Asymmetrically localized proteins stabilize basal bodies against ciliary beating forces

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Asymmetrically localized proteins stabilize basal bodies against ciliary beating forces

Brian A. Bayless, Domenico F. Galati, Anthony D. Junker, Chelsea B. Backer, Jacek Gaertig, and Chad G. Pearson

Basal bodies are radially symmetric, microtubule-rich structures that nucleate and anchor motile cilia. Ciliary beating produces asymmetric mechanical forces that are resisted by basal bodies. To resist these forces, distinct regions within the basal body ultrastructure and the microtubules themselves must be stable. However, the molecular components that stabilize basal bodies remain poorly defined. Here, we determine that Fop1 functionally interacts with the established basal body stability components Bld10 and Poc1. We find that Fop1 and microtubule glutamylation incorporate into basal bodies at distinct stages of assembly, culminating in their asymmetric enrichment at specific triplet microtubule regions that are predicted to experience the greatest mechanical force from ciliary beating. Both Fop1 and microtubule glutamylation are required to stabilize basal bodies against ciliary beating forces. Our studies reveal that microtubule glutamylation and Bld10, Poc1, and Fop1 stabilize basal bodies against the forces produced by ciliary beating via distinct yet interdependent mechanisms.

Introduction

Motile cilia produce hydrodynamic forces for diverse functions including cellular motility and fluid flow (Marshall and Kintner, 2008). The ciliary axoneme generates hydrodynamic force by using axonemal dynein to slide nine doublet microtubules arranged around two single microtubules (Satir et al., 2014). This activity leads to a two-phase, asymmetric beat stroke. During the power stroke, the extended cilium moves perpendicular to the cell surface, whereas during the recovery stroke, the bent cilium moves parallel to the cell surface, thereby returning the cilium for another cycle. Although axonemal dynein is sufficient to undulate cilia, to generate effective fluid flow, ciliary forces must be coupled to the cell cortex.

Basal bodies (BBs) nucleate motile cilia and anchor them to the cell cortex. They comprise nine radially symmetric triplet microtubules organized around a cartwheel structure. Because the A and B tubules of BBs are continuous with the axoneme tubules, BBs directly experience cilia-generated forces (Dippell, 1968; Allen, 1969; Dirksen, 1971). In sperm flagella, where axonemal mechanics are best understood, the nine microtubule doublets effectively form two segments. As flagella beat, these segments slide ∼110 nm relative to each other in a piston-like fashion, which leads to symmetric compression and tension at the cilary base (Lindemann et al., 1992; Vernon and Woolley, 2004; Riedel-Kruse et al., 2007). However, during ciliary beating, which is asymmetric, the greatest compressive and tensile forces are predicted to occur at the sides of the BB where the power stroke terminates and initiates, respectively. The BB structures and molecules that compensate for the forces produced by motile cilia are poorly defined.

Despite the persistent exposure to ciliary forces, BBs are stable (Brinkley and Cartwright, 1975; Bobinnec et al., 1998a; Abal et al., 2005; Kunimoto et al., 2012). This stability stems from two conserved aspects of BB organization. First, symmetrically localized proteins, such as Bld10 and Poc1, link the radially symmetric BB structural domains. Bld10 connects the spokes of the BB cartwheel to each of the nine triplet microtubules, whereas Poc1 is crucial for linking triplet microtubules to one another (Bayless et al., 2012; Meehl et al., 2016). In bld10Δ and poc1Δ Tetrahymena thermophila cells, ciliary beating causes BB disassembly and triplet microtubule loss from sites that experience the greatest compressive and tensile forces. The second aspect of BB stability stems from microtubule-associated proteins and tubulin posttranslational modifications (PTMs), such as glutamylation, that directly stabilize triplet microtubules (Magiera and Janke, 2014). Depletion of the microtubule-associated protein CAP350 renders centriole triplet microtubules sensitive to nocodazole (Le Clech,
2008). Antibodies targeting microtubule glutamylation, which is prominent at axonemes and BBs, cause centriole fragmentation (Bobinnec et al., 1998a), whereas loss of the Tetrahymena tubulin glutamylases TTLL1 and TTLL9 leads to fewer ciliary rows and defects in BB maturation (Wloga et al., 2008). However, the interdependence of the various proteins and tubulin PTMs that stabilize BBs remains poorly defined. Moreover, it is unclear how these stability factors compensate for the asymmetric distribution of ciliary forces.

We identify the Tetrahymena FGFR1 oncogenic partner (Fop1) as a Bld10- and Poc1-associated protein required for BB stability. Both Fop1 and microtubule glutamylation are asymmetrically enriched at triplet microtubules predicted to experience the greatest compression forces from ciliary beating. Poc1 is necessary for normal Fop1 localization to BBs, whereas Poc1 and Fop1 levels affect BB microtubule glutamylation. These results suggest that cooperation between distinct pathways stabilizes BBs against ciliary forces.

Results and discussion

Fop1 is a BB stability protein
Candidate Tetrahymena BB stability proteins were identified by isolating Bld10- and Poc1-associated proteins (Tables S1, S2, and S3). 26 potential stability factors, including Fop1 (TTH ERM_00537420), whose orthologues localize to centrosomes and function in microtubule anchoring, were identified (Mikolajka et al., 2006; Yan et al., 2006). To assess whether Fop1 stabilizes BBs, we depleted FOP1 from Tetrahymena cells and quantified the linear density of BBs within the medial half of ciliary rows (Fig. 1, A and B; and Fig. S1, A–C). FOP1 knockdown reduces BB frequency by 16%, and, as observed for other BB stability proteins, increasing temperature exacerbates this defect (Fig. S1, C–E; 37°C; Pearson et al., 2009; Bayless et al., 2012). Because the number of ciliary rows remains constant, the reduced BB frequency represents actual BB loss (Fig. S1 G; Nanney and Chow, 1974).

BB loss could arise from reduced BB assembly through the cell cycle (Bayless et al., 2012). To test this, FOP1 knockdown cells were starved, which arrests cells in G1 and halts new BB assembly (Figs. 1 C and S1 F). In selected G1 cells, there are 16% fewer BBs, which suggests that a fraction of BBs disassemble upon FOP1 knockdown. To determine whether the remaining BBs exhibit structural defects, we examined the BB ultrastructure using EM. 37% of FOP1 knockdown BBs have a reduced diameter or longitudinal tapering distal to the cartwheel (Figs. 1 D and S1 H). Moreover, 54% of BBs have C tubules that are missing or disconnected from their associated B tubules (Figs. 1 E and S1 I). Thus, Fop1 likely stabilizes BBs by promoting C-tubule formation and/or maintenance.

Figure 1. Fop1 is a BB stability protein. (A) Whole-cell BB immunofluorescence (α-centrin). FOP1 knockdown (KD) cells have fewer BBs compared with wild type (WT). The cell’s anterior is defined by the oral apparatus (arrow). Bar, 10 µm. (B) Schematic of a 10-µm segment of BBs along a ciliary row that represents the region used for BB frequency analysis. Bar, 1 µm. (C, top) FOP1 KD cells have fewer BBs than WT cells. BB loss is rescued by the reintroduction of FOP1. Representative fluorescence images of BB rows in WT, FOP1 KD, and FOP1 KD with FOP1 rescue at 30°C and 37°C in cell cycle arrested cells. (bottom) Quantification of BBs per 10 µm. Mean ± SEM; n = 300 rows; *, P < 0.01. (D and E) FOP1 KD causes constricted BBs with defective C tubules (arrows). Electron micrographs of longitudinal (D) and cross-sectional (E) views of FOP1 KD BBs at 37°C. n = 50 BBs for each orientation. Bars, 100 nm.
BBs (Goto et al., 1982; Pearson et al., 2009). Both high temperature and viscous, polyethylene oxide (PEO)–containing media exacerbate BB loss in FOP1 KD cells (Fig. 2, A and B). Conversely, NiCl2 treatment rescues BB loss caused by shifting FOP1 knockdown cells to 37°C (Fig. 2 C). These results suggest that Fop1 stabilizes BBs against the forces derived from ciliary beating.

### Fop1 localizes asymmetrically to BBs

The Tetrahymena ciliary power stroke moves from the cell’s anterior toward the posterior. Thus, the BB posterior face may experience asymmetric compressive force, whereas the anterior face experiences asymmetric tensile force. Because Fop1 stabilizes BBs, we asked whether Fop1 is asymmetrically positioned to reinforce the BB posterior and/or anterior faces. Endogenously tagged Fop1::mCherry localizes posteriorly relative to the centrally localized BB component Sas6a (Fig. 3 A). To quantify this localization, we determined the mean Fop1 signal relative to Sas6a, Cen1, and Poc1, which have known positions within the BB architecture (Fig. 3, B and C; and Fig. S2, A and B; Culver et al., 2009; Pearson et al., 2009). Indeed, fluorescence image averaging confirms that Fop1 is enriched at the posterior face of BBs. Furthermore, colocalization of N- and C-terminal–tagged Fop1 shows that the Fop1 protein is not extended or arranged in a polarized orientation (Fig. S2 C).

To examine the radial distribution of Fop1 at BBs, we used structured illumination microscopy (SIM; Fig. 3 D–F; and Fig. S2, D and F). Surprisingly, Fop1 variably localizes as horseshoe and ring shapes (Fig. 3 E). The angular displacement of the horseshoe varies and we predict that these localization patterns reflect dynamics of Fop1 protein within BBs. When the Fop1 SIM signal is averaged, the apex of the horseshoe preferentially localizes to the BB posterior face and, depending on the experiment, is slightly offset to triplet microtubules 1, 2, 8, and 9 (Fig. 3, B, D, and F; and Fig. S2, D and F). Linear and radial linescans reveal a second peak of lower intensity toward the anterior-facing triplet microtubules 3–5 that are often directly opposite to the posterior Fop1 signal. Subtle differences between imaging approaches may have resulted from minor deviations in sample alignment during averaging. Consistent with this result, immuno-EM shows a similar distribution (Figs. 3 G and S2 G). Collectively, Fop1 variably accumulates at BB regions predicted to experience the greatest compressive and tensile forces produced by ciliary beating.

A remaining question is how Fop1 is asymmetrically positioned to the posterior and anterior-facing triplet microtubules. One possibility is that the unequal forces generated by ciliary beating establish asymmetric cues for Fop1 localization. However, Fop1 is asymmetrically localized before ciliogenesis, although the level of Fop1 asymmetry increases with BB maturation and ciliary forces leading us to hypothesize that ciliary beating increases the asymmetric Fop1 localization (Fig. S2 E). Fop1 remains asymmetrically localized even when ciliary beating is inhibited (Fig. S2 F). This suggests that ciliary forces do not initiate or maintain asymmetric Fop1 localization. New BB assembly occurs from a predetermined triplet microtubule at the BB proximal end (O’Toole and Dutcher, 2014; Pearson, 2014), and BBs attach to the cellular cytoskeleton through structures that are asymmetric in their distribution (Bayless et al., 2016). Thus, the rotationally symmetric BB has inherent functional and structural asymmetries, and we predict that these preexisting asymmetries contribute to the asymmetric localization of Fop1.

### Poc1 is necessary for normal incorporation of Fop1 into BBs

Poc1 uniformly localizes around the BB circumference; yet, in poc1Δ cells, posterior-facing triplet microtubules preferentially disassemble (Pearson et al., 2009; Meehl et al., 2016). Because Fop1 localizes to these same posterior-
facing triplet microtubules, Poc1 and Fop1 may cooperatively stabilize this BB domain. To test such cooperativity, we measured Fop1: mCherry levels in poc1Δ cells. Fop1’s levels do not change in poc1Δ cells, and, similarly, Poc1: mCherry levels do not change upon FOPI knockout (Fig. 4 A). Thus, Poc1 and Fop1 are not required for each other’s BB localization. To determine whether Poc1 and Fop1 can promote each other’s BB incorporation, Poc1 or Fop1 was overexpressed and the BB levels of the reciprocal protein were measured. Poc1 overexpression increases Fop1 levels, whereas Fop1 overexpression does not change Poc1 BB levels (Figs. 4, B and C; and Fig. S2 I). This suggests that Poc1 promotes, but is not required for, Fop1 BB incorporation, and this functional interaction may impart stability to the posterior- and anterior-facing triplet microtubules.

There is a time delay between the start of BB assembly and the nucleation of a cilium, which provides a window to stabilize BBs before ciliary beating (Nanney, 1971; Bayless et al., 2012). Therefore, Poc1 may influence the timing of Fop1 incorporation to enhance BB stability before ciliary beating. To test this hypothesis, we measured the onset and subsequent maturation of endogenously tagged protein incorporation into BBs (Fig. 4, D–F). Daughter BBs form next to existing mother BBs and separate anteriorly as they mature. Thus, the distance between mother and daughter BBs is a proxy for BB maturation, and the protein fluorescence intensity ratio between the mother and daughter BBs reflects how much protein is incorporated during BB maturation. Fop1 incorporates early during BB assembly and levels increase rapidly during maturation, whereas Poc1 incorporates later and more slowly.
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Both Poc1 and Fop1 reach complete protein levels before achieving the mean separation distance of BB ciliogenesis (2.6 ± 0.1 µm). This suggests that these stability factors incorporate before ciliary force generation (Fig. S2 H). Interestingly, Fop1 incorporates more slowly in poc1Δ cells, whereas Poc1 incorporation is not affected by FOP1 knockdown (Fig. 4, E and F). These data are consistent with a model whereby a low level of symmetrically localized Poc1 facilitates the asymmetric recruitment of Fop1 to stabilize the posterior- and anterior-facing triplet microtubules of the BB in preparation for ciliary beating.

Microtubule glutamylation is asymmetrically enriched at and stabilizes BBs

BB and ciliary microtubules are posttranslationally modified by glutamylation. In flagellar axonemes, glutamylation asymmetrically localizes to the doublet microtubules that are in the plane of the ciliary beat stroke (Fouquet et al., 1996). It is not known whether such asymmetry exists at BBs, as observed for Fop1. To address this, we evaluated microtubule glutamylation via GT335 antibody staining. Glutamylated tubulin is found along the entire longitudinal length of the triplet microtubules. However, glutamylation is radially concentrated at the posterior-facing triplet microtubules.
microtubules (Fig. 5, A and B; and Fig. S3, A–D). SIM imaging of individual BBs reveals that glutamylation localizes in horseshoe- and foci-shaped profiles (Fig. 5 B). When averaged, glutamylation is concentrated at the BB posterior face (triplet microtubules 1, 2, 8, and 9) and at the anterior face (triplet microtubule 3, 4, and 5; Fig. 5 C and Fig. S3, A and C), which is consistent with immuno-EM localization (Fig. S3 B). The asymmetric positioning of tubulin glutamylation is not affected by the inhibition of ciliary beating (Fig. S3 C). Additionally, glutamylation is more centrally localized compared with Fop1 (Fig. S3 D). Despite variability in their precise localization, the overall asymmetric distributions of Fop1 and microtubule glutamylation are, to our knowledge, the first characterized molecular asymmetries with respect to the BB microtubule radial axis.

To determine when BB microtubules are glutamylated, we tracked tubulin glutamylation levels during BB maturation relative to the incorporation of BB stability factors (Fig. 5 D). BB glutamylation commences coincident with Fop1 and Bld10 incorporation but earlier than Poc1 incorporation. Furthermore, BB glutamylation incorporates more rapidly than Poc1. These data collectively indicate that the maturation of a stable BB occurs in multiple stages. Thus, asymmetric localizing proteins and PTMs incorporate early during BB maturation, and this is critical for BB stability.
Glutamylation has been reported to both stabilize and destabilize ciliary axonemes, whereas it primarily stabilizes cytoplasmic microtubules and centrioles (Bobinnec et al., 1998a,b, 1999; Wloka et al., 2010). At BBs, it is unknown whether glutamylation stabilizes microtubules against ciliary beating forces. To address this, BB frequency was quantified in cells lacking the two tubulin glutamylases, TTLL1 and TTLL9 (ttll1Δ and ttll9Δ), which specifically glutamylate BB microtubules but have no measurable effect on axonemal microtubule glutamylation (Fig. 5 E and F; and Fig. S3 E). BB frequency is decreased in cell cycle–arrested and cycling ttll1Δ cells (Figs. 5 G and S3 F). Moreover, BB loss in ttll1Δ,9Δ cells is repressed by ciliary inhibition with NiCl₂ treatment (Fig. 5 H). The rescue of BB loss by inhibition of ciliary beating suggests that glutamylation, like Poc1 and Fop1, stabilizes BBs against ciliary-dependent forces.

**Poc1, Fop1, and microtubule glutamylation act in distinct pathways to stabilize BBs**

Poc1 promotes the incorporation of Fop1 during BB maturation, indicating that there is interdependence among the stability factors for their incorporation into BBs. To determine whether such interdependence extends to microtubule glutamylation, BB glutamylation levels were quantified in poc1Δ and FOP1Δ knockout strains. Surprisingly, Poc1 or Fop1 loss increases BB glutamylation, whereas their overexpression decreases glutamylation (Figs. 5 I and S3 G). This suggests that BB glutamylation does not require Poc1 or Fop1. Instead, the increased glutamylation upon Poc1 or Fop1 loss might serve to compensate for BB instability in their absence. Alternatively, loss of Poc1 and Fop1 may promote the accessibility of TTLL-modifying enzymes to allow for additional tubulin glutamylation, which facilitates microtubule severing through katanin and spastin (Sharma et al., 2007; Lacroix et al., 2010; Valenstein and Roll-Mecak, 2016). Nonetheless, we find that microtubule glutamylation and BB stability factors have an inverse relationship with respect to their BB incorporation.

If BB glutamylation compensates for the loss of Poc1, then it follows that decreased glutamylation in poc1Δ cells should exacerbate BB instability. To test this hypothesis, and to exclude the possibility that increased glutamylation catalyzes BB instability in stability factor mutants, we created cells that were null for POC1, TTLL1, and TTLL9. Surprisingly, the linear BB frequency in cycling poc1Δ, ttll1Δ, ttll9Δ triple mutants is increased (Fig. S3 H). However, the total number of BBs per cell is drastically reduced, because the number of ciliary rows is decreased (Fig. S3 I). To specifically test whether existing BBs are less stable in the triple mutant, we assessed BB frequency in cell cycle–arrested cells. At 30°C, the triple mutant cells exhibit a 36%, 18%, and 24% decrease in BB frequency compared with wild-type (WT), poc1Δ, or ttll1Δ,9Δ cells, respectively (Fig. 5 J). Thus, BBs use temporally distinct but functionally overlapping strategies to maintain the normal complement of BBs in the cell.

**Conclusion**

How BBs manage the mechanical stress imposed upon them by asymmetric ciliary undulations remains poorly understood. This study illuminates an overlapping series of events required for the assembly of stable BBs capable of resisting such stress. These events rely on the timely incorporation of stabilizing proteins and microtubule PTMs at specific domains of the assembling BB. Bld10 and Poc1 symmetrically localize with respect to the BB rotational axis. These BB stability factors exhibit unique incorporation profiles that reflect their distinct functions in assembly and stability. Despite their symmetric localization, loss of triplet microtubules in bld10Δ and poc1ΔΔ mutants occurs asymmetrically along the plane of ciliary beating, suggesting that these regions experience greater mechanical stress (Bayless et al., 2012; Meehl et al., 2016). Together with the symmetrically localizing stability proteins, Fop1 and microtubule glutamylation reinforce these regions likely by asymmetrically localizing to the radial BB architecture. The coordination between uniquely incorporated and localized stability proteins reveals a dynamic and multidomain architecture of the BB that is responsible for resisting asymmetric ciliary beating.

**Materials and methods**

**T. thermophila cell culture**

T. thermophila strains were grown in 2% SPP media (2% protease peptone, 0.2% glucose, 0.1% yeast extract, and 0.003% Fe-EDTA) to mid-log phase at 30°C, unless otherwise indicated. Cells were considered mid-log phase at a density of ~3 × 10⁵ cells/ml as determined using a Coulter Coulter Z1 (Beckman Coulter). All temperature-shift experiments were performed for 24 h. Strains were cell cycle arrested in starvation media (10 mM Tris-HCl, pH 7.4). Perturbations that affect ciliary beating (3% PEO at all temperatures or 250 and 100 µM NiCl₂ at 30°C and 37°C, respectively; Larsen and Satir, 1991) were introduced 24 h after cell cycle arrest. The frequency of BBs was quantified 24 h after either PEO or NiCl₂ treatment. Ciliary inhibition by NiCl₂ treatment was confirmed by reduced cellular swimming.

**Mass spectrometry**

Proteins isolated by immunoprecipitation were identified by using an ion trap mass spectrometer (LTQ XL; Thermo Fisher Scientific) with a reverse phase gradient over C18 resin (Phenomenex). Analysis was performed using SEQUEST software as described previously (Washburn et al., 2001).

**Plasmids**

The Fop1:mCherry strain was generated by transforming cells with p4T2-1:FOP1:mCherry. This cassette integrates into the endogenous FOP1 locus and remains under control of the endogenous promoter. p4T2-1:FOP1:mCherry was generated by PCR amplification of an 825-bp fragment of FOP1 immediately upstream of the TGA stop codon (5'-CGGGTACCCATATCTCCT-3' and 5'-CGGAAATCTTAACTTTCAACAT-3') that was cloned into p4T2-1-mCherryLAP to generate p4T2-1-FOP1US-mCherryLAP (Winey et al., 2012). An 807-bp fragment downstream of the TGA stop codon (5'-CGGGATCCGAGAGTCCTTTTTT-3' and 5'-CGGAGCTCTTTGATCTCATC-3') was then cloned into p4T2-1-FOP1US-mCherryLAP plasmid intermediate to create p4T2-1:FOP1:mCherry. This plasmid contains the NEO2 drug selection marker. A FOP1:GFP construct was generated as described for p4T2-1:FOP1:mCherry using the p4T1-1-GEPLASp plasmid (Cheeseman and Desai, 2005). The fop1 knockout cassette was created by replacing the entire FOP1 open reading frame with the NEO2 resistance gene. This was achieved by cloning an 835-bp fragment of the FOP1 5' UTR (5'-CGGGTACCCATATCCTA-3' and 5'-CGAGCTGAGCTTTAATGC-3') and a fragment of the FOP1 3' UTR (5'-CGGGATCCGAGAGTCCTTTTTT-3' and 5'-CGGAAGCTCTTTGATCTCATC-3') into the p4T2-1 plasmid.

The FOP1 rescue construct (pBSMTTGF:FOP1) was generated by creating a genomic clone of GFP fused to FOP1 under control
of an inducible MTT promoter (Shang et al., 2002). The 1185-bp *FOP1* cDNA fragment was RT-PCR amplified (5′-CGTACGTAATA GAGAGATCAC-3′ and 5′-CGACTAGTCTATTAACCTTCA-3′) and cloned into pBSMTT:GFP:gtw (provided by D. Chalker, Washington University, St. Louis, MO) via a pENTR intermediate. This rescue cassette was transformed into *FOP1* knockout cells, integrating into the *rpl29* locus.

The overexpression constructs pBSMTTGF:PAS6a and pBSMTTGF:POC1 were generated as described for the pBSMTTGF:*FOP1*, by PCR amplifying (5′-CGGATCCGATGCTAGTTC ATC-3′ and 5′-CGAAGCTTCACTATTATTGTT-3′, and 5′-CGGATCCGATGCTAGTTC ATC-3′ and 5′-CGAAGCTTCACTATTATTGTT-3′, respectively). These genomic clones were then cloned into pBSMTTGFCHX.

**Macronuclear transformation**

Biological transformation was used to transform the p4T2-1-LAP constructs, pBSMTTGF:CHX overexpression constructs, and the *fop1* knockout cassette into the macronucleus of *Tetrahymena* cells (Bruns and Cassidy-Hanley, 2000). 200 µg/ml paromomycin was used to select for the *NEO2* gene, and 7.5 µg/ml cycloheximide was used to select for the *CHR* resistance gene (Gaertig et al., 1994; Hai et al., 2000). To increase the copy number of the endogenously tagged GFP and *mCherry* constructs, strains were selectively sorted by incrementally increasing the concentration of paromomycin in the media.

All GFP and *mCherry* constructs were assayed and maintained with paromomycin concentrations that maintained a strong signal to noise of the fluorescent tag. To ensure equal copy number of LAP constructs in experiments where *FOP1:mCherry* and *POC1:mCherry* levels were assessed in the presence or absence of *POC1* or *FOP1* (Fig. 4 A), respectively, strains were grown at the same paromomycin concentration for multiple passages. In experiments where *Sas6a, Poc1*, or *Fop1* were overexpressed by induction of the MTT promoter and levels of *Poc1* or *Poc1* were measured, “WT” denotes the identical MTTGF strains that were not induced with CdCl2 (Fig. 4, B and C).

**FOP1 knockout**

Biological transformation was used to insert the *fop1* knockout cassette into the macronuclear *FOP1* gene. The entire open reading frame of *FOP1* was targeted and replaced with the *NEO2* gene (Hai et al., 2000). Allelic assortment of the *NEO2* gene was performed by increasing the paromomycin concentration. Increased assortment of the knockout cassette decreases the copy number of WT *FOP1* alleles. Knockdown was verified by genomic PCR (42% reduction) and RT-PCR (58% reduction; Fig. S1, A and B).

**Immunoprecipitation**

BLD10:*mCherryLAP* or *POC1:*mCherryLAP constructs were transformed into B2086 *Tetrahymena* cells. Immunoprecipitation was performed as previously described (Cheeseman et al., 2004; Cheeseman and Desai, 2005). In brief, 2 l of confluent cells (10⁶ cells/ml) were washed in PBS and then resuspended in lysis buffer (50 mM Hepes, pH 7.4, 1 mM EGTA, 1 mM MgCl₂, 100 mM KCl, 10% glycerol, and 0.05% NP-40) with protease inhibitors. The whole-cell solution was dropped into LiNi{sub}₂ to produce drops of cell lysate. 5 g of cell drops was ground in a cryogrinder and resuspended in 1.5x lysis buffer and sonicated. Samples were centrifuged at 21,600 g, and the supernatant was collected before centrifugation at 135,000 g. Samples were immunoprecipitated with α-RFP–coated beads and eluted with 0.1 M glycine, pH 2.6 (Cheeseman et al., 2001).

**Light microscopy**

Fluorescence imaging was performed as previously described (Bayless et al., 2012) using a Nikon 100x Plan-Apochromat numerical aperture 1.4 objective at 25°C. Images were captured with a CMOS camera (Xyla 4.2; Andor Technology). All images were acquired using NIS Elements imaging software (Nikon). Acquisition times ranged between 50 and 500 ms, depending on the experiment. BB frequency was quantified as previously described (Bayless et al., 2012).

**Quantification of BB frequency**

Quantification of BB frequency was performed by counting the number of BBs along a 10-µm region along a ciliary row in the medial half of the cell. Ciliary rows around the entire circumference of the cell were quantified. All experiments used five measurements per cell over 20 cells. All experiments were repeated in triplicate for a total of 300 counts. All counts were corrected for cell length by multiplying the number of BBs by the ratio of mutant to WT cell length. In cycling cells grown in SPP, only nondividing, stage I cells were selected for analysis. In cell cycle–arrested cells, during cell starvation, a small fraction of cells become arrested in G2; however, the cells that are the focus of this study arrest in G1, assessed by cell size and BB frequency. These G1–arrested cells were used for all BB quantification in starvation experiments.

**Immunofluorescence**

Immunofluorescence was performed as described previously (Bayless et al., 2012). Cells were washed in PHEM buffer (60 mM 1,4-piperazinediethanesulfonic acid, 25 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid, 10 mM EGTA, and 2 mM MgCl₂, pH 6.9) and then fixed in formaldehyde fixative (3.2% paraformaldehyde and 0.2% Triton X-100, in PHEM buffer) for 5 min. Cells were washed three times in 0.1% BSA in PBS (BSA-PBS) before a 24-h incubation at 4°C with primary antibody diluted in 1.0% BSA-PBS. Primary antibodies used in this study were α-TiCen1 (Stemm-Wolf et al., 2005) and α-glutamylation (GT335; Adipogen; Wolff et al., 1992). Cells were then washed three times in 0.1% BSA-PBS before incubation for 2 h at 25°C in secondary antibody diluted in 1.0% BSA-PBS. Secondary antibodies used in this study were Alexa Fluor 488, 594, or 647 goat α-rabbit IgG, Alexa Fluor 488, 594, or 647 goat α-mouse IgG (Invitrogen). Cells were then washed three times in 0.1% BSA-PBS, and 1 µl of cell pellet was added to coverslip and mounted to a slide with Citifluor mounting media (Ted Pella). Samples were then sealed using clear nail polish.

**SIM**

SIM imaging was performed using the N-SIM system (Nikon) on a Ti Eclipse inverted microscope equipped with a 100x CF160 Apo-Chromat superresolution/TIRF NA 1.49 objective. Images were captured with an iXon DU897 X3 512 × 512 EMCCD (Andor Technology). Samples were excited using an N-SIM dual band filter cube for 488 and 561 nm excitation. SIM imaging was also performed on a DeltaVision OMX Blaze (GE Healthcare) superresolution system. The microscope was equipped with a 60x 1.42 NA point spread function objective. All images were acquired at 25°C using NIS Elements or DeltaVision software, respectively. Subtle resolution differences were observed between the N-SIM and OMX Blaze imaging systems. These likely accounted for the improved visualization of glutamylation tubulin fluorescent signal rings with the OMX Blaze system. Image analysis was performed using NIS Elements and ImageJ.
Fluorescence image averaging

Fluorescence averaging was performed using ImageJ’s built-in plugins and the ImageJ macro language. To generate the mean fluorescence intensity from widefield images, individual BB rows were rotated along a common anterior–posterior axis and boxes were manually centered over maximum intensity projections of each BB. Once the approximate central position of each BB was identified, boxes were automatically recentered over the highest intensity pixel and used to crop the images. These images were subsequently background subtracted and normalized so that the brightest pixel was set equal to 1. The resulting centered and normalized images were compiled into a new image stack, and the mean intensity image was generated using the mean intensity projection method in ImageJ. The method used to generate the mean fluorescence intensity from SIM images was identical, except the images were all manually centered over the area of lowest intensity within the interior portion of the ring- or horseshoe-shaped signal. To generate the linescans of mean widefield data, a line was centered over the mean BB image, and the resulting pixel intensities were fit to a Gaussian function using the ImageJ curve fitting plugin. To generate linescans of mean SIM data, a line or a circle was centered over the mean ring and the individual pixel intensities across all images were retrieved, which allowed statistical analyses to be performed across the linescan. All linescans were generated without resizing the images. However, some figures show smoothed images that were digitally enlarged with bicubic interpolation and displayed with the ImageJ “physics” lookup table.

Transmission EM

For immuno-EM, a C-terminal FOP1::GFP fusion under endogenous expression was prepared for immuno-EM using high-pressure freezing and freeze substitution (Dahl and Staehelin, 1989; Meehl et al., 2009). Incubation in rabbit-generated α-GFP antibodies, followed by incubation with α-rabbit secondary antibodies conjugated to 15-nm gold particles was used to localize Fop1-GFP. Fop1 was then localized in 60-nm sections by transmission EM. Images were acquired using a CM10 electron microscope (Philips) with a BioScan2 CCD camera (Gatan). For EM analysis of FOP1 knockdown BB structural defects, FOP1 knockdown cells were subjected to high-pressure freezing and freeze substitution as previously described (Pearson et al., 2009). Images were acquired using a Tecnai G2 (FEI) equipped with a Gatan Ultrascan digital camera. All images were processed for figures using Corel Draw.

Online supplemental material

Fig. S1 describes genomic and transcript levels of Fop1 in FOP1 knockdown cells, quantification of BB loss in cycling cells, quantification of BB loss with alternative BB markers, histograms of linear BB frequency and ciliary row counts corresponding to Fig. 1 C, and additional examples of FOP1 knockdown transmission EM phenotypes observed in Fig. 1 (D and E). Fig. S2 displays Fop1 localization relative to Cen1, Poc1, and Fop1’s N- and C-terminal–tagged proteins, SIM averaging of Fop1 localization using the Delavation OMX system, the asymmetric shift of Fop1 relative to separation distance between mother and daughter BBs, N-SIM averaged localization of Fop1 after inhibition of ciliary beating (Nicl treatment), longitudinal immuno-EM localization of Fop1::GFP, mean timing of ciliogenesis, Cen1::GFP incorporation profile, and quantification of ciliary glutamylation in ttl1Δ, poc1Δ triple mutants. The authors would like to thank Dr. Thomas H. Giddings for EM expertise and Dr. Iain Cheeseman for generous support with discussions and mass spectrometry analyses. We greatly appreciate Dr. Marisa Ruehle for critical comments on the manuscript and Kristin Dahl for generating the fop1Δ and FOP1::mCherry constructs. Thanks to Andrew Cahill and Nikon and Robert Moody and GE Healthcare for technical expertise with SIM microscopy.

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