Crescerin uses a TOG domain array to regulate microtubules in the primary cilium

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ABSTRACT Eukaryotic cilia are cell-surface projections critical for sensing the extracellular environment. Defects in cilia structure and function result in a broad range of developmental and sensory disorders. However, mechanisms that regulate the microtubule (MT)-based scaffold forming the cilia core are poorly understood. TOG domain–containing proteins ch-TOG and CLASP are key regulators of cytoplasmic MTs. Whether TOG array proteins also regulate ciliary MTs is unknown. Here we identify the conserved Crescerin protein family as a cilia-specific, TOG array-containing MT regulator. We present the crystal structure of mammalian Crescerin1 TOG2, revealing a canonical TOG fold with conserved tubulin-binding determinants. Crescerin1’s TOG domains possess inherent MT-binding activity and promote MT polymerization in vitro. Using Cas9-triggered homologous recombination in Caenorhabditis elegans, we demonstrate that the worm Crescerin family member CHE-12 requires TOG domain–dependent tubulin-binding activity for sensory cilia development. Thus, Crescerin expands the TOG domain array–based MT regulatory paradigm beyond ch-TOG and CLASP, representing a distinct regulator of cilia structure.

INTRODUCTION

Eukaryotic cilia and flagella are specialized microtubule (MT)-based organelles. Motile cilia are classically associated with cellular movement or direction of extracellular fluid flow (Satir and Christensen, 2008). In addition, most mammalian cell types contain a primary cilium—a solitary, antenna-like extension that receives extracellular chemical and mechanical stimuli and transduces a signal to the cell body to elicit a response (Marshall and Nonaka, 2006; Satir and Christensen, 2007; Christensen et al., 2008; Veland et al., 2009; Hoey et al., 2012). Genetic mutations in ciliary proteins underlie diseases collectively termed ciliopathies. In many of these diseases, the structural integrity of the primary cilium is compromised (Marshall, 2008).

The structural core of the cilium and flagellum is the axoneme, composed of parallel MTs arranged in ninefold radial symmetry. The axonemal MTs are polymers of αβ-tubulin heterodimers. Nine axonemal doublet MTs arise from the basal body, a modified centriole docked at the plasma membrane, which templates the radial MT symmetry and anchors the MT minus ends. The MT doublets, composed of A and B tubules, span the length of the cilium middle zone. At the middle/distal zone transition, B tubules terminate, and singlet A tubules extend into the distal zone. The axonemal MTs provide structural support, serve as tracks along which anterograde and retrograde intraflagellar transport complexes traffic ciliary components, and, in motile cilia, serve as the substrate for axonemal dynein motors that drive motility (Scholey, 2003, 2008; Ishikawa, 2012; Kobayashi and Takeda, 2012). Doublet MTs in the middle zone are stable and resistant to MT-depolymerizing agents, including colchicine (Tilney and Gibbins, 1968). In contrast, singlet MTs in the distal segment are dynamic and change their length in response to external stimuli (Marshall and Rosenbaum, 2001; Mukhopadhyay et al., 2008; Rich and Clark, 2012). The plus ends of both the A and B tubules exhibit rapid tubulin turnover (Hao et al., 2011), and this localized dynamism is necessary to maintain proper...
axoneme structure. However, the identities and mechanisms of MT-associated proteins (MAPs) involved in regulating ciliary MT organization and dynamics are not well understood.

Whereas MT polymerization–dependent GTP hydrolysis confers the αβ-tubulin polymer with inherent dynamics, a host of cytoplasmic MAPs regulate MT dynamics in space and time to facilitate dynamic cellular changes as required during mitosis, polarized cell migration, and ciliogenesis. Colonic and hepatic tumor overexpressed gene (ch-TOG) and cytoskeletal linker protein 170–associated protein (CLASP) represent two distinct MAP families that promote MT polymerization and pause, respectively (Akmanova et al., 2001; Brittle and Ohkura, 2005; Drabek et al., 2006; Sousa et al., 2007; Widlund et al., 2011). Both ch-TOG and CLASP use arrayed tubulin-binding TOG domains to regulate MT dynamics (Al-Bassam et al., 2007; Al-Bassam and Chang, 2011; Slep, 2009). The number of TOG domains in each array differs (Figure 1A), as do the architecture and tubulin-binding activity of each TOG domain. Higher eukaryotic ch-TOG family members use a pentameric TOG domain array to promote MT polymerization (Widlund et al., 2011), whereas CLASP members use a trimeric TOG domain array to stabilize MTs and promote MT pause (Al-Bassam et al., 2010; Patel et al., 2012).

A TOG domain consists of six pairs of antiparallel HEAT loops that buttress the tubulin-binding determinants have diverged in sequence, both within a given TOG array and across the ch-TOG and CLASP families, making it challenging to identify TOG domain–containing proteins using BLAST searches. Using structure-based methodology, we previously predicted and confirmed the existence of a cryptic TOG domain (TOG2) in CLASP (Slep and Vale, 2007; Leano et al., 2013), establishing a TOG array–based paradigm for ch-TOG and CLASP family MT regulatory activity. Here we asked whether we could identify another TOG domain–containing protein family in eukaryotes based on our knowledge of TOG structural features and
RESULTS

Crescerin defines a conserved protein family with a predicted TOG domain array

Given that the ch-TOG and CLASP families play central roles regulating MT dynamics, we asked whether any other conserved protein families might use a TOG domain array to regulate MT dynamics. We performed structure-based BLAST searches using individual TOG domains from ch-TOG and CLASP family members and scanned weak hits for sequence identity in the discontinuous TOG domain intra-HEAT loop determinants that bind tubulin. Searching the human proteome, we were surprised to find an uncharacterized protein, FAM179B (KIAA0423), that showed discontinuous sequence similarity to CLASP TOG1 (BLAST E-value of 1e-09). This novel protein is a member of a protein family conserved in ciliated/flagellated eukaryotes ranging from mammals to unicellular eukaryotes, including Chlamydomonas (SSA6) and Tetrahymena, but is absent in nonciliated eukaryotes, including yeast and Dictyostelium (Figure 1A and Supplemental Table S1). We designate this protein family and its unnamed members Crescerin. The only family member characterized is the C. elegans member CHE-12. The single study examining CHE-12 showed that it is expressed in a subset of amphid neurons possessing simple rod-like cilia and two phasmid sensory neurons under the transcriptional control of DAF-19, a transcription factor that controls numerous ciliary genes (Bacaj et al., 2008). Overexpressed CHE-12::green fluorescent protein (GFP) localizes to cilia, and mutations in che-12 impair cilia formation (Bacaj et al., 2008). Concordant with a role in cilia structure, che-12 mutant worms do not properly chemotax in response to sodium chloride (Bacaj et al., 2008).

On further investigation, we found that mammals contain two closely related Crescerin family genes, FAM179A and FAM179B (Figure 1A). Sequence analysis and secondary structure prediction of mouse FAM179B revealed four conserved domains—an N-terminal pair and a C-terminal pair—connected by a long central linker (Figure 1A). Each of the four domains contains 12 predicted α-helices that are predicted to form six tandem HRs (Supplemental Figures S1 and S2) with intra-HEAT loops displaying similarity to the intra-HEAT loop tubulin-binding determinants found in ch-TOG and CLASP family TOG domains. The mammalian FAM179A protein contains only two conserved TOG-like domains, which share high sequence similarity with the FAM179B C-terminal pair of TOG domains. A number of Crescerin family members from different species were found to contain fewer than four TOG domains. We classified all proteins that include TOG-like domains similar to the mammalian FAM179B TOG1 or TOG2 into the Crescerin1 subfamily and those that only contain FAM179B TOG3 and TOG4-like domains (i.e., more similar to FAM179A) into the Crescerin2 subfamily (Figure 1A and Supplemental Table S1). Dendrogram analysis of the TOG and TOG-like domains from the ch-TOG, CLASP, and Crescerin families suggests that the predicted Crescerin1 TOG-like domains TOG2, TOG3, and TOG4 are most similar to CLASP family TOG domains TOG1, TOG2, and TOG3, respectively (Figure 1B). In contrast, Crescerin1 TOG-like domain TOG1 is not closely related to TOG domains from either the CLASP or the ch-TOG family and is the least conserved TOG domain across the Crescerin family. We sought to determine whether the Crescerin protein family represents a third TOG array-containing protein family and plays a role distinct from the ch-TOG and CLASP protein families potentially specific to cilia structure and function.

Crescerin1 localizes to cilia in C. elegans amphid and phasmid neurons and mammalian cells

The C. elegans Crescerin homologue CHE-12 was previously reported to be required for the structure and function of cilia in the amphid neurons, a subset of sensory neurons in the head (Perkins et al., 1986; Bacaj et al., 2008). Previous studies used extrachromosomal arrays to show that the che-12 promoter is active in amphid neurons and phasmid neurons, that overexpressed CHE-12 localizes to cilia, and that this localization was dependent on intraflagellar transport (IFT) particle B (Bacaj et al., 2008). To characterize endogenous CHE-12 expression and localization, we used Cas9-triggered homologous recombination (Dickinson et al., 2013) to insert gfp into the che-12 locus, generating GFP::CHE-12 and CHE-12::GFP fusion proteins expressed under the control of all native regulatory elements (Figure 2, Supplemental Figure S3, and Supplemental Table S2). We additionally expressed mCherry under control of the gcy-5 promoter to label the ASER neuron, an amphid neuron with a single, rod-like cilium (Figure 2A). GFP::CHE-12 and CHE-12::GFP both localized to multiple sensory cilia in the head, including the bundle of parallel amphid channel cilia (Figure 2, B and C). CHE-12 was also expressed in the phasmid cilia in the tail region, where it localized along the phasmid cilia with slight enrichment near the cilia base (Figure 2D). CHE-12 GFP constructs showed punctate localization along the amphid and phasmid cilia, which may reflect association with IFT particles.

The localization of CHE-12 to amphid and phasmid neuronal cilia in C. elegans parallels the localization of endogenous Crescerin1 at the primary cilium and basal body in mammalian cells, as observed by staining mouse IMCD3 cells with an antibody we raised against mouse Crescerin1 (Figure 3A), suggesting a conserved role for the Crescerin family in cilia structure and/or function. In addition, GFP-tagged, full-length mouse Crescerin1 also localized to primary cilia in human HEK-293 cells (Figure 3, B and C). Crescerin1-GFP localized along the length of the primary cilium and at the basal body at low expression levels (Figure 3B). At high expression, Crescerin1-GFP accumulated at the basal body and distal tip of primary cilium, as well as along cytoplasmic MTs (Figure 3C). Cells were fixed and stained with anti–acetylated tubulin antibody (red) to label cilia and anti-pericentrin antibody (blue) to label basal bodies and centrosomes. Localization of overexpressed Crescerin1-GFP to the distal cilia tip may imply coupling to IFT machinery. The potential for this is supported by the punctate localization we observed for CHE-12 GFP constructs along the amphid and phasmid cilia, as well as previous work that found CHE-12 localization to depend on IFT particle B (Bacaj et al., 2008). We never observed Crescerin1-GFP at the plus ends of cytoplasmic MTs, suggesting that Crescerin1 does not preferentially localize to the distal cilia tip through recognition of a GTP-MT cap.

Structure of the Crescerin1 second conserved domain reveals a bona fide TOG domain

To determine whether the Crescerin family represents a novel TOG domain array-containing protein family, we cloned and purified all four conserved putative TOG domains from mouse Crescerin1. We obtained crystals of the second conserved domain (residues 332–620) in space group P2₁2₁2₁. We collected a native x-ray diffraction
Crescerin TOG array regulates cilia MTs

data set to 2.2-Å resolution and a single-wavelength anomalous dispersion data set on selenomethionine-substituted protein to 2.37-Å resolution. We solved and built the structure, refining against the native data to 2.2-Å resolution, yielding R and Rfree values of 0.18 and 0.24, respectively (Supplemental Table S3). The model contains one molecule of Crescerin1 in the asymmetric unit and encompasses residues 345–594.

The Crescerin1 TOG2 structure reveals a bona fide TOG domain fold consisting of 12 helices that pair into tandem HRs (A–F) and stack, forming a 60-Å oblong solenoid (Figure 4A). TOG domain HRs can be grouped into two triads: the first triad contains HRs A–C, and the second triad contains HRs D–F. These HR triads have a canonical handedness and relative positioning similar to that observed in ch-TOG family TOG domains (Al-Bassam et al., 2007; Slep and Vale, 2007; Fox et al., 2014). The HR A–C triad has a right-handed twist, whereas the HR D–F triad has a right-handed twist between repeats D and E but transits to a left-handed twist between repeats E and F. Between the two HR triads, there is a lateral shift relative to the axis of the solenoid, a common feature found in all other TOG domain structures determined to date. The Crescerin1 TOG2 structure has inter-HR α-helices α2β1, situated between HR A and HR B, also found in hCLASP1 TOG2, and α2E1, situated between HR D and HR E, also found in the hCLASP1 TOG2 structure, as well as in ch-TOG family structures of Drosophila Msps TOG2 and TOG4. To map conserved residues across the Crescerin family onto the structure, we aligned 13 diverse species and contoured conservation at 75% identity (green) and 75% similarity (yellow, Figure 4B and Supplemental Figure S2). The majority of conserved residues map to the intra-HEAT loops that span one face of the domain. These conserved intra-HEAT loops are similar in composition and structure to the tubulin-binding intra-HEAT loops found in ch-TOG and CLASP protein family TOG domain structures (Figure 4C and Supplemental Figure S2). In addition, the electrostatic charge distribution across the first five intra-HEAT loops (HR A–E) is basic (Figure 4B) and, like other TOG domain structures determined to date, would complement the acidic nature of the αβ-tubulin TOG-binding site. Overall, the structure of the Crescerin1 second conserved domain conforms to a canonical TOG architecture with conserved intra-HEAT loop determinants, homologous in composition and charge to the tubulin-binding determinants in ch-TOG and CLASP TOG domains. We thus designate this domain a TOG domain and refer to it hereafter as TOG2. This confirms our prediction that Crescerin represents a third, unique TOG domain–containing protein family.

Crescerin1 TOG2 has a unique C-terminal β-hairpin that promotes domain stability

Although Crescerin1 TOG2 has overall structural similarity to previously determined ch-TOG and CLASP family TOG domain structures, it also has architecturally unique components that differentiate it from these other TOG domain structures. One striking feature is the presence of a 21-residue C-terminal β-hairpin that runs alongside the second HR triad and occupies the area created by the lateral shift between the HR A–C and D–F triads. The β-hairpin forms multiple hydrophobic interactions and hydrogen bonds with conserved residues in the α2C′, α2D′, and α2E′ helices (we designate the helices of each HR as X and X′, as in Slep and Vale, 2007, with the numeral 2 indicating TOG2). An invariant asparagine (N491) in α2D′ forms a hydrogen bond with a conserved tyrosine residue (Y588) on the β2′ strand of the β-hairpin. The β-hairpin is further locked alongside the TOG domain by a network of hydrogen bonds formed between the A589 backbone amide and a conserved glutamate side chain (E531) in α2E′, as well as the T576 backbone carbonyl group and a conserved arginine side chain (R571) in helix α2F′ (Figure 4D). Commensurate with a stable, bona fide interaction
Acetylated tubulin

Pericentrin

Merge

A

Crescerin1

IMCD3

B

Crescerin1-GFP

HEK-293

C

Crescerin1-GFP

HEK-293

**FIGURE 3:** Crescerin1 localizes to the primary cilium in mammalian cells. (A) Antibody staining against endogenous Crescerin1 in IMCD3 cells shows localization to the primary cilium and basal body. (B) GFP-tagged mouse Crescerin1 localizes along the primary cilium and to the basal body in HEK-293 cells. (C) At high expression levels, GFP-tagged Crescerin1 accumulates at the basal body (yellow arrowhead) and distal tip (yellow arrow) of the primary cilium, as well as along the lattice of cytoplasmic microtubules. Primary cilia are labeled using antibody against acetylated tubulin (red), and basal bodies are labeled using antibody against pericentrin (blue). Scale bars, 5 μm.

between the β-hairpin and the central Hs, the β-hairpin has comparatively low B-factors that parallel those of the Hs with which it interacts. To assess further the contribution of the C-terminal β-hairpin to domain stability, we created a shorter TOG2 construct lacking the β-hairpin (TOG2-ΔβHP, residues 332–576) and performed secondary structure analysis by circular dichroism (CD). Although the overall α-helical signature was similar for both constructs (Figure 3E), the TOG2-ΔβHP construct had a lower melting temperature (T_m = 45.5°C) than the TOG2 construct containing the β-hairpin (residues 332–620; T_m = 57.5°C), indicating that the β-hairpin promotes TOG2 domain stability (Figure 3F). Secondary structure elements that flank TOG domains have been shown to stabilize other TOG domains (Leano et al., 2013). Crescerin family sequence alignments and secondary structure prediction analyses on the regions flanking TOG domains 1, 3, and 4 did not show conservation or similar predicted secondary structure, suggesting that the TOG2 C-terminal β-hairpin is a unique stabilizing feature for Crescerin1 TOG2. Whether this β-hairpin plays additional functional roles remains to be determined.

The Crescerin1 TOG2 tubulin-binding surface is similar to Stu2 TOG1

TOG domain structures from the ch-TOG and CLASP families reveal distinct structural variations that predict diversified interactions with αβ-tubulin. To determine the TOG structures to which Crescerin1 TOG2 is most similar, we performed pairwise structural alignments using the DALI server (Holm and Rosenström, 2010), aligning the first HR triads as a point of reference, since this region displays the highest conservation across TOG domains (Figure 5A). The Crescerin1 TOG2 intra-HEAT loop surface assumes a straight conformation that closely resembles Stu2 TOG1 and contrasts with the bent architectures observed in ch-TOG TOG4, MAST TOG1, and hCLASP1 TOG2 (Figure 5A, red arrows; Slep and Vale, 2007; Ayaz et al., 2012; De La Mora-Rey et al., 2013; Leano et al., 2013; Fox et al., 2014).

To gain insight into whether Crescerin1 TOG2 could form contacts with tubulin as observed in the Stu2 TOG1–αβ-tubulin structure, we docked Crescerin1 TOG2 on the tubulin heterodimer by structurally aligning Crescerin1 TOG2 HRs A–C (first HR triad) and Hs D–F (second HR triad) to the coordinates of Stu2 TOG1 in the Stu2 TOG1–αβ-tubulin complex (Ayaz et al., 2012; Figure 5, B–D and E–G, respectively). Many of the Stu2 TOG1 tubulin-binding residues are conserved and similarly positioned in Crescerin1 TOG2 (Figure 5, B–G). Prime β-tubulin binding determinants reside in Stu2 TOG1 HR A, including W23 and K24. Although electron density for the Crescerin1 TOG2 HR-A loop was poor and precluded modeling of the homologous residues Y364 and K365, the HR A helices are similarly positioned, and we anticipate that Y364 and K365 form homologous contacts with β-tubulin T107 and S400, respectively (T109 and G410 in mammalian β-tubulin; Figure 5C). In the Crescerin1 TOG2 HR-B and HR-C intra-HEAT loops, N406 and K447 occupy positions analogous to Stu2 N68 and R116 respectively, both of which interact with yeast β-tubulin Y106 (Y108 in mammalian β-tubulin; Figure 5D). In Crescerin1 TOG2 HR-D, K481 and R484 are positioned similarly to Stu2 K151 and K154, which interact with β-tubulin E158 and E157, respectively (E160 and E159 in mammalian β-tubulin; Figure 5F). Similarly, the highly conserved R526 in Crescerin1 TOG2 HR-E is positioned analogously to Stu2 R200, which interacts with α-tubulin V410, and E415 (V409 and E414 in mammalian α-tubulin; Figure 5G). The composition and conformational similarity between Stu2 TOG1 and Crescerin1 TOG2 intra-HEAT loops suggests that Crescerin1 TOG2 might be involved in binding free αβ-tubulin and/or MTs to regulate MT dynamics.

**Crescerin1 TOG domains promote MT polymerization in vitro**

To determine whether Crescerin might regulate the MT cytoskeleton, we first tested whether Crescerin1 TOG domains could influence MT polymerization in vitro. We mixed different purified TOG domains (Figure 6A) with tubulin and monitored tubulin polymerization at 37°C using a light scattering assay. The intrinsic polymerization activity of tubulin in the absence of other proteins is reflected as a characteristic sigmoidal increase in light scattering signal with an initial lag phase of ~500 s (Figure 6B, black curve). Addition of TOG1 or TOG3 to the reaction mixture did not affect the MT polymerization rate. In contrast, addition of either TOG2 or
FIGURE 4: The Crescerin1 second conserved domain is a bona fide TOG domain with unique features. (A) Crystal structure of Crescerin1 TOG2. Each HR (A–F) and the C-terminal β-hairpin are shown in a different color. See Supplemental Table S3 for crystallographic data, phasing, and refinement statistics. (B) Crescerin1 TOG2 shown in surface representation, oriented as in A, with cross-species conservation (left) or electrostatic charge distribution (right) mapped on the surface. See Supplemental Figure S2 for multiple sequence alignment of Crescerin TOG2 showing conservation across species. (C) Crescerin1 TOG2 conserved residues (colored as in B) map to the tubulin-binding intra-HEAT loops and to the C-terminal β-hairpin–HR interface. (D) Expanded view of the Crescerin1 TOG2 C-terminal β-hairpin–HR interface. (E, F) Circular dichroism spectra (E) of Crescerin1 TOG2 (blue) and TOG2-ΔβHP (red) at 20°C (solid lines) and 94°C (dotted lines) and thermal melt curves (F) monitored using 208-nm (solid lines) and 222-nm (dotted lines) light.
TOG4 dramatically increased the rate of MT polymerization. To determine whether TOG2 and TOG4 potentiated tubulin polymerization using canonical tubulin-binding determinants, we mutated a conserved aromatic residue in the HR A intra-HEAT loop in each TOG domain to glutamate (Y364E in TOG2 and F1559E in TOG4). Homologous mutations in other TOG domains have been shown to abrogate tubulin binding without affecting the TOG domain fold (Slep and Vale, 2007; Leano et al., 2013). We performed CD spectrum analysis on these constructs and confirmed that Y364E and F1559E mutations in TOG2 and TOG4, respectively, did not affect the α-helical fold and thermal stability of these domains (Supplemental Figure S4). The Y364E and F1559E mutations abrogated the ability of Crescerin1 TOG2 and TOG4, respectively, to accelerate tubulin polymerization, suggesting that both TOG2 and TOG4 use...
of Crescerin1 TOG domains to MT lattice-association activity. We compared cells with similar Crescerin1-GFP expression levels and classified MT lattice binding as strong, weak, or no lattice binding when Crescerin1-GFP colocalized with MTs completely, partially, or was diffuse in the cytoplasm, respectively (Figure 7, A–C). Full-length Crescerin1 was found to strongly colocalize with MTs in 72.3 ± 2.4% of transfected cells (Figure 7D). Mutating the intra-HEAT HR A loop tubulin-binding region in TOG1 and TOG2 (P109E and Y364E, respectively) did not appreciably alter MT-association activity. In contrast, mutating the intra-HEAT HR A–loop tubulin-binding region in TOG3 and TOG4 to glutamate (W1274E and F1559E, respectively) completely abrogated MT association. An N-terminal Crescerin1 truncation construct containing the central linker and TOG domains 3 and 4 (L-TOG-34, residues 577–1776) also had robust MT-association activity (Figure 7D). Crescerin1 L-TOG-34 MT association was fully ablated when the TOG4 HR A loop was mutated. A smaller Crescerin1 N-terminal truncation construct containing only a small portion of the central linker and TOG domains 3 and 4 (L–TOG-34, residues 1119–1776) showed minimal MT association, which was dependent on TOG3 and TOG4 HR A–loop tubulin-binding determinants. These data suggest that Crescerin1 TOG3 and TOG4 use canonical TOG domain intra-HEAT loop determinants to engage the straight tubulin conformation found in MTs and function synergistically with determinants in the central linker region to bind MTs. This also suggests that whereas the Crescerin1 N-terminal region encompassing TOG domains 1 and 2 has MT polymerization activity, it does not contribute dramatically to MT lattice association in our cellular overexpression assay, a characteristic that is shared with the polarized pentameric TOG domain array of the ch-TOG protein family (Currie et al., 2011, Fox et al., 2014).

The MT-regulatory function of CHE-12 is necessary for proper cilia development in C. elegans

We next sought to investigate how Crescerin affects cilia structure and function. Because it is difficult to exercise precise temporal control on Crescerin expression in mammalian cells to match endogenous levels at different cell cycle stages, we turned to C. elegans as a model system for these experiments, where we have the tools to perform structure/function experiments by modifying the endogenous che-12 locus. As discussed earlier, the C. elegans Crescerin family member CHE-12 localizes to amphid and phasmid sensory cilia (Bacaj et al., 2008; our results). Two che-12 mutant alleles, mn389 and mn399, were previously shown to cause shortened amphid cilia (Bacaj et al., 2008). However, these two mutations truncate the CHE-12 protein after the second TOG domain and may not be molecular null mutations. To determine the effect of a complete loss of CHE-12 function, we used Cas9-triggered homologous recombination to delete the entire che-12 coding sequence and replace it with gfp (Supplemental Figure S3). As expected, the resulting Δche-12 animals expressed GFP primarily in the amphid and phasmid sensory neurons, with GFP signal observed in the cell body and dendrite, overlapping with Pgcy-5::mCherry expression in the ASER neuron (Supplemental Figure S5C). To probe the role of CHE-12 in cilia structure, we first scored cilium length in ASER neurons using the Pgcy-5::mCherry reporter to visualize the ASER dendrite and cilium (Figure 8, A and B). ASER cilia in the wild-type (WT) N2 strain were 7.0 ± 0.6 μm. The phenotype of the che-12 alleles mn389 and mn399 that truncate CHE-12 after TOG2 are milder, with a ∼30% decrease in cilia length for the che-12(mn399) mutant compared with WT (Bacaj et al., 2008).
The more severe 54% decrease in cilia length that we observed in the Δche-12 animals suggests that the mn399 strain may retain partial CHE-12 function and highlights the efficacy of Cas9-triggered homologous recombination to generate complete gene deletions with full loss-of-function phenotypes.

We also measured cilia length in the GFP-tagged CHE-12 strains. Both CHE-12::GFP and GFP::CHE-12 knock-in animals had cilia that were slightly shorter than WT (6.0 ± 1.5 and 5.9 ± 1.5 μm, respectively), suggesting that the terminal tag might have interfered slightly with CHE-12 function, although not as dramatically as the che-12 deletion. A control strain, expressing a labeled nonneuronal protein (tagRFP::PH) in the same genetic background and containing the same unc-119 rescue marker used in all Cas9-mediated recombination experiments in this study had amphid cilia lengths that did not deviate statistically from WT (Figure 8, A and B).

We next asked whether CHE-12 uses canonical tubulin-binding determinants across its tetra-TOG domain array to promote proper cilia length. To answer this, we mutated a conserved hydrophobic residue on the predicted tubulin-binding surface of each TOG domain HR A intra-HEAT loop (F34, A281, W815, W1073) to glutamate (denoted TOG-1′2′3′4′) as we did for Crescerin1 in vitro and cellular studies. We used Cas9-triggered homologous recombination to generate complete gene deletions with full loss-of-function phenotypes.

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Previous electron microscopy work examining the mutant strain che-12(mn389) found ultrastructural defects in amphid cilia structure and the organization of MTs in these cilia (Bacaj et al., 2008). To determine what effect complete loss of CHE-12 function has on cilia ultrastructure, we examined cilia in WT, Δche-12, and GFP::CHE-12 TOG-1′2′3′4′ mutant worms using electron microscopy. In WT worms, the proximal sections of each of the 10 amphid channel cilia run parallel to each other, forming a well-organized bundle, with most cilia exhibiting the canonical ring of nine doublet MTs surrounding a variable number of central singlet MTs (Figure 8C). These bundled cilia become smaller and more tightly packed toward their distal ends.

![Figure 8](image_url)

**Figure 8:** Deleting che-12 or mutating tubulin-binding residues in CHE-12 TOG domains results in shorter and disorganized cilia in *C. elegans* amphid neurons. (A) Images of the Pgcy-5::mCherry–labeled ASER dendrite with sensory cilia (bracket) from WT and mutant strains. Scale bars, 5 μm. (B) ASER cilia length for strains shown in A. Values in parentheses indicate number of cilia measured/strain; error bars indicate SD. (C) Cross-sectional transmission electron micrographs of amphid sensilla in WT, Δche-12, and GFP::CHE-12 TOG-1′2′3′4′ strains. Left, expanded views of smaller, red-boxed regions on the right. Scale bars, 200 nm.
both the ch-TOG and CLASP families regulate spindle dynamics, facilitating proper segregation of sister chromatids between daughter cells. Both ch-TOG and CLASP require tubulin-binding activity across their TOG domain arrays in order to regulate MT dynamics. Although the activities of ch-TOG, CLASP, and a host of other cytoplasmic regulators of MT dynamics have been extensively studied, little is known about factors that regulate ciliary MTs. We identified Crescerin as a novel TOG domain–containing protein family through structural characterization of its second TOG domain. We find that Crescerin1 localizes to primary cilia in mammalian cells and uses its TOG domain array to promote MT polymerization in vitro. In C. elegans, we showed that the Crescerin family member CHE-12 requires canonical TOG tubulin-binding determinants for proper sensory cilia development. This highlights the emerging paradigm of MT regulators that mechanistically employ arrayed TOG domains with distinct structures and tubulin-binding activities to differentially regulate MT dynamics, whether during interphase or mitosis or in specialized organelles like the primary cilium.

Concordant with the ability of Crescerin1 to promote MT polymerization in vitro, we find that Crescerin1 TOG2 is most similar to TOG1 from the ch-TOG family of MT polymerases. However, Crescerin1 TOG2 does have unique features, including the presence of a β-hairpin at its C-terminal region that makes extensive contacts with HRs C–E and contributes to domain stability. The primary sequences of Crescerin’s four TOG domains bear low sequence identity to one another and to TOG domains from ch-TOG and CLASP families. Comparing TOG domain structures, we find that the tubulin-binding surface of many of these domains adopts different conformations, which imply different tubulin-binding modes, as well as potential effects on the curvature of the tubulin heterodimer. Our data demonstrate that Crescerin1 TOG domain pairs promote tubulin polymerization in vitro. Studies analyzing the ability of ch-TOG family TOG domains to promote tubulin polymerization in vitro found that a pair of TOG domains (TOG1 and TOG2 from yeast Ste2 or Drosophila Msps) are required for this activity, whereas the individual TOG domains are not sufficient (Slep and Vale, 2007; Ayaz et al., 2012; Fox et al., 2014). In contrast, a single TOG domain (TOG2) from human CLASP1 was sufficient to promote tubulin polymerization in vitro, suggesting that it either stabilized a conformation of tubulin more amenable for polymerization or aided in lattice contacts through a direct or allosteric mechanism (Leano et al., 2013). Of interest, Crescerin TOG2 and TOG4 are also able to individually increase MT polymerization rates in vitro, similar to human CLASP1 TOG2 (Leano et al., 2013). The structural mechanism underlying the ability of these individual TOG domains to promote tubulin polymerization remains to be determined. In cells, we find that the central linker with the C-terminal TOG(3+4) pair, when overexpressed, can bind the lattice of cytoplasmic MTs. This parallels similar effects observed when a linker-TOG3-4 and a linker-TOGS construct from the Drosophila ch-TOG family member Msps were overexpressed, yielding MT lattice association that was dependent on determinants in both the preceding linker and the TOG domain(s) (Currie et al., 2011). In contrast, Msps TOG1 and TOG2 do not appear to promote MT lattice association but have been shown to interact with free tubulin heterodimers over gel filtration, an activity that Msps TOG3 and TOG4 lack (Slep and Vale, 2007; Fox et al., 2014). This suggests that different TOG domains in the Msps array recognize different tubulin states: free tubulin versus polymerized tubulin. Collectively, this leads us to hypothesize that the Crescerin TOG array may also be polarized with differential tubulin-binding activity, in which the N-terminal TOG domains bind and promote tubulin incorporation into growing MTs, whereas the C-terminal TOG pair affords MT association.
Why is Crescerin specific to cilia, and how does this compare and contrast with the regulation of MT dynamics in the cytoplasm? Extensive work has characterized MAPs that regulate cytoplasmic MTs, including investigations that probe the roles and mechanisms of the TOG array proteins ch-TOG and CLASP. Although MAPs affect MT dynamics, a key underlying factor is the concentration of tubulin and its ability to diffuse throughout the cytoplasm. This contrasts with components at the cilium tip, where the singlet microtubule plus ends reside. In this diffusion-limited space, a small fraction of the tubulin that is present diffuses in from the cytoplasm while the major fraction is transported to the cilia distal tip via IFT to reach the concentration required for MT polymerization, particularly during the nonequilibrium phase of ciliogenesis, when the flux of IFT-coupled tubulin is greatest (Craft et al., 2015). Previous studies identified cilia-specific tubulin isoforms in C. elegans (Hao et al., 2011), as well as in mammals (Péchart et al., 1999; Jensen-Smith et al., 2003). These tubulin isoforms are coupled to IFT and are actively transported to the distal tip (Hao et al., 2011; Bhogaraju et al., 2013). Little is known about the critical concentration of these tubulin isoforms or their dynamics, but models that predict how cilia of fixed length are generated and maintained rely on specific parameters that include the flux of tubulin to and from the cilia tip, as well as the parameters of ciliary microtubule dynamics (Ludington et al., 2015). Recent work found that the tubulin diffusion rate is slower at the cilia tip than at other ciliary regions, which suggests that nonpolymerized tubulin engages in macromolecular interactions with ciliary tip components (Craft et al., 2015). The differential nature of tubulin isotype and availability, as well as the difference in MT dynamics in the cytoplasm versus the distal cilia tip, suggests that additional, cilia-specific MT regulators are required to regulate ciliary MTs. We suggest.

FIGURE 9: Deleting che-12 or mutating tubulin-binding residues in CHE-12 TOG domains affects amphid cilia function. (A) Schematic diagram of C. elegans chemotaxis assay on a 0–60 mM NaCl gradient (red–green) or 30 mM NaCl isocratic (yellow) agar plates. Worms are placed at the plate center and migration scored after 30 min. WT animals preferentially migrate toward 60 mM NaCl on the gradient plate but distribute uniformly on an isocratic plate. (B) Bar graph showing the normalized density (see Materials and Methods) of worms in different sections of the plate. Error bars indicate SDs over the course of four or five experiments. (C) Quantitated results of Dil uptake assay. Error bars indicate SDs from three independent experiments. (D, E) Images showing strong dye-filling in WT (D) C. elegans amphid neurons (yellow arrow) and no dye filling in GFP::CHE-12 TOG-1’2’3’4’ amphid neurons (E). Gut staining due to feeding: yellow arrowhead. Scale bars, 50 μm.
that these differential requirements underlie the need for Crescerin in the cilium. Just as ch-TOG and CLASP are part of larger regulatory complexes, we anticipate that Crescerin interacts with other regulators of ciliary MT dynamics, coordinating phases of polymerization and depolymerization, driving polymerization during cilio-genesis, or oscillating between short phases of polymerization and depolymerization during the maintenance phase to retain a cilium of proper length.

With these observations, we reclassify Crescerin as a third conserved TOG domain–containing protein family found in ciliated and flagellated eukaryotes, specialized for MT regulation in cilia. In C. elegans, CHE-12 expression is restricted to a small set of sensory neurons that have a simple rod-shaped cilium. Further work is necessary to fully understand Crescerin1 and Crescerin2 expression and function in mammalian systems, where different tissues have diversified cilia that perform highly specialized functions. Future experiments will examine whether Crescerin is associated with IFT particles, how it affects cilia MT dynamics, and whether it serves to deliver cilia-specific tubulin isoforms to the distal cilia tip.

**MATERIALS AND METHODS**

**Sequence analysis and phylogenetic tree**

Homologues of Crescerin from different organisms in the National Center for Biotechnology Information (NCBI) database were identified using the BLASTP algorithm (Altschul et al., 1990). The NCBI accession numbers of Crescerin homologues used for sequence alignments are given in Supplemental Table S1. Secondary structure predictions were performed using Jpred 3 (Cole et al., 2008) and PSIPRED servers (Jones, 1999; Buchan et al., 2013). Phylogenetic and molecular evolutionary analyses were conducted using integrated algorithms at the server phylogeny.imm.fr/phyllo.cgi/index.cgi (Dereeper et al., 2008) and MEGA, version 6 (Tamura et al., 2013). Sequence alignment was carried out using the MUSCLE algorithm with the neighbor-joining method of clustering in MEGA. We used structural data from previously solved TOG domain structures to further optimize the alignment in the intra-HEAT loops. We constructed the initial tree using the maximum likelihood method implemented in the PhyML program (version 3.0). The LG substitution model was selected assuming an estimated proportion of invariant sites and four gamma-distributed rate categories to account for rate heterogeneity across sites. The gamma shape parameter was estimated directly from the data. Reliability for internal branch was assessed using the bootstrapping method (100 bootstrap replicates). We used this initial tree for tree inference using the Subtree-Pruning-Regrafting (extensive) algorithm with a very weak branch swap filter in MEGA, version 6. Subsequent reliability for internal branch was assessed using the bootstrapping method (1000 bootstrap replicates). The final tree was prepared for publication using FigTree, version 1.4.0.

**Cloning and expression**

Mouse Crescerin1 TOG2 (residues 332–620) was cloned into pET28b (Novagen, EMD Millipore, Darmstadt, Germany), generating a thrombin-cleavable N-terminal hexahistidine (His6) tag. Crescerin1 TOG2 was transformed into BL21 DE3 Escherichia coli and grown in 6 l of Luria broth (50 μg/ml kanamycin) at 37°C to an optical density of 0.6 at 600 nm. The temperature was lowered to 15°C and protein expression induced with 100 μM isopropyl-β-D-thiogalactoside (final concentration) for 16 h. Cells were centrifuged at 23,000 × g for 45 min. Supernatant was loaded onto a 10-ml Ni2+-nitriloacetic acid column (Qiagen, Venlo, Netherlands) and washed with 300 ml of buffer A, and TOG2 was eluted over a 250-ml linear gradient between buffers A and B (buffer B = buffer A supplemented with 290 mM imidazole). TOG2-containing fractions were pooled, CaCl2 was added to 2.5 mM, and 0.1 mg of bovine α-thrombin (Haematologic Technologies, Essex Junction, VT) was added to remove the His6 tag, leaving an N-terminal Gly-Ser-His-Met cloning artifact. Thrombin digest proceeded for at least 16 h at 4°C. Digested protein was filtered over 0.5 ml of benzamidine-Sepharose (GE Healthcare, Buckinghamshire, United Kingdom) and exchanged into buffer C (25 mM 4-[(2-hydroxyethyl)-1-piperazineethanesulfonic acid [HEPES], pH 7.0, 0.1% β-ME) by dialysis. Protein was loaded onto a 10 ml of SP-Sepharose fast flow column (GE Healthcare), washed with 200 ml of buffer C, and eluted over a 250-ml linear gradient between buffers C and D (buffer D = buffer C supplemented with 1 M NaCl). Protein fractions were pooled, concentrated, and exchanged into 10 mM HEPES, pH 7.5, 100 mM NaCl, and 0.1% β-ME in a Millipore Ultrafree 10,000 MWCO concentrator to 15 mg/ml, frozen in liquid nitrogen, and stored at −80°C. Purification of selenomethionine-substituted protein proceeded according to the aforementioned protocol.

**Crystallization**

Crescerin1 TOG2 was crystallized using the hanging-drop vapor diffusion method. A 2-μl amount of protein (native and selenomethionine-substituted protein) at 10 mg/ml was added to an equal volume of a well solution containing 25% polyethylene glycol 3350, 0.25 M magnesium chloride, and 0.1 M HEPES, pH 7.5, 0.1% β-ME in a Millipore Ultrafree 10,000 MWCO concentrator to 15 mg/ml, frozen in liquid nitrogen, and stored at −80°C. Purification of selenomethionine-substituted protein proceeded according to the aforementioned protocol.

**Data collection, structure determination, and refinement**

Isomorphous Crescerin1 TOG2 native and selenium peak single-wavelength anomalous dispersion data sets were collected on single crystals to resolutions of 2.1 and 2.3 Å, respectively. Diffraction data were collected at the Advanced Photon Source 22-ID beamline, Argonne National Laboratory, at 100 K. Crescerin1 TOG2 crystals belong to the space group P212121, with one molecule in the asymmetric unit. Data were indexed, integrated, and scaled using HKL2000 (Otwinowski and Minor, 1997). Selenium sites were identified and used to generate initial experimental, density-modified electron density maps (PHENIX; Grosse-Kunstleve and Adams, 2003; Terwilliger, 2004, 2005; McCoy et al., 2007; Moriarty et al., 2009; Adams et al., 2010; Chen et al., 2010; Afonine et al., 2012). Initial models were built using AutoBuild (PHE-NIX), followed by iterative manual building in Coot and refinement runs using phenix.refine (PHENIX). The selenomethionine-substituted TOG2 structure was refined against a ML2 target.
function, and the model was converted to native methionine and refined against native data using a ML target function to 2.2-Å resolution. Refinement was monitored using a Free R, based on 10% of the data randomly excluded from refinement. No clear electron density was evident for N-terminal residues 332–344, HR A loops residues 361–365, HR F residues 558–562, and the C-terminal residues 595–620. Lyase K380 was modeled as alanine. Information regarding data statistics, model building, and refinement is given in Supplemental Table S3. Atomic coordinates have been deposited in the Protein Data Bank under accession code 5DN7.

Circular dichroism

The CD spectra of Crescerin1 TOG domains were collected at 20 and 94°C using a Chirascan-plus CD spectrometer (Applied Photophysics, Leatherhead, United Kingdom). Each protein sample was diluted to a final concentration of 0.1 mg/ml in 10 mM sodium phosphate buffer (pH 7.5) and 50 mM NaF. The spectra were recorded from 260 to 190 nm with a step size of 0.5 nm using a cuvette of 1 mm path length. The time per point was kept at 1.25 s. A baseline control CD spectrum was taken and subtracted from each spectrum. All spectra were smoothed in the Chirascan-plus software with a moving window of five points. In CD thermal melt experiments, data were collected at 1°C intervals from 20 to 94°C at 208- and 222-nm wavelengths.

In vitro MT polymerization assay

Crescerin1 constructs for MT polymerization assays (TOG1, 67–317; TOG2, 332–620; TOG2-Y364E; TOG3, 1243–1491; TOG4, 1533–1776; TOG4-F1559E; TOG1(+2), 67–620; and TOG3(+4), 1243–1776) were cloned into E. coli expression vectors, expressed, and purified in a manner similar to that for TOG2 as detailed earlier. Tubulin was purified from bovine brain following protocols established by the Mitchison lab (http://mitchison.med.harvard.edu/protocols/microtubules/Large%20Scale%20Tubulin%20Preparation.pdf). Tubulin polymerization was monitored using a SPEX Fluorolog-3 spectrophotometer (Horiba Jobin Yvon, Edison, NJ) in T-format, high-voltage mode. The excitation and detection wavelengths were set at 350 nm and the excitation and emission slits at 0.5 and 0.75 nm, respectively. The cuvette holder was maintained at 37°C. Clarified tubulin samples (18 μM final concentration) were prepared in the presence or absence of Crescerin1 TOG domain constructs each at 3 μM final concentration in polymerization buffer (50 mM 2-(N-morpholino)ethanesulfonic acid [MES], pH 6.6, 3.4 M glycerol, 5 mM dithiothreitol, 1 mM ethylene glycol tetraacetic acid [EGTA], 5 mM MgSO4, 1 mM GTP) and incubated on ice for 8 min. The 300-μl reactions were loaded onto a 10% SDS-PAGE gel and stained with Coomassie blue. Tubulin polymerization was monitored using a SPEX Fluorolog-3 spectrophotometer (Horiba Jobin Yvon, Edison, NJ) in T-format, high-voltage mode. The excitation and detection wavelengths were set at 350 nm and the excitation and emission slits at 0.5 and 0.75 nm, respectively. The cuvette holder was maintained at 37°C. Clarified tubulin samples (18 μM final concentration) were prepared in the presence or absence of Crescerin1 TOG domain constructs each at 3 μM final concentration in polymerization buffer (50 mM 2-(N-morpholino)ethanesulfonic acid [MES], pH 6.6, 3.4 M glycerol, 5 mM dithiothreitol, 1 mM ethylene glycol tetraacetic acid [EGTA], 5 mM MgSO4, 1 mM GTP) and incubated on ice for 8 min. The 300-μl samples were injected into a quartz cuvette of 4 mm pathlength at room temperature and then immediately placed into the 37°C cuvette holder. Scattering was recorded at 350 nm in 1-s intervals over 1500 s. Spikes in scattering within the first 100 s were the result of samples equilibrating to 37°C.

Mammalian expression plasmids

For constitutive expression in mammalian cells driven by the cytomegalovirus (CMV) immediate early promoter, full-length and truncated mouse Crescerin1 constructs (Figures 3, B and C, and 7) were cloned into pcDNA-DEST47 and pcDNA-DEST53 plasmids using the Gateway cloning system (Invitrogen, Thermo Fisher Scientific, Waltham, MA), generating C-terminal and N-terminal GFP fusion proteins, respectively.

Cell culture and transfection

IMCD3 cells were cultured in DMEM:F-12 (1:1) (Gibco, Thermo Fisher Scientific, Waltham, MA) supplemented with 10% fetal bovine serum (FBS; Gibco) and 1x Anti-Anti (Gibco) at 37°C and 5% CO2. HEK-293 cells were cultured in DMEM (Gibco) supplemented with 10% FBS and 1x Anti-Anti at 37°C and 5% CO2. For transfection, IMCD3 and HEK-293 cells were seeded in six-well plates and transfected when they reached 70–80% confluence. At 1 h before transfection, the media in the cell culture plates were replaced with either Opti-MEM (IMCD3) or nonsupplemented DMEM (HEK-293). Cells were transfected with 2.5–3 μg of the plasmid DNA per well of a six-well plate, using either polyethylenimine or Lipofectamine 3000 (Invitrogen) transfection reagents for HEK-293 and IMCD3 cells, respectively. After 6–8 h of transfection, the cells were seeded on glass coverslips in low-serum–supplemented (0.25% FBS) DMEM. HEK-293 cells were serum starved for 24–48 h and IMCD3 cells were serum starved for 4 d to promote growth arrest and induce ciliogenesis. For MT lattice–binding assays, HEK-293 cells (not serum starved) were fixed 24 h after transfection.

Immunofluorescence microscopy of mammalian cells

For visualization of primary cilia in mammalian cells, the cells were first treated with PHEM buffer (60 mM 1,4-piperazinediethanesulfonic acid [PIPES], 21 mM HEPES, 10 mM EGTA, 2 mM MgCl2, pH 7.5) containing 4% formaldehyde for 20 s, followed by permeabilization in PHEM buffer supplemented with 0.2% Triton X-100 for 2 min. The cells were then fixed in fresh PHEM buffer supplemented with 4% formaldehyde for 20 min and then blocked with 10 mg/ml BSA solution dissolved in PHEM buffer supplemented with 0.2% Triton X-100. Polyclonal antibodies against mouse Crescerin1 were raised in guinea pig (Pocono Rabbit Farm and Laboratory, Canadensis, PA) using purified fragments of TOG1 and TOG2 as antigen. Mouse monoclonal anti-acetylated tubulin antibody (clone 6-11B-1, T7451; Sigma-Aldrich) was used for labeling the primary cilium, and anti-pencillin-tin antibody raised in rabbit (ab4448; Abcam, Cambridge, United Kingdom) was used for labeling centrioles and basal bodies. Mouse monoclonal anti-α-tubulin antibody (clone D1MA1, T6199; Sigma-Aldrich) was used for labeling microtubules. Nuclei were stained with 4’,6-diamidino-2-phenylindole (Molecular Probes, Invitrogen). All samples were mounted with mounting medium (20 mM Tris, pH 8.5, 90% glycerol, 5 mg/ml n-propyl gallate). Images were collected on an inverted epifluorescence microscope (Eclipse Ti; Nikon, Tokyo, Japan) using a 100× objective. Images were analyzed using NIS-Elements imaging software and ImageJ (National Institutes of Health, Bethesda, MD).

C. elegans culture and strain construction

C. elegans strains were kept at 20°C and fed E. coli OP50 bacteria except where noted. Strain N2 was used as WT. To label the ASER neuron for imaging, we generated extrachromosomal arrays by injecting a plasmid encoding Pgcy-5:mCherry (a gift from Shai Shaham, Rockefeller University) together with the plasmid pRF4, which carries the dominant rol-6(su1006) marker (Mello et al., 1991). Supplemental Table S2 shows a list of all strains generated and used in this study. All new che-12 alleles reported in this study were generated using Cas9-triggered homologous recombination (Dickinson et al., 2013). For each modification, we built a homologous repair template comprising the desired edited nucleotide sequence flanked on each side by 1.5 kb of unmodified genomic sequence. Each repair template also contained an uncl-119(+)-selectable marker flanked by LuxP sites. In addition, we generated a Cas9–sgRNA construct by inserting the desired target sequence into the vector pDD162 (Dickinson et al., 2013) using site-directed mutagenesis. Cas9 target sites were selected using the CRISPR design tool available at crispr.mit.edu/ and are listed in Supplemental Table S2.
Young adults carrying an unc-119(ed3) mutant allele were injected with a plasmid mixture containing 50 ng/µl Cas9–sgRNA plasmid, 10 ng/µl homologous repair template, 10 ng/µl pMA122 (Phsp16-41::PEEL-1), 10 ng/µl pGH8 (Prab-3::mCherry), 5 ng/µl pCFJ104 (Pmyo-3::mCherry), and 2.5 ng/µl pCFJ90 (Pmyo-2::mCherry). Injected animals were placed at 25°C for 8–12 d and then heat shocked for 4 h at 34°C to induce expression of the PEEL-1 toxin, which kills animals carrying extrachromosomal arrays. Plates carrying non-uncoordinated animals that survived heat shock and lacked the mCherry coinjection were considered candidate knockout/knock-in founders, and single worms from these plates were picked to establish lines. Genomic DNA was isolated from each line, and the presence of the desired genome modification was confirmed by PCR and sequencing. Only a single line from each independent plate was kept.

Supplemental Figure S3 shows the strategy for modifying the che-12 locus using homologous recombination. To generate che-12-knockout alleles, we designed a homologous repair template to delete the entire che-12 coding sequence and replace it with gfp. This construct was injected into young adults of strain HT1593 for Cas9-triggered homologous recombination as described. To generate GFP-tagged WT and mutant che-12 alleles at the endogenous locus, we first removed the unc-119(+)/ selectable marker from the che-12-knockout strain LP177 by injecting Cre recombinase (Dickinson et al., 2013). The resulting uncoordinated animals were used for a second round of Cas9-triggered homologous recombination to reintroduce the WT or mutant che-12 gene.

Fluorescence microscopy of C. elegans

For fluorescence microscopy of C. elegans strains, adult worms were mounted on thin agar pads made of 2.5% agar dissolved in M9 medium. We used polystyrene nanoparticle beads (2.5% by volume, 0.1-µm diameter; Polysciences, Warrington, PA) to immobilize the worms on the agar pad as described by Kim et al. (2013). The worms were imaged using a Nikon Eclipse Ti microscope equipped with 100X, 1.4 numerical aperture (NA) objective and Yokogawa (Tokyo, Japan) CSU-10 spinning disk head.

Transmission electron microscopy of C. elegans

Adult C. elegans hermaphrodites were washed off agar plates with M9 buffer and collected into 1.5-ml microcentrifuge tubes. The worms were allowed to settle to the bottom of the tube and washed in M9 buffer to wash off the bacteria and then were fixed in 2% paraformaldehyde/2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer, pH 7.4, with 0.05% CaCl₂. Samples were microwave irradiated using a PELCO BioWave Pro microwave system (Ted Pella, Redding, CA) at 450 W to a temperature of 40°C, allowed to cool to room temperature, and stored at 4°C for several days before processing. After several washes in 0.1 M sodium cacodylate buffer, the samples were postfixed in 1% osmium tetroxide/1.25% potassium ferrocyanide in 0.1 M sodium cacodylate buffer for 1 h, followed by rinses in deionized water. (LP177 worms were encapsulated in agarose before the next step.) Samples were dehydrated in a graded series of ethanol solutions (30, 50, 75% 90, 100, 100) followed by two changes in propylene oxide, and infiltration and embedding in Spurr’s low-viscosity epoxy resin (Polysciences). Ultrathin sections (70–80 nm) were cut using a diamond knife, mounted onto 200 mesh copper grids or Formvar carbon–coated copper slot grids, and stained with 4% aqueous uranyl acetate and Reynolds’s lead citrate. Samples were observed using a LEO EM910 transmission electron microscope operating at 80 kV (Carl Zeiss Microscopy, Peabody, MA), and digital images were acquired using a Gatan Orius SC1000 CCD Digital Camera with Digital Micrograph 3.11.0 (Gatan, Pleasanton, CA).

Chemotaxis assays

Chemotaxis plates with a linear NaCl gradient were prepared according to an established protocol (Luo et al., 2014). Briefly, 2% agar solutions in chemotaxis buffer (5 mM phosphate, 1 mM CaCl₂, 1 mM MgSO₄), with or without 60 mM NaCl, were prepared and sterilized by autoclaving. Before pouring, each Petri dish (circumference, 8.5-cm diameter) was labeled to mark the high-salt and low-salt ends and six parallel sections of equal width perpendicular to the gradient axis. The plates were uniformly elevated at the low-salt end. Then 20 ml of high-salt agar (containing 60 mM NaCl) was poured into the plate and allowed to harden. This created a sloped agar block with maximum height at the high-salt end and barely covering the surface at the low-salt end. The plates were then laid flat, and 20 ml of the low-salt agar (0 mM NaCl) was poured over the high-salt agar block to create a horizontal surface. Plates were incubated for 16 h to allow NaCl to diffuse, creating a 0–60 mM NaCl gradient across the agar surface. Control isocratic chemotaxis plates were prepared using sterile chemotaxis agar supplemented with 30 mM NaCl.

The C. elegans strains were grown for 3 d on nematode growth media plates seeded with bacteria to obtain a synchronous adult population. The worms were washed off the plates in chemotaxis buffer and collected in microcentrifuge tubes. The worms were allowed to settle to the bottom of the tube and washed three times with chemotaxis buffer to wash off the bacteria, embryos and juveniles. From 150 to 300 adult worms, resuspended in 20 µl of chemotaxis buffer, were then pipetted into the plate center. Timing for the chemotaxis assay was initiated when the buffer evaporated completely and the worms started crawling on the agar surface. After 30 min, the plates were cooled to 4°C to paralyze the worms, and the number of worms in each of six sectors along the plate’s linear NaCl gradient were counted. We determined the normalized worm density for each sector, which we define as follows:

\[
\text{Normalized density} = \frac{\text{Area of sector } X}{\text{Total area of the plate}} \times \frac{\text{Number of worms on sector } X}{\text{Total number of worms on the plate}}
\]

By this measure, when the animals are uniformly distributed throughout the plate, the normalized density for each sector will be equal to 1.

Dye-uptake assay

Adult C. elegans hermaphrodites were washed off agar plates with M9 buffer and collected into 1.5-ml microcentrifuge tubes. The worms were allowed to settle to the bottom of the tube and washed three times in M9 buffer to wash off the bacteria. The worms were then resuspended in M9 buffer containing 1,1′-dioctadecyl-3,3,3′,3′-tetramethylindocarbocyanine perchlorate (DiI; Anaspec, Fremont, CA) at a final concentration of 10 µg/ml and incubated at 20°C for 3 h. Postincubation, the worms were washed again in M9 buffer and transferred on a new agar plate to remove excess dye. After 5 min, the worms were mounted on thin agar pads made of 2.5% agar dissolved in M9 medium supplemented with 10 mM NaNO₃ to immobilize the animals. The worms were imaged using a Nikon Eclipse Ti microscope equipped with a 100×/1.4 NA objective and a Yokogawa CSU-10 spinning disk head. A total of ~100 worms were imaged for each strain over the course of three experiments.
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