 10 µm.

Figure 5. HNA-linked mutation R88W impairs MT bundling and neurite asymmetry. (A and B) Gels show supernatant (S) and pellet (P) fractions from high (A; 39,000 g) and low (B; 8,000 g) speed MT-pelleting assays. (C and D) Graphs show percentages of total protein co-pelleted with MTs (C) and percentage of total tubulin pelleted (D) in three independent experiments. (E) Images show X-hodamine–labeled MTs with recombinant SEPT9$_{i3}$ and SEPT9$_{i3}(R88W)$. Bars, 10 µm. (F and G) Plots show the intensity of X-hodamine fluorescence per micron of MT (F; n = 100) and the length of MTs (G; n = 100) per condition. (H) Images of dermal cells from healthy individual (control 1) and HNA patient with the R88W genotype stained for $\alpha$-tubulin. Bars, 10 µm. (I) Phase-contrast images show PC12 cells transfected with GFP and GFP-tagged wild-type and R88W SEPT9$_{i3}$. Insets show GFP fluorescence in inverted monochrome. Graph shows percentage of cells (n = 90) with one or more neurites. Pooled data from three independent experiments are shown. Bars, 10 µm.
Expression and purification of recombinant proteins

Plasmids encoding for recombinant proteins were transformed into Escherichia coli BL21 (DE3) (Invitrogen). After bacterial cultures reached an O.D.600 of 0.8, protein expression was induced with 0.5 mM IPTG for 16 h at 18°C. Bacteria were centrifuged at 5,000 rpm for 5 min at 4°C. Cells were resuspended in buffer containing 1% Triton X-100, 50 mM Tris, pH 8.0, 150 mM NaCl, 10% glycerol, and 10 mM imidazole, and lysed using a French pressure cell (1,280 psi). Cell lysates were clarified by centrifugation at 14,000 g for 30 min. Protein expression was induced with 0.5 mM IPTG for 16 h at 18°C. Bacteria were centrifuged at 5,000 rpm for 5 min at 4°C. Pellets were resuspended in buffer containing 1% Triton X-100, 50 mM Tris, pH 8.0, 150 mM NaCl, 10% glycerol, and 10 mM imidazole, and lysed using a French pressure cell (1,280 psi). Cell lysates were clarified by centrifugation at 14,000 g for 30 min. Supernatant and pellets were treated for 5 min. In high speed sedimentation assays, untreated, mock, or subtilisin-treated MTs were incubated with recombinant SEPT9 proteins (10 µM) or SEPT9(61–113) peptides (30 µM) for 20 min at room temperature. Each reaction was placed on cushion buffer and centrifuged at 39,000 g (high speed) for 20 min at 25°C in an ultracentrifuge (Optima TL100, Beckman Coulter). In low speed sedimentation assays, MTs were incubated with recombinant SEPT9 proteins (2 µM), SEPT9(61–113) peptides (30 µM), or SEPT9(62/67, SEPT9(61–113) complexes and SEPT9(61–113) complexes (0.2 µM; Fig. S1, J and K) for 10 min at room temperature. Each reaction was placed on cushion buffer and centrifuged at 8,000 g for 5 min. Pellets were resuspended in PBS of the same volume with the supernatant. Equal volumes of pellet and supernatant were loaded onto 10% SDS-PAGE and gels were stained with Coomassie Brilliant Blue. Gels were scanned and protein band densities were quantified with the Odyssey infrared scanning system (LI-COR Biosciences).

Visualization of MT bundling by fluorescence and electron microscopy

X-hodamine-labeled bovine brain tubulin (10 µM; Cytoskeleton, Inc.) was polymerized in G-PEM buffer containing 80 µM paclitaxel. MTs were incubated with recombinant proteins (20 nM) in G-PEM buffer containing 20 µM paclitaxel for 10 min at room temperature. An aliquot (5 µl) of each reaction mix was mounted on a slide and sealed with a glass coverslip and nail polish. Slides were imaged on a fluorescent microscope (Axio Observer. Carl Zeiss) equipped with a Plan-Apo 63×/1.40 NA objective and a deep-cooled CCD camera (ORCA-AG; Hamamatsu Photonics), and Slidebook 5.0 software (Intelligent Imaging Innovations). All images were taken in the TRITC channel at 50-s exposures. MT lengths and intensities were quantified by masking individual MTs (>5 µm). MT length and average fluorescence intensity per pixel after background subtraction were measured using Slidebook 5.0 software.

For visualization of MT bundling by negative stain electron microscopy, pre-polymerized MTs (Cytoskeleton, Inc.) were mixed with recombinant SEPT9-F9 and incubated with glow-discharged 300 mesh holey carbon copper grids covered with a thin layer of carbon. Grids were washed with 2% uranyl formate and subsequently blotted and air dried. Images were collected on a transmission electron microscope (Tecnai 11G Spirit; FEI) equipped with a CCM-300 300-kV microscope (UltraScan 4000; Gatan, Inc.) operating at accelerating voltage of 120 kV, defocus value of ~950 nm, and magnification of 52,000.

Western blots and overlay assays

Bovine brain tubulin (>99% pure; Cytoskeleton, Inc.) was separated by 7.5% SDS-PAGE and transferred to PROTRAN B85 nitrocellulose membrane (Whatman), which were stained with Poncova 5 (Sigma-Aldrich) and scanned on a Canon LIDE 200 (Canon). After electrotransfer, grids were blocked for 1 h at 4°C in blocking buffer (10 mM Tris HCl, pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.1% Tween 20, 5% nonfat dry milk, and 0.05% BSA). Subsequently, membranes were overlayed with His-SEPT9 antibodies.
(200 nM) in blocking buffer for 2 h. Membranes were washed with TBS containing 0.1% Tween 2D and incubated in the same buffer containing 2% BSA and mouse antibody against 6×His tag (1:5,000, R&D Systems) followed by secondary Alexa Fluor 680 goat anti-mouse IgG (1:15,000; Invitrogen). Blots were imaged with the Odyssey infrared imaging system (LI-COR Biosciences).

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