An ARL3–UNC119–RP2 GTPase cycle targets myristoylated NPHP3 to the primary cilium

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The membrane of the primary cilium is a highly specialized compartment that organizes proteins to achieve spatially ordered signaling. Disrupting ciliary organization leads to diseases called ciliopathies, with phenotypes ranging from retinal degeneration and cystic kidneys to neural tube defects. How proteins are selectively transported to and organized in the primary cilium remains unclear. Using a proteomic approach, we identified the ARL3 effector UNC119 as a binding partner of the myristoylated ciliopathy protein nephrocystin-3 (NPHP3). We mapped UNC119 binding to the N-terminal 200 residues of NPHP3 and found the interaction requires myristoylation. Creating directed mutants predicted from a structural model of the UNC119–myristate complex, we identified highly conserved phenylalanines within a hydrophobic sandwich to be essential for myristate binding. Furthermore, we found that binding of ARL3-GTP serves to release myristoylated cargo from UNC119. Finally, we showed that ARL3, UNC119b (but not UNC119a), and the ARL3 GAP Retinitis Pigmentosa 2 (RP2) are required for NPHP3 ciliary targeting and that targeting requires UNC119b myristoyl-binding activity. Our results uncover a selective, membrane targeting GTPase cycle that delivers myristoylated proteins to the ciliary membrane and suggest that other myristoylated proteins may be similarly targeted to specialized membrane domains.

Keywords: ARL3; NPHP; UNC119; myristoylation

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Ciliopathies are rare, autosomal recessive disorders with phenotypes including cystic kidneys, retinal degeneration, neural tube defects, polydactyly, and obesity (for review, see Hildebrandt et al. 2011). The renal ciliopathy nephronophthisis (NPHP) and associated retinal–renal disorder Senior-Loken syndrome (SLS) are caused by mutation of any of 13 currently known disease genes (Wolf and Hildebrandt 2011). The proteins encoded by these genes localize to the microtubule-based primary cilium or the basal body/centrosome at the base of the cilium, leading to the hypothesis that defects in ciliary function underlie the pathology of NPHP/SLS.

The NPHP3 gene is mutated in ~1% of cases of NPHP (Olbrich et al. 2003). It encodes a protein with a conserved position 2 glycine, which we predict here to be N-terminally myristoylated, and has also been shown to localize to a distinct region of the primary cilium (Shiba et al. 2010). Loss-of-function mutation of Nphp3 results in embryonic lethality, and a hypomorphic mutation of Nphp3 is the causative mutation in the pcy mouse, a classic cystic kidney disease model (Olbrich et al. 2003). Directed protein–protein interaction studies have found that NPHP3 associates with other NPHP proteins, notably NPHP1 (Olbrich et al. 2003) and Inversin/NPHP2 (Bergmann et al. 2008). NPHP3 has been suggested to play a role in both canonical and noncanonical Wnt signaling, although the molecular details of these functions are not understood (Bergmann et al. 2008; Zhou et al. 2010).

The highly conserved small GTPase Arf-like 3 (ARL3) was identified in several screens to identify ciliary genes using different methods [Avidor-Reiss et al. 2004; Efimenko et al. 2005; Pazour et al. 2005]. ARL3 localizes to the primary cilium (Zhou et al. 2006), and knockout mice show renal and retinal involvement, a striking resemblance to...
Senior-Loken syndrome (Schrick et al. 2006). While the guanine nucleotide exchange factor (GEF) for ARL3 is unknown, the protein encoded by the retinitis pigmentosa disease gene Retinitis Pigmentosa 2 (RP2) has been shown to function as an ARL3-specific GTPase-activating protein (GAP) [Veltel et al. 2008]. Interestingly, RP2 has been shown to localize to the primary cilium (Hurd et al. 2011) and is implicated in trafficking proteins from the Golgi to the ciliary membrane [Evans et al. 2010]. ARL3 has three known GTP-specific interacting proteins [effectors]: BART, PDE6D, and UNC119 [Linari et al. 1999; Sharer and Kahn 1999; Van Valkenburgh et al. 2001]. Interestingly, UNC119, initially identified in Caenorhabditis elegans as an uncoordinated mutant showing broad neural deficiencies (Maduro and Pilgrim 1995), was identified in several ciliary screens as well, suggesting that ARL3 and UNC119 may function together in some aspect of ciliary biology (Avidor-Reiss et al. 2004; Li et al. 2004; Liu et al. 2007). UNC119 shares extensive sequence homology with PDE6D, which binds to and helps localize prenylated proteins in photoreceptors [Zhang et al. 2004, 2007], and a very recent study demonstrates that UNC119 shares many of the same biochemical properties, including binding to and extracting lipid-modified proteins from membranes [Zhang et al. 2011]. Vertebrate genomes contain two UNC119 genes, UNC119a and UNC119b, although the functional differences between the two genes have not been explored. The encoded proteins share a highly similar PDE6D-like domain, while the N-terminal domains have diverged considerably. UNC119a mutant mice show retinal degeneration [Ishiba et al. 2007], and a heterozygous truncation of UNC119a has been found in a patient with a cone–rod dystrophy [Kobayashi et al. 2000].

N-terminal myristoylation of proteins is thought to confer reversible membrane binding activity to the modified protein [Resh 1999]. How myristoylated proteins are targeted to the appropriate cellular locations remains unclear. Although the existence of a specific myristoyl–binding “receptor” has been proposed, such a protein has not been identified. Here we report that UNC119 is a myristoyl-binding protein that binds to a subset of myristoylated proteins, including NPHP3 and cystin, and that the small GTPase ARL3 regulates myristoyl–cargo binding. By using a structural model of the UNC119–myristoyl–peptide complex to guide mutagenesis, we identified key residues important for myristate binding. Cell-based NPHP3 ciliary localization assays indicate that ARL3, UNC119b and its myristoyl-binding activity (but not UNC119a), and the ARL3 GAP RP2 are required for proper ciliary targeting of NPHP3. We also showed that the myristoyl-binding activity of UNC119 is required in vivo as well. Furthermore, we identified C5orf30 as an UNC119-binding protein that preferentially binds unliganded UNC119. Finally, morpholino (MO)-mediated knockdown experiments in zebrafish are consistent with a role for the pathway in ciliary function. Based on our results, we propose that ARL3–UNC119b–RP2 constitute a GTPase cycle that targets a specific set of myristoylated proteins to the ciliary membrane.

Results

**UNC119 binds the N terminus of ciliopathy protein NPHP3 in a myristoyl-dependent manner**

We recently generated a protein–protein interaction network for the NPHP–JBTS–MKS disease spectrum, which included tandem affinity purification of an N-terminally tagged NPHP3 [Sang et al. 2011]. Sequence analysis predicts that NPHP3 is N-terminally myristoylated, which would be blocked by our N-terminal fusion. We sought to determine whether a C-terminally tagged, myristoylated NPHP3 would localize differently. Toward this end, we generated a stable retinal pigmented epithelial cell line (RPE) expressing C-terminally LAP-tagged murine NPHP3. In contrast to our N-terminally tagged construct (data not shown), the C-terminally tagged NPHP3 was able to localize to the primary cilium (Fig. 1A), suggesting, as seen for other ciliary proteins, including cystin [Tao et al. 2009], that myristoylation aids in proper NPHP3 targeting. Several attempts to generate a nonmyristoylated, G2A mutant NPHP3 stable cell line suitable for tandem affinity purification were unsuccessful, likely due to poor stability of the mutant protein. An intriguing possibility is that the myristoylation-dependent targeting may be important for accumulating NPHP3 in cilia.

To identify the factors that target NPHP3 to cilia, we used tandem affinity purification of our C-terminally tagged NPHP3. Purification of LAP-NPHP3 yielded several strong interacting proteins [Fig. 1B]. Excision and tandem mass spectrometry identified the 33-kDa and 31-kDa bands as UNC119a and UNC119b, respectively. We next confirmed the NPHP3:UNC119 interaction by GST pull-down using bacterially expressed GST-UNC119a and myc-tagged NPHP3 in vitro translated in wheat germ extract, a system known to be competent for myristoylation [Fig. 1C; Heuckeroth et al. 1988]. Using this assay, we subsequently localized the UNC119-binding domain of NPHP3 to the N-terminal 200 residues [Fig. 1C] and found that when expressed in RPE cells, this domain is sufficient for ciliary localization [Fig. 1E]. Attempts to determine the localization of the G2A mutant of the N-terminal NPHP3 fragment were unsuccessful due to poor stability, similar to results obtained with the full-length mutant construct. We next asked whether the interaction required myristoylation. Remarkably, mutation of the myristoylated position 2 glycine of NPHP3 to alanine abrogated binding [Fig. 1D]. These data suggested that UNC119 functions as a myristoyl-binding protein.

**UNC119 is a myristoyl-binding protein**

To further investigate whether UNC119 was most centrally a myristoyl-binding protein and to determine other functions of UNC119, we wanted to purify the two UNC119 isoforms and identify their interacting proteins. To identify these proteins, we generated stable RPE cell lines expressing LAP-tagged UNC119a and UNC119b, and also a murine inner medullary-collecting duct cell line (IMCD3) expressing LAP-tagged Unc119a [data not shown]. Interestingly, the LAP-UNC119a localized to the
centrosome, as determined by colocalization with pericentrin [Fig. 2A]. More importantly, the LAP-UNC119b was enriched at the transition zone and extended into the proximal end of the cilium [Fig. 2A]. Silver staining of the tandem affinity-purified UNC119a and UNC119b complexes shows an overlapping set of interactors as well as unique interactors [Fig. 2B]. Strikingly, one of the most abundant proteins present in each complex is NPHP3, reciprocally validating the NPHP3–UNC119a/b interactions. An affinity purification–mass spectrometry approach was taken for the IMCD3 LAP-Unc119a cell line in an effort to identify less abundant interacting proteins. In addition to Nphp3, the tandem affinity-purified Unc119a complex contained a strong enrichment of proteins (10 in total) either known or predicted to be N-terminally myristoylated [Fig. 2C]. These included the known ciliary protein cystin, which is mutated in the mouse cystic kidney model cpk [Hou et al. 2002]. Cystin also requires myristoylation for proper ciliary membrane localization [Tao et al. 2009]. To validate the interaction with cystin in cells, we generated a stable RPE cell line expressing the first 20 amino acids of cystin tagged on its C terminus, which is sufficient for targeting cystin to the cilium [Supplemental Fig. S1]. Tandem affinity purification of this cystin polypeptide yielded virtually stoichiometric amounts of UNC119a and UNC119b [Fig. 2D]. To confirm this was a direct interaction in vitro and to characterize the interaction, we synthesized C-terminally biotinylated peptides corresponding to cystin residues 2–20 either unmyristoylated (cystin) or myristoylated (Myr-cystin) on the N terminus to test for direct interaction. As shown in Figure 2E, the myristoylated cystin peptide efficiently coprecipitated in vitro translated UNC119a, whereas the nonmyristoylated peptide failed to interact with UNC119a. We next asked whether the NPHP3:UNC119a interaction could be competed by addition of the Myr-cystin peptide. Addition of increasing amounts of the Myr-cystin peptide, but not the unmyristoylated peptide, to a GST pull-down reaction containing GST-UNC119a and in vitro translated NPHP3 disrupted the NPHP3:UNC119 interaction [Fig. 2F]. These data strongly support the idea that UNC119 may function as a highly selective cargo adaptor for myristoylated proteins.

C5orf30 is an UNC119-binding protein that preferentially binds unliganded UNC119

In addition to the multiple myristoylated proteins that copurified with UNC119, we found an uncharacterized protein called C5orf30 that lacks a position 2 glycine and would not be predicted to be myristoylated [Fig. 2B]. Unlike the ARL3 and UNC119 genes, C5orf30 appears to be only found in vertebrate genomes. When GFP-tagged, C5orf30 localizes to the transition zone and proximal cilium in addition to being found throughout the cytoplasm [Supplemental Fig. S2A], consistent with a possible role in directing ciliary trafficking. In situ hybridization of mouse retinal sections show that C5orf30 is highly expressed in photoreceptors [Supplemental Fig. S2B], suggesting a role for C5orf30 in trafficking of proteins into the photoreceptors. We confirmed that C5orf30 is a direct UNC119-binding protein by GST pull-down assay with in vitro translated C5orf30 and GST-UNC119a [Fig. 2G].
regulate UNC119 binding to myristoylated proteins. We first determined whether C5orf30 binding to UNC119 was disrupted by the Myr-cystin peptide. Indeed, the UNC119a:C5orf30 interaction was disrupted by addition of the Myr-cystin peptide, but not the nonmyristoylated peptide (Fig. 2G). We wanted to consider the question of whether C5orf30 might reciprocally block myristoylated protein binding. We have yet to produce a form of the C5orf30 protein that would allow a simple competition assay. However, we did compare the relative efficiency of the binding among C5orf30, UNC119, and NPHP3 in our LAP purifications of each protein (Figs. 1B, 2B; data not shown). Consistent with our predictions, NPHP3 bound efficiently to UNC119, but we did not detect C5orf30 by silver staining. Reciprocally, purified C5orf30 binds UNC119, but we could not detect NPHP3. LAP-purified UNC119 bound to both C5orf30 and NPHP3 (Fig. 2A), which we believe to be a mixed population of C5orf30–UNC119 and NPHP3–UNC119 complexes. These data suggest that C5orf30 binds preferentially to unliganded UNC119. An intriguing possibility is that C5orf30 may act as a recycling factor to help extract UNC119 from the membrane after myristoylated cargo release.

Phenylalanine residues lining the hydrophobic β clamshell of UNC119 mediate myristate binding

The crystal structure of UNC119a was solved by the Northeast Structural Genomics Consortium [Protein Data Bank [PDB] 3GQQ], revealing a hydrophobic β sandwich...
fold, similar to other lipid-binding proteins such as PDE6D and RhoGDI. The length of the hydrophobic core is \( \sim 20 \text{ Å} \), which forms a tunnel that would readily accommodate the 14-carbon myristate moiety. Most striking are the 11 highly conserved phenylalanine residues lining the hydrophobic \( \beta \) sandwich [Fig. 3A]. We used structural modeling to place a myristate into the hydrophobic core [Fig. 3B], based on positioning with the tunnel domain circumscribed by these phenylalanines. Our model is remarkably similar to, and in full agreement with, the recently published cocrystal structure of UNC119a and a lauroylated peptide [Zhang et al. 2011]. Based on our model, we hypothesized that the phenylalanine residues lining the hydrophobic “tunnel” would be important for myristate binding, and that these phenylalanines cluster into four hydrophobic binding sites. To test our hypothesis, we generated four UNC119b mutant constructs, changing two to four phenylalanines to alanines in each of the sites. These constructs were in vitro translated and tested for myristoyl-binding activity using the biotinylated Myr-cystin peptide. As shown in Figure 3C, all four mutants displayed reduced binding to the Myr-cystin peptide, with the Mut-4 mutant [F144A, F148A, and F207A] having the greatest effect. Importantly, each of the phenylalanine mutants were able to bind to ARL3 QL as efficiently as wild-type UNC119b, supporting that the overall architecture of UNC119b is not disrupted by our mutagenesis (Fig. 3C). Next, we mutated each of the phenylalanines in the Mut-4 site individually. We found that while F144 contributes little to Myr-cystin binding, F148 and F207 are critical for binding to the Myr-cystin peptide [Fig. 3D], validating our structural model.

Ciliary small GTPase ARL3 binds UNC119 and membranes in a GTP-specific fashion in RPE cells

UNC119 is a known GTP-specific effector of ARL2 and ARL3 [Van Valkenburgh et al. 2001]. We focused our attention on ARL3 because it localizes to the primary cilium in RPE cells [Zhou et al. 2006], while ARL2 does not [data not shown]. We generated stable RPE cell lines expressing LAP-tagged GDP-mimicking ARL3 TN and the GTP-mimicking ARL3 QL mutant proteins. In contrast to other ciliary small GTPases, such as Rab8, both the GDP- and GTP-mimicking mutant forms of ARL3 localized to the primary cilium (Supplemental Fig. S3),

Figure 3. Mutations in the highly conserved phenyl groove of UNC119 disrupt myristoyl–peptide binding. (A) Sequence alignment of human UNC119b and the Chlamydomonas homolog POC7. Phenylalanine residues mutated in C are indicated and color-coded. (B) Structural model of UNC119a and the myristoylated N-terminal three amino acids of cystin based on the UNC119a crystal structure 3GQQ. The four groups of conserved phenylalanines mutagenized in C are shown and color-coded. Residue numbers correspond to the UNC119b sequence. Portions of the structure were removed to allow visualization of the myristoyl-binding pocket. (C, top immunoblot) Biotinylated Myr-cystin-binding assays were performed as in Figure 2E only using in vitro translated wild-type [wt] or mutant Myc-tagged UNC119b. The bottom immunoblot shows the results of a binding reaction between GST-ARL3 QL and in vitro translated Myc-tagged UNC119b. (D) Biotinylated Myr-cystin-binding assays were performed as in Figure 2E only using in vitro translated wild-type [wt] or mutant Myc-tagged UNC119b.
suggesting that the ARL3 GTPase cycle may function to release proteins within the cilium, rather than regulating ciliary entry. Tandem affinity purification of LAP-ARL3 TN yielded no strong interacting proteins, while the LAP-ARL3 QL copurified with six additional protein bands (Fig. 4A). Mass spectrometry was used to identify each of the proteins. We found all of the known ARL3 effectors (BART, PDE6D, and UNC119) as well as two novel effectors of unknown function (CCDC104 and C20orf194) and also the ARL3 GAP RP2 (Fig. 4A). The ARL3 effectors were assayed by GST pull-down using either GST-ARL3 TN or GST-ARL3 QL along with the in vitro translated effector (Fig. 4B), and, indeed, each of these candidate effectors bound in a GTP-dependent manner. These novel effectors will not be discussed further and will be characterized in future work.

ARL3 QL releases myristoylated peptides from UNC119

We next considered whether the ARL3 interaction would regulate UNC119 interactions with myristoylated proteins. First, we tested whether the cystin N-terminal peptide would disrupt the known interaction between UNC119 and the GTP-bound small GTPase ARL3. As shown in Figure 4D, addition of the cystin peptide to a binding reaction containing GST-UNC119a and in vitro translated ARL3 QL had no effect. This result confirms the specificity of the competitive binding seen between NPHP3 and the myristoylated cystin peptide. Moreover, like many ARF/ARL family proteins, ARL3 contains a position 2 glycine and would itself be predicted to be myristoylated. Myristoylation of these factors would help direct the ARF/ARL GTPase to the membrane. Here, it appears that ARL3 binds UNC119 only via its GTP effector function and not via a myristoylation signal.

We next tested whether ARL3 QL would release myristoylated cargo from UNC119. We preformed complexes of in vitro translated UNC119a or UNC119b and the biotinylated Myr-cystin peptide on streptavidin-sepharose beads and then added recombinant GST-ARL3. Remarkably, addition of increasing amounts of GST-ARL3 QL, but not the addition of GST-ARL3 TN, leads to a dose-dependent release of UNC119 from the immobilized Myr-cystin peptide (Fig. 4E). Interestingly, we found that despite binding to UNC119 similarly to ARL3 QL, ARL2 QL was unable to release UNC119 from the Myr-cystin peptide (Fig. 4E). These data suggest that ARL3 QL facilitates release of myristoylated cargo from UNC119 into the appropriate membrane domain. Given that PDE6D shares sequence similarity with UNC119...
and also binds to ARL2/ARL3 and lipid-modified proteins, it is likely that ARL2 or ARL3 will similarly regulate PDE6D cargo binding.

The ARL3 pathway is required for NPHP3 ciliary localization

Having established a biochemical relationship between ARL3, UNC119, and myristoylated ciliary proteins (NPHP3 and cystin), we sought to test the model that ARL3/UNC119 are responsible for targeting myristoylated ciliary proteins to the primary cilium. To test this idea, we generated a stable RPE cell line expressing LAP-tagged NPHP3 1–200. We chose this N-terminal fragment of NPHP3 over the full length because it exhibited notably more efficient and consistent ciliary localization, making this variant more suitable for a ciliary targeting assay. We transfected these cells with either control scrambled siRNA or siRNAs targeting ARL2, ARL3, UNC119a, UNC119b, or RP2 (Fig. 5A,B). We used two distinct siRNAs for ARL3, UNC119a, UNC119b, and RP2 to minimize the chance of off-target effects. All siRNAs depleted the endogenous transcripts to <15% of control levels [Supplemental Fig. S4]. As expected, depletion of ARL2 had no effect. Strikingly, knockdown of ARL3, UNC119b, and RP2 each dramatically reduced the percentage of cilia with localized NPHP3 1–200. We observed very low amounts of our LAP-NPHP3 1–200 protein in knockdown cilia when imaged at longer exposures (<10-fold wild-type levels), likely reflecting a combination of the level of overexpression of our reporter construct and possibly incomplete knockdown of the targeted protein. Interestingly, depletion of UNC119a had no effect even though it is able to bind NPHP3. We suspect that this is due to the difference in localization between UNC119a and UNC119b in RPE cells, but may reflect more broadly on functional differences between UNC119a and UNC119b [see the Discussion]. Depletion of ARL3 and UNC119b in a stable cell line expressing GFP-tagged cystin 1–20 yielded similar results (data not shown). Further, knockdown of ARL3, UNC119a, UNC119b, or RP2 showed no effect on the trafficking of ciliary G-protein-coupled receptors [Supplemental Fig. S5], indicating that delivery of nonmyristoylated proteins to the cilium is not disrupted by the depletion of ARL3 pathway components.
Myristoyl-binding activity of UNC119b is required to localize NPHP3 to the primary cilium

Having established that the N-terminal fragment of NPHP3 requires UNC119b for proper ciliary localization, we wondered whether this localization would require UNC119b myristoyl-binding activity. To test this hypothesis, we performed a series of siRNA/rescue experiments in our stable LAP-NPHP3 1–200 cell line using tdTomato-tagged UNC119b constructs and a siRNA targeting the UNC119b 3′ untranslated region (UTR) (Fig. 5C,D). As expected, the wild-type tdTomato-UNC119b construct was able to completely rescue localization of NPHP3 to the cilium in UNC119b-depleted cells. Strikingly, the myristoyl-binding mutant tdTomato-UNC119b Mut-4 failed to rescue to the same extent, indicating that NPHP3 ciliary targeting indeed requires the myristoyl-binding activity of UNC119b. Interestingly, the myristoyl-binding mutant UNC119b was found only at the basal body/transition zone (Fig. 5C, open arrowhead), whereas the wild-type version was present in the proximal cilium (Fig. 5C, closed arrowhead) as well as the basal body/transition zone. An intriguing possibility is that when lacking its myristoylated protein cargo, UNC119b may only traffic to the basal body, and that cargo binding may trigger ciliary entry of UNC119b.

Myristoyl-binding activity of UNC119b is required in C. elegans

Unc-119 mutant worms exhibit defects in locomotion (Maduro and Pilgrim 1995), and sensory neurons in these mutants fail to fill with lipophilic dye (see below). We asked whether human UNC119B with mutations in the myristoyl-binding site (Mut-4) would be able to rescue C. elegans unc-119 phenotypes. Wild-type human UNC119B, but not Mut-4, was able to rescue both the locomotor and head sensory neuron dye-filling defects of unc-119 mutant animals, while Mut-4 appeared to have a dominant-negative activity in dye-filling sensory neurons in the tail (Fig. 6A,B). These data suggest that UNC119 family proteins require myristoyl-binding activity in vivo.

Ciliary function in zebrafish requires nphp3, unc119a, and c5orf30

Zebrafish have become a useful model organism used to study basic ciliary function. MO-mediated knockdown of genes involved in ciliary function typically result in curved body, defects in the size of the ciliated embryonic structure known as the Kupffer’s vesicle (KV), and reduced visual function (Yen et al. 2006; Pretorius et al. 2010). Zebrafish embryos injected at the one- to four-cell stage with gene-specific MOs were assessed for overall morphology, KV size, and visual function. Previous work has noted ciliary roles for nphp3 and unc119a (most similar to human UNC119B) (Manning et al. 2004; Zhou et al. 2010). We found that knockdown of zebrafish nphp3, unc119a, and c5orf30 each resulted in a MO dose-dependent curved body axis and KV defects (Fig. 6C,E). Moreover, we found that nphp3 and unc119a knockdown results in reduced visual function (Fig. 6D,E). Consistent with the results from our cell-based NPHP3 targeting assay and the differences in localization seen between human UNC119a and UNC119b, knockdown of zebrafish unc119b (most similar to human UNC119a) yielded only a mild KV defect but no other obvious cilia-related phenotypes. Finally, in zebrafish, we identified three genes with high homology with ARL3: arl3 (95%), the syntenic gene to human ARL3, and two arl3-like factors, arl3l1 (75%) and arl3l2 (77%), that are not found in humans or mice. Knockdown of the syntenic arl3 using two independent MOs resulted in no obvious cilia-related defects, suggesting possible subfunctionalization of arl3 family members within the zebrafish (data not shown). Taken together, these data demonstrate that nphp3, unc119a, and c5orf30 have a strong role in ciliary function in the zebrafish.

Discussion

We describe here data showing that the highly conserved UNC119 proteins recognize the myristoylated N terminus of a subset of proteins, including the ciliopathy protein NPHP3 and cystin. Mutagenesis based on our structural model of the UNC119–myristate complex allowed us to identify four groups of conserved phenylalanine residues that are important for myristoyl–peptide binding, with two key phenylalanines being critical for binding. We then turned our attention to the ciliary small GTPase ARL3. By using a proteomic approach, we identified a comprehensive list of GTP-specific effector proteins, including the known ARL3 effectors BART, PDE6D, and UNC119, as well as two proteins of unknown function (CCDC104 and C20orf194). By fractionating our cell lines expressing GDP- and GTP-mimicking forms of ARL3, we found that, similar to other ARF and ARF-like family members, ARL3 bound to membranes GTP-specifically. Next, we found that ARL3 binding, but not ARL2 binding, released a myristoylated peptide from UNC119. Based on knockdown experiments using our cell-based assay monitoring ciliary localization of the myristoylated N-terminal UNC119-binding fragment of NPHP3, we suggest that the ARL3 GTPase pathway (ARL3-GTP, UNC119b, and GAP RP2) is required to target myristoylated ciliary proteins to the ciliary membrane.

Model of myristoylated ciliary protein trafficking

Based on the data presented, we propose a model (Fig. 7) in which UNC119b first binds a subset of myristoylated cargo proteins in the cytoplasm. Because myristoylation occurs cotranslationally (Wilcox et al. 1987), this binding may happen shortly after the initiation of translation to prevent mistargeted myristate–membrane interactions (see below). The UNC119b–myristoylated cargo complex would then be targeted to the cilium by a currently unknown mechanism. Also working at an early step in the process, the ciliary small GTPase ARL3 would be activated by a currently unknown GEF. This activation would potentially occur within the cilium because both the

![Figure 6](https://example.com/fig6.png)

**Figure 6** Ciliary localization of the myristoylated N-terminal component of the UNC119 family. **A**. ARL3-GAP RP2-GFP (green) localization in human cells expressing LAP-NPHP3 1–200. **B**. ARL3-GAP RP2-GFP localization in human cells expressing LAP-NPHP3 1–200 and siRNA targeting UNC119. **C**. ARL3-GAP RP2-GFP localization in human cells expressing LAP-NPHP3 1–200 and siRNA targeting nphp3. **D**. ARL3-GAP RP2-GFP localization in human cells expressing LAP-NPHP3 1–200 and siRNA targeting unc119a. **E**. ARL3-GAP RP2-GFP localization in human cells expressing LAP-NPHP3 1–200 and siRNA targeting c5orf30.

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GTP- and GDP-locked forms are ciliary, although the site of this activity is not truly clear. Once in the cilium, membrane-bound ARL3-GTP binds to UNC119b, releasing the myristoylated cargo into the cilium/ciliary membrane. The ARL3 GAP RP2, which is known to reside within the ciliary membrane (Hurd et al. 2011), activates the ARL3 GTPase activity, thus releasing UNC119b (Veltel et al. 2008). Interestingly, we obtained multiple peptides identifying RP2 as present in our purification of NPHP3 (data not shown). It is tempting to speculate that the cargo itself may lend specificity to target membrane deposition by containing sequences that bind the GAP.

Based on our competitive binding experiments with C5orf30 and the Myr-cystin peptide, we propose that C5orf30 binds preferentially to unliganded UNC119. Based on this activity, and our observation that C5orf30 can reside in the primary cilium (Supplemental Fig. S3A), C5orf30 may aid in either the release of cargo or recycling of UNC119b.

As we were preparing our paper, Zhang et al. (2011) published a study reporting that UNC119 binds the myristoylated Ga protein Transducin and showed biochemically that UNC119 facilitates its extraction from rod outer segment membranes. While we considered a possible GDI-like “extraction” mechanism for UNC119, our results showing that NPHP3 fails to localize to the primary cilium when UNC119b is depleted, rather than accumulating in the ciliary membrane, suggest a membrane

**Figure 6.** Myristoyl-binding activity is required to rescue *C. elegans unc-119 (ed)* mutants, and knockdown of nphp3, unc119a, and c5orf30 in zebrafish results in cilia-related phenotypes. [A] Locomotor activity of wild-type, unc-119 (ed3) mutant, and unc-119 (ed3) mutant worms rescued with either wild-type or Mut-4 human UNC119b. An asterisk (*) and hash mark (#) denote values that are significantly different ([P] < 0.05) from wild-type and unc-119(ed3) values, respectively. [B] The ability of head and tail sensory neurons of animals of the indicated genotypes to fill with Dil dye was analyzed and graphed. An asterisk (*) and hash mark (#) denote values that are significantly different ([P] < 0.05) from wild-type and unc-119(ed3) values, respectively. [C] Representative images of MO-injected zebrafish embryos at 48 h post-fertilization (hpf). [D] Representative images of low-dose MO-injected zebrafish at 5 dpf post-fertilization (dpf) used in the vision assay. Note that a lower amount of MO was necessary, since embryos with a curved body cannot be used in the vision assay. [E] Table reporting results from KV analysis and vision assay. (*) [P] < 0.01; (**) [P] < 0.05, Fisher’s exact test.
targeting function. It will be interesting to determine what factors dictate UNC119’s role in membrane extraction versus delivery.

**UNC119a versus UNC119b**

The UNC119a and UNC119b proteins share a highly similar PDE6D-like domain, comprising more than three-quarters of each protein, which we showed here binds to the myristate moiety of a subset of N-terminally myristoylated proteins. Despite this high degree of similarity, we found that UNC119b, but not UNC119a, is required for proper ciliary NPHP3 localization. In contrast, the N-terminal ~50 residues of the two orthologs have diverged considerably. Interestingly, we found that when expressed in RPE cells, UNC119a localized to the centrosome, while UNC119b localized to the transition zone and proximal cilium. The ciliary targeting sequence thus may lie within the divergent N terminus of UNC119b, although experiments to test this hypothesis will be necessary. We also noted differences in the banding pattern of our silver-stained gels comparing the purified UNC119a and UNC119b complexes. The divergent N-terminal domains may additionally contribute to the selectivity of myristoyl–peptide binding, accounting for the differences observed between the two purified complexes. Furthermore, in zebrafish, knockdown of unc119a lead to three cilia-related phenotypes (curved body, KV defect, and vision impairment), while unc119b depletion displayed only a mild KV defect, supporting differential roles for unc119a and unc119b in vivo. Additional studies addressing the binding partners, specificity, and mouse knockout phenotype of UNC119b would be important follow-on studies.

**UNC119 specificity and functional orthologs**

Approximately 0.5% of the human genome is predicted to be myristoylated (Maurer-Stroh et al. 2004), and >50 proteins in the flagellar and basal body (FABB) proteome, a list of candidate ciliary proteins (Li et al. 2004), possess position 2 glycines (our observation). A subset of these myristoylated proteins contains a cysteine at position 3, predicted to also be palmitoylated, including RP2. Based on our structural model, the β sandwich fold is unlikely to accommodate a second long hydrophobic lipid chain, and thus doubly acylated proteins are unlikely to be candidates for the ARL3–UNC119 targeting pathway. Notably, all of the predicted myristoylated proteins identified in our UNC119 purifications lacked a position 3 cysteine. Our purifications of UNC119a and UNC119b suggest that they recognize only a subset of the total myristoylated protein population, although future purifications from different cell types or tissues may expand the current repertoire. It is intriguing to consider whether there are functional equivalents of UNC119 for other myristoylated proteins.

**UNC119 as a chaperone**

In addition to acting as a ciliary targeting factor for a subset of myristoylated proteins, UNC119 may also act as a chaperone, burying the myristate until the cargo reaches its appropriate target membrane. Interestingly, other myristoylated proteins, such as HIV gag and c-abl, are able to bury their myristate moieties within an internal protein domain (Hantschel et al. 2003; Tang et al. 2004). In the “myristate switch” mechanism, the HIV gag myristate is exposed only when the protein reaches its destination at the apical membrane (Tang et al. 2004). It may be that proteins unable to bury their myristate will require UNC119 or UNC119-like proteins for proper membrane localization. Here, especially as these proteins are conserved to *Chlamydomonas*, the function of an independent myristate-binding protein would appear to be an ancient device for membrane targeting.
Strong evolutionary conservation of the ARL3 pathway

The ARL3 and UNC119 genes are conserved from humans to the green algae Chlamydomonas (called ARL3 and POC7, respectively). The Chlamydomonas genome also contains at least two RP2-like proteins [Stephan et al. 2007]. While the ciliary cargos identified here (NPHP3 and cystin) are not obviously conserved, myristoylated proteins appear to be enriched in biochemical preparations of Chlamydomonas flagellar proteins [Pazour et al. 2005]. This remarkable degree of conservation suggests an ancient origin of the myristoylation-based membrane targeting mechanism described here.

New candidates for renal and retinal degeneration (NPHP/SLS) genes

NPHP and SLS are autosomal recessive genetic disorders characterized by cystic kidneys and cystic kidneys plus retinal degeneration, respectively. Currently, there are 13 genes known to cause NPHP or SLS when mutated, with mutation of several of these genes leading to both diseases. These known disease alleles account for only ~30% of diagnosed cases of NPHP, suggesting that many more disease genes remain to be identified. Based on the data presented here, we propose that UNC119b and C5orf30 are strong candidates for NPHP or SLS disease genes. Additionally, patients with retinitis pigmentosa or the pediatric retinal disease Leber’s congenital amaurosis would be candidates, having lesions in these genes. Future studies will be necessary to confirm these two genes as bona fide ciliopathy genes.

Materials and methods

Plasmids

Gateway entry clones for human ARL3, BART, BBS1, CCDC104, PDE6D, and UNCI19a were obtained from Invitrogen. C5orf30, cystin 1–20, C20orf194, and UNCI19b were synthesized by GeneWiz. Tandem affinity purifications were performed essentially as described (Mukhopadhyay et al. 2010). Briefly, cells stably expressing a LAP-tagged construct were lysed in 50 mM HEPES (pH 7.4), 300 mM KCl, 1 mM EGTA, 1 mM MgCl2, 0.5 mM DTT, 0.3% NP-40, 0.2 mM PMSF, and 1× Roche Complete protease inhibitors. Lysates were cleared at 27,000 × g, and then again at 100,000 × g. Complexes were immunoprecipitated with rabbit anti-GFP antibodies and eluted with either TEV [LAP6] or PreScission [LAP5 and LAP7] protease. Complexes were again captured by using S protein agarose (Novagen), and finally eluted in 1× SDS sample buffer.

Recombinant proteins

Recombinant GST, GST-ARL3 TN, GST-ARL3 QL, and GST-UNC119a were expressed in Rosetta 2 cells [EMD Biosciences], induced with 0.2 mM IPTG at overnight 18°C, and purified using glutathione-sepharose according to the manufacturer’s recommendations [GE Healthcare Life Sciences]. In vitro translated proteins were generated by using the TNT HighYield Wheat Germ Protein expression system following the manufacturer’s recommendations [Promega].

GST pull-down assays

For each pull-down reaction, 2.5 μg of GST or GST fusion protein was added along with 5–10 μL of in vitro translated protein and 5 μL of glutathione-sepharose in binding buffer (300 μL 25 mM HEPES at pH 7.4, 150 mM NaCl, 2 mM MgCl2, 1% Triton X-100) for 1 h at 4°C. The beads were washed five times with the same buffer and eluted with SDS sample buffer. The eluates were resolved by SDS-PAGE and analyzed by immunoblotting with anti-myc- or anti-His antibodies. For competition assays, binding reactions were performed as described above, only 0.1 μM, 1 μM, 5 μM, or 10 μM peptide was included.

ARL3-UNC119 targets NPHP3 to the cilium

line containing a stably integrated FRT cassette was transfected with pOG44 encoding FLP recombinase [Invitrogen] and the appropriate LAP vector. Cells were then selected with the appropriate antibiotic.

Microscopy

Cells were plated on coverglass and induced to ciliate by withdrawing serum. Twenty-four hours later, the cells were cold-treated for 15 min on ice to depolymerize nonciliary microtubules, fixed 10 min in 4% formaldehyde in PBS, and washed three times with PBS. Coverslips were blocked in immunofluorescence buffer [PBS plus 30 mg/mL BSA, 0.1% NP-40] plus 5% normal donkey serum [Jackson ImmunoResearch]. Cilia and basal bodies/centrosomes were visualized by immunostaining with 6-11b-1 anti-acetylated tubulin [1:2000, Sigma] and rabbit anti-pericentrin [1:2000, Abcam] antibodies in immunofluorescence buffer, respectively. Cy3 anti-rabbit and Cy5 anti-mouse secondary antibodies [Jackson ImmunoResearch] were added at 1:2000 in immunofluorescence buffer for 30 min after five washes with immunofluorescence buffer. Hoechst 33342 [AnaSpec, Inc.] was added for 5 min, and the coverslips were washed five times with immunofluorescence buffer and mounted in Fluoromount G. Images were acquired on an Everest deconvolution workstation [Intelligent Imaging Innovations] equipped with a Zeiss AxioImager.Z1 microscope, a CoolSnapHQ cooled CCD camera [Roper Scientific], and a 40× PlanApochromat NA 1.3 objective or 63× PlanApochromat NA 1.4 objective.

Tandem affinity purification

Tandem affinity purifications were performed essentially as described [Mukhopadhyay et al. 2010]. Briefly, cells stably expressing a LAP-tagged construct were lysed in 50 mM HEPES (pH 7.4), 300 mM KCl, 1 mM EGTA, 1 mM MgCl2, 0.5 mM DTT, 0.3% NP-40, 0.2 mM PMSF, and 1× Roche Complete protease inhibitors. Lysates were cleared at 27,000 × g, and then again at 100,000 × g. Complexes were immunoprecipitated with rabbit anti-GFP antibodies and eluted with either TEV [LAP6] or PreScission [LAP5 and LAP7] protease. Complexes were again captured by using S protein agarose [Novagen], and finally eluted in 1× SDS sample buffer.
Peptide-binding assays

Peptides were synthesized and purified by reverse-phase chromatography by Elim Biopharmaceuticals, corresponding to the human cystin protein. More than 95% of the Myr-cystin peptide contains the myristoyl moiety. Peptide-binding reactions were performed by adding 450 fmol of C-terminally biotinylated peptide with 5–10 μL of Wheat Germ Extract IVT (Promega) protein and 5 μL of streptavidin-sepharose (GE Healthcare Life Sciences) in 300 μL of binding buffer for 1 h at room temperature. The beads were washed five times with binding buffer and eluted with SDS sample buffer. For the peptide release assays, 450 fmol of biotinylated Myr-cystin peptide was incubated with 5–10 μL of in vitro translated protein and 5 μL of streptavidin-sepharose in 300 μL of binding buffer for 1 h at room temperature. After two washes with binding buffer, GST-ARL2 or GST-ARL3 was added to 5.5 nM, 11 nM, 22 nM, or 44 nM, with a final volume of 300 μL. After a 1-h incubation, the complexes were washed and analyzed as described above.

Structural modeling

To build our structural model, we used the structure of truncated UNC119 (PDB code 3GQQ) as a reference. Docking of the myristoylated peptide was performed in two steps. First, the myristoyl group was placed into the cavity surrounded by the phenyl grove of UNC119, and energy was minimized using the interactive modeling program Benchware3D (Tripos). The choice of orientation of the myristoyl polar tail was made so that space is available to connect the peptide. The peptide was then built out of the myristate ester and extended on the protein surface to complement the surface charge and shape of UNC119. The peptide was then truncated back to the five N-terminal residues to yield the model presented here. The illustration of the model was made using MacPyMOL.

Cellular fractionation

Stable cell lines expressing LAP-tagged ARL3 TN or ARL3 QL were fractionated by first lysing the cells in hypotonic buffer and dialyzing. Nuclei were pelleted at 1400g and discarded. The supernatant was then separated into a soluble fraction (S100) and an insoluble pellet (P100) by centrifugation at 100,000g and supernatant was then separated into a soluble fraction (S100) and an insoluble pellet (P100) by centrifugation at 100,000g. Equal amounts of each were then separated by SDS-PAGE and analyzed by immunoblotting with an anti-GFP antibody (Abcam).

NPHP3 trafficking assay and rescue experiments

The LAP7-NPHP3 1–200 stable cell line was plated on coverslips overnight and transfected with siRNA with Lipofectamine RNAi Max [Invitrogen] following the manufacturer’s recommendations. After 48 h, the cells were serum-starved with 0.2% FBS-containing medium. Twenty-four hours later, the cells were processed for immunofluorescence as described above. Images from each sample were collected and analyzed using identical settings. Percent ciliary NPHP3 was determined by first counting the number of NPHP3-positive cilia (viewing only the Cy3 channel for pericentrin and GFP channel for NPHP3), and then counting the total number of cilia as determined by Cy5 acetylated tubulin immunostaining. Rescue experiments were performed by first transfecting cells with the rescue plasmid using the Amaxa Nucleofection Kit V and plating them on coverslips. The following day, the cells were transfected with siRNAs as described above. After 24 h, the cells were serum-starved for 24 h to induce ciliation, processed for imaging, and analyzed as described above. The following siRNAs used were purchased from Dharmaco: ON-TARGETplus nontargeting control, ON-TARGETplus SMARTpool targeting ARL2, ON-TARGETplus UNC119a oligos J-009734-07 and J-009734-08, ON-TARGETplus ARL3 oligos J-011813-06 and J-011813-07, ON-TARGETplus UNC119b oligos J-035010-15 and J-035010-16, and ON-TARGETplus RP2 oligos J-012350-08 and J-012350-09. UNC119b oligo J-035010-15 targets the 3’ UTR and was used in the rescue assay.

C. elegans studies

Strains The wild-type strain was C. elegans Bristol, strain N2. Additional strains were WP38 unc-119(ed3) (Caenorhabditis Genetics Center) and CX3553 lin-15(n765ts), klyl104 [str-1p::gfp+lin-15(+)] (Troemel et al. 1997).

Molecular biology Ce unc-119p::Hs unc119b cDNA::Ce unc-119 3’ UTR constructs were generated by sequential PCR fusions (Hobert 2002). The fusions contain 2.56 kb and 1.27 kb of sequences upstream and of downstream from, respectively, the Ce unc-119 coding region [Maduro et al. 2000]. Primer sequences are available on request. PCR fusion products were sequenced to confirm the absence of PCR-induced mutations. Transgenic strains were generated by injecting plasmids at 100 ng/μL together with unc-122::dsRed as the coinjection marker.

Dye-filling assay Animals were dye-filled with DiI as described previously [Perkins et al. 1986; Herman and Hedgecock 1990]. Individual animals were placed on agaroose slides and examined under 400X magnification on a Zeiss Axioplan microscope.

Tracking of locomotion Well-fed 1-d-old adult animals were placed on a freshly seeded plate and their movements were recorded for 15 min as described (Biron et al. 2008; Beverly et al. 2011). Velocities of individual animal movements were extracted using custom-written scripts in MatLab (The MathWorks, Inc.).

Statistical analyses Statistical analyses were performed using the SPSS 19 statistical analysis package.

Zebrafish studies

The following site antisense MOs were designed and purchased from GeneTools: unc119a, AGCTCTGACTTCTCAT CTCTCGGAA; unc119b, GCTGTTAACAAGATAGCCTATTATA CT; Cson80, AGCTCTCACCTGACATCATTCCAT; and nphp3, TCACCAGAGAAGTCGGCCGTACCCAT. The MOs were air pressure-injected into one- to four-cell embryos at the amounts indicated in Figure 6.

Analysis of KV size The size of the KV was assessed on live embryos at the six- to 12-somite stage of development as previously described [Yen et al. 2006]. Briefly, embryos with a KV smaller than the width of the notochord were scored as abnormal.

Vision startle response assay The light-induced vision startle response behavioral assay was performed as previously described [Pretorius et al. 2010]. In brief, 5-d-post-fertilization [dpf] embryos were exposed to a 1-sec block in bright light intensity as a visual stimulus. A positive visual response was recorded if the embryo made an abrupt change in swimming behavior immediately upon light change. Five trials were performed, spaced 30 sec apart, followed by the mechanical stimulus of probing embryos with a blunt needle. Only embryos that responded to the mechanical stimulus and had a straight body were included in the vision analysis.
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