Title
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Permalink
https://escholarship.org/uc/item/9jr0q3tt

Journal
The Journal of cell biology, 214(3)

ISSN
0021-9525

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Publication Date
2016-08-01

DOI
10.1083/jcb.201604002

Peer reviewed
Assembly and activation of dynein–dynactin by the cargo adaptor protein Hook3

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Metazoan cytoplasmic dynein moves processively along microtubules with the aid of dynactin and an adaptor protein that joins dynein and dynactin into a stable ternary complex. Here, we examined how Hook3, a cargo adaptor involved in Golgi and endosome transport, forms a motile dynein–dynactin complex. We show that the conserved Hook domain interacts directly with the dynein light intermediate chain 1 (LIC1). By solving the crystal structure of the Hook domain and using structure-based mutagenesis, we identify two conserved surface residues that are each critical for LIC1 binding. Hook proteins with mutations in these residues fail to form a stable dynein–dynactin complex, revealing a crucial role for LIC1 in this interaction. We also identify a region of Hook3 specifically required for an allosteric activation of processive motility. Our work reveals the structural details of Hook3’s interaction with dynein and offers insight into how cargo adaptors form processive dynein–dynactin motor complexes.

Introduction

Eukaryotic cells use molecular motors to transport and spatially organize organelles, proteins, and mRNAs within the cytoplasm. Cytoplasmic dynein is a molecular motor that carries cargo toward microtubule minus ends (Allan, 2011). Dynein is a large homodimer composed of two ∼500-kD heavy chains that contain the ATPase motor domain (Schmidt, 2015; Bhabha et al., 2016). The N-terminal portion of the heavy chain binds additional subunits known as the dynein tail subunits, which include the light intermediate chain (LIC), intermediate chain (IC), and light chains (LCs; LC8, Tctex1, and LC7/roadblock; Pfister et al., 2006; Pfister and Lo, 2012). This tail complex is responsible for linking dynein to cargo (Pfister and Lo, 2012). Mammalian dynein is not constitutively active; rather, its motility is regulated by cargo interaction (McKenney et al., 2014; Schlager et al., 2014).

The mammalian LICs, encoded by two closely related gene products, LIC1 and LIC2 (Hughes et al., 1995; Tyan et al., 2000), are involved in several different types of cargo interactions and dynein-based movements, including endosomal and lysosomal transport, ER export, Golgi transport, and axonal vesicle trafficking (Koushika et al., 2004; Palmer et al., 2009; Horgan et al., 2010; Tan et al., 2011; Kong et al., 2013; Brown et al., 2014). The domain structure of the LIC allows it to interact with cargo adaptors while integrated into the dynein holoenzymes. The LIC’s highly conserved N-terminal G protein–like domain binds directly to the dynein heavy chain, and the less conserved C-terminal domain binds adaptor proteins (Schroeder et al., 2014; Fig. 1 A). These cargo adaptors are themselves multifunctional proteins that can bind to a protein (e.g., a Rab GTPase) on a membranous cargo (Fu and Holzbaur, 2014; Cianfrocco et al., 2015; Carter et al., 2016).

In addition to binding dynein LIC and cargo, adaptor proteins promote an interaction between dynein and dynactin, a 12-subunit protein complex (Schroer, 2004). For mammalian dynein, the formation of this tripartite complex is important for long-distance movement (processivity) along microtubules (McKenney et al., 2014; Schlager et al., 2014). This mechanism has been best studied for Bicaudal D2 (BicD2), an adaptor that links dynein–dynactin to Rab6 on Golgi-derived vesicles (Dienstbier and Li, 2009). The N terminus of BicD2 consists of a 270-residue coiled coil that sits in a groove of the Arp1 filament of dynactin and also interacts with the N-terminal region of the dynein heavy chain; this dynein heavy chain–BicD2-Arp1 interaction was proposed to stabilize the tripartite complex (Urnavicius et al., 2015). The mechanism by which this interaction promotes motility is less clear. One possibility is that cargo adaptors activate an autoinhibited state of dynactin (McKenney et al., 2014), enabling it to bind to microtubules and initiating motility. Mammalian dynein also may be locked in an autoinhibited conformation (Torisawa et al., 2014), and cargo adaptors and dynactin may release this autoinhibited state and reposition the motor domains of the dynein dimers for motility (Urnavicius et al., 2015). These models, however, have not proposed a role for a LIC–adaptor protein interaction. Furthermore,
it is unclear whether the assembly of the tripartite motor complex and activation of motility are separable functions.

One cargo adaptor that has been shown to assemble and activate dynein–dynactin is Hook3, although its mechanism has been less studied compared with BicD2. The Hook proteins, first identified for their role in endocytic cargo sorting in Drosophila melanogaster (Krämer and Phistry, 1999), are a widely expressed class of dynein-associated cargo adaptor proteins (Bielska et al., 2014; Zhang et al., 2014). Drosophila and fungi have a single Hook gene, whereas mammals have three Hook genes. The most conserved region of the Hook genes is found at the N-terminal domain (aa 1–160; Fig. 1 A). Without this Hook domain, Hook can no longer interact with dynein and dynactin alone (Hook31–160) also bound LIC1 389–523, albeit more weakly than Hook31–239 (Fig. 1, D and E; and Fig. S1 B). Hook31–160 lacks the predicted coiled coil found in Hook31–239, and thus the stronger interaction of Hook31–239 might be because it is a dimer. We therefore tested an artificially dimerized coiled-coil region of Hook3 protein to test for interactions with LIC1. GFP-tagged Hook31–552 was incubated with beads coated with GST-tagged versions of either full-length LIC1, the LIC N-terminal G-domain (LIC11–389), or the C-terminal domain (LIC1389–523); the beads and any interacting proteins were centrifuged, and the protein composition of the pull-down was analyzed by immunoblot. The results revealed that Hook31–552 cosedimented with full-length LIC1 and the LIC C terminus alone, but not with the N-terminal LIC1 G domain (Fig. 1, B and C; and Fig. S1 A). Thus, similar to the other cargo adaptors RILP, BicD2, and FIP3 (Schroeder et al., 2014), Hook3 also directly binds to LIC1389–523.

We truncated Hook31–552 to identify a smaller fragment that might bind LIC1389–523. The shorter truncation Hook31–239 bound to LIC1389–523 in the pull-down assay (Fig. 1, D and E; and Fig. S1 B), and the two proteins co-eluted as a stable complex by gel filtration chromatography (Fig. S1 C). The Hook domain alone (Hook31–160) also bound LIC1389–523, albeit more weakly than Hook31–239 (Fig. 1, D and E; and Fig. S1 B). Hook31–160 lacks the predicted coiled coil found in Hook31–239, and thus the stronger interaction of Hook31–239 might be because it is a dimer. All mammalian Hook isoforms form a complex with Fused Toes and the Fused Toes– and Hook-interacting protein; fungal homologs of these proteins are important for dynein-mediated early endosome transport by linking Hook to the cargo (Xu et al., 2008; Yao et al., 2014). Here, we sought to understand the mechanism by which Hook3 interacts with dynein and dynactin and activates processive motility. We discuss the crystal structure of the Hook domain and show that this domain binds directly to the C-terminal region of LIC1. Structure-based mutagenesis studies revealed two conserved surface residues that are essential for this interaction. Abrogation of the LIC interaction renders Hook3 unable to join dynein and dynactin in a stable complex. Interestingly, although the N-terminal 239 residues of Hook3 are sufficient for forming a stable complex with dynein–dynactin, this tripartite complex is immobile; activation of motility requires a more distal coiled-coil region of Hook3. This result reveals that complex assembly and activation of motility are separable activities. Our data suggest a model for how Hook3 joins dynein and dynactin into a motile complex.

**Results**

The Hook domain of Hook3 binds to the dynein LIC

Hook3 is comprised of the N-terminal, highly conserved Hook domain (Walenta et al., 2001), followed by three coiled coils and a C-terminal cargo-binding region (Fig. 1 A). A yeast two-hybrid assay revealed an interaction between aa 1–236 of Caenorhabditis elegans Hook and the LIC (Malone et al., 2003). We sought to confirm a direct interaction between Hook3 and LIC1 using purified proteins, as we demonstrated previously for the adaptor proteins RILP, BicD2, and FIP3 (Schroeder et al., 2014). Previous work showed that Hook31–552 is sufficient to produce a highly processive dynein–dynactin–Hook3 complex (McKenney et al., 2014), and thus we used this slightly truncated Hook3 protein to test for interactions with LIC1. GFP-tagged Hook31–552 was incubated with beads coated with GST-tagged versions of either full-length LIC1, the LIC N-terminal G-domain (LIC11–389), or the C-terminal domain (LIC1389–523); the beads and any interacting proteins were centrifuged, and the protein composition of the pull-down was analyzed by immunoblot. The results revealed that Hook31–552 cosedimented with full-length LIC1 and the LIC C terminus alone, but not with the N-terminal LIC1 G domain (Fig. 1, B and C; and Fig. S1 A). Thus, similar to the other cargo adaptors RILP, BicD2, and FIP3 (Schroeder et al., 2014), Hook3 also directly binds to LIC1389–523.

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indicate that the N-terminal Hook domain can bind specifically to the C-terminal region of LIC1 and that the region between aa 160 and 239 strengthens this interaction.

**The Hook domain contains a calponin homology fold with an extended α-helix**

We attempted to co-crystallize LIC1389–523 with either Hook31–239 or Hook31–160, but crystals were obtained only for Hook31–160. A 1.7-Å dataset was obtained from one of the crystals, and a polyalanine model based on a nuclear magnetic resonance (NMR) solution structure of mouse Hook1 1–160 (PDB 1WIX) was used for molecular replacement. After multiple rounds of refinement, the final structure has an Rwork of 18.4 and Rfree of 21.4 (Table 1). Two copies of the protein are present in the asymmetric unit and interact through an antiparallel arrangement of their C-terminal α-helices (helix H described later). This interaction may not be physiological because Hook31–160 is monomeric, as determined by static light scattering (unpublished data).

The Protein Homology/analogY Recognition Engine (PHYRE), which predicts a protein’s tertiary structure based on homology, previously predicted that the Hook domain is comprised of a calponin homology (CH) fold (Zhang et al., 2014). Our structure indeed exhibits a canonical seven-helix CH fold (Fig. 2 A). However, the crystal structure reveals an additional eighth α-helix (helix H, aa 132–158; Fig. 2 A), which was not expected from prior secondary structure prediction (Drozdetskiy et al., 2015). This same α-helix also appears in the NMR structure of the Hook domain of mouse Hook1 (PDB 1WIX), but it is bent in the middle and folded back on itself (Fig. S2 A). Thus, it appears that helix H is able to adopt different conformations; the extended conformation that we have observed may be stabilized by protein–protein interactions in the asymmetric unit.

We next mapped the conserved surface residues in the Hook domain onto our crystal structure using an alignment of 19 Hook domain sequences ranging from fungal to mammalian species (Fig. S3). Strikingly, one side of the structure is much more highly conserved than the other (Fig. 2 B). This contrast is even more evident in the map of conserved residues between the

![Figure 2. The structure of the Hook domain exhibits an extended α-helix and restricted conservation. (A) The 1.7-Å structure of the Hook domain (aa 9–158) from human Hook3 with the helices labeled A–H. Colors (helices A–G) denote the canonical CH domain. (B) The conservation of residues on the surface of the structure in A is shown with red representing the most conserved and white depicting the least conserved. Highly conserved residues are labeled.](image-url)
three human Hook genes (Fig. S2, B and C). Several highly conserved residues lie within the extended helix H, including the universally conserved Q147 and nearby conserved hydrophobic residues. Two other prominent patches of conservation lie on this same face of the CH domain—one cluster consists mainly of hydrophobic residues (S15, W19, and L123), and the other consists of charged residues (K77, D102, E108, and E114).

Two highly conserved residues mediate the Hook3-dynein interaction and are critical for dynein-dynactin motility

The surface conserved residues could be part of a binding interface with the dynein light intermediate chain. To test which region of the Hook domain might be involved in binding LIC1, we made four proteins with different clusters of alanine mutations: (1) Q147A/M151A/I154A, (2) I136A/I139A/M142A, (3) N68A/W69A/K77A, and (4) D102A/E108A (Fig. 3 A). These mutations were made in the construct Hook31–239 because of its higher binding affinity to LIC1 than Hook31–160. The triple and double mutations produced monodisperse protein with a similar gel filtration pattern to the wild-type (WT) protein (Fig. S4 A). We tested each mutant Hook3 protein for binding to GST-LIC1389–523 using the bead pull-down assay (Fig. 3, B and C; and Fig. S4 B). The triple mutants Q147A/M151A/I154A and I136A/I139A/M142A exhibited little or no detectable binding. In contrast, the triple mutant N68A/W69A/K77A and the double mutant D102A/E108A showed little difference in binding (Fig. 3, B and C; and Fig. S4 B). Because patches Q147A/M151A/I154A and I136A/I139A/M142A both lie within helix H, these results suggest that the highly conserved helix H contains the main LIC1 binding interface.

We investigated the more solvent-exposed Q147A/M151A/I154A patch in more depth with single-point mutants. The gel filtration of the Q147A, M151A, and I154A mutants also show monodisperse protein, as shown with the triple mutants (Fig. S4 A). Strikingly, the single I154A and Q147A mutations each led to a dramatic reduction in the Hook3–LIC1 interaction (Fig. 4, A and B; and Fig. S4 C). In contrast, the Hook3 mutant M151A could still bind LIC1 as well as WT (Fig. 4, A and B; and Fig. S4 C).

We tested whether these single-point mutants affected the binding of Hook31–552 to intact dynein and dynactin in a porcine brain lysate (McKenney et al., 2014). WT Hook31–552 pulled down dynein–dynactin as previously reported (McKenney et al., 2014), but no detectable endogenous BicD2 (another cargo adapter that can bind dynein–dynactin; Fig. S4, D and E). In contrast, the Hook3 single-point mutants Q147A and I154A bound very little or no dynein and no detectable dynactin, whereas the M151A mutant bound dynein–dynactin in a manner similar to WT (Fig. 4, C and D; and Fig. S4 E). These results indicate that the highly conserved residues Q147 and I154 in helix H of Hook3 both play critical roles in binding LIC1 and forming a stable dynein–dynactin complex.

We next investigated the ability of Hook3 mutant proteins to stimulate dynein–dynactin motility (McKenney et al., 2014). Dynein and dynactin, purified from a human RPE-1 cell line (Fig. S4 F), were preincubated with GFP-tagged Hook3 constructs. The mixture was then added in the presence of ATP to glass-immobilized microtubules, and interactions of GFP-Hook3 with microtubules were examined by total internal reflection fluorescence (TIRF) microscopy. Processive movement of dynein–dynactin and WT GFP-Hook3 was observed as previously described (McKenney et al., 2014). The point mutant M151A produced a similar number of motile dynein–dynactin–Hook3 molecules compared with WT GFP-Hook3 (Fig. 4, E and F), and the velocities of the molecules were in a similar range as WT Hook3 (Fig. S5 C). In contrast, Q147A and I154A GFP-Hook3 constructs did not elicit processive runs (Fig. 4, E and F), presumably because they did not bind to and form a complex with dynein and dynactin. Thus, Q147 and I154 are each essential for Hook3’s interaction with LIC1 and for the formation of a processive dynein–dynactin complex.

Hook3 truncations that assemble dynein–dynactin do not elicit processive motility

We sought to define the roles that the Hook domain and the extended coiled coil domains of Hook3 play in assembling dynein and dynactin into a complex. Previous work on the 270-residue coiled-coil domain of BicD2 showed that it sits in the groove of the dynactin Arp1 filament and creates a binding interface with the dynein heavy chain (Urnavicius et al., 2015). We made two constructs that consisted primarily of the Hook domain (aa 1–160 and 1–239), truncations that excluded the Hook domain (aa 160–552 and 239–552), and a truncation that excluded just the CH domain but contained helix H of the Hook domain (aa 130–552; Fig. 1 A). These constructs, bound to Strep-Tactin resin, were incubated with porcine brain lysate and then assessed for their ability to pull down the endogenous dynein–dynactin complex by immunoblotting for the dynein IC and the dynactin subunit p150. The construct Hook31–239 pulled...
Two conserved Hook3 residues are critical for the assembly and motility of dynein–dynactin. (A) Single-point mutations Q147A, M151A, and N263A in Strep-Tactin–Hook3Δ239 were compared with WT and tested for binding to human GST-LIC1389–523 as in Fig. 3B (representative of triplicate experiments). Negative control lacks LIC1 on the beads. (B) Ratio of band intensity to the WT Hook31–552 signal in A; mean and SD from n = 3 independent experiments. (C) StrepII-Hook3 constructs, bound to Strep-Tactin resin, were incubated with porcine brain lysate; and the resin analyzed by immunoblotting for the dynein intermediate chain (IC) and the p150 subunit of dynactin. The amount of each Hook3 for the dynein intermediate chain (IC) and the p150 subunit of dynactin. The relative amounts of dynein pulled down by Hook31–552 (Fig. 5, A and B; and Fig. S5 A). However, lengthening the coiled-coil domain from residue 239 to 552 did not significantly change the amount of dynein that was pulled down with Hook3 from the brain lysate (Fig. 5, C and D; and Fig. S5 A). However, lengthening the coiled coil resulted in a progressive increase in the amount of interacting dynactin (Fig. 5, C and D; and Fig. S5 A). These results suggest that it is not essential for the Hook3–dynactin–LIC1 interaction. However, a longer Hook3 coiled coil is able to increase the affinity of dynein–Hook3 for dynactin.

We tested the microtubule binding ability and motility of the dynein–dynactin complex with Hook31–239, Hook31–402, Hook31–440, and Hook31–552. In this experiment, dynein and dynactin were first purified by affinity chromatography (see Methods) and then incubated with these truncated Hook proteins. Surprisingly, in the presence of ATP and dynein–dynactin, all of the truncations induced poor single-molecule motility compared with Hook31–552 (Fig. 5, E and F). Hook31–239 produced no processive motility at all, and even the longer constructs Hook31–402 and Hook31–440 produced very few motile events (Fig. 5, E and F). The few complexes that were motile with Hook31–402 and Hook31–440 exhibited similar velocities to Hook31–552 (Fig. S5 C). We tested whether the smallest truncation Hook31–239 might be unstable after addition of ATP, but found only a slight (18%) dissociation of dynein from the tripartite complex on beads after a 1-h incubation with 2.5 mM ATP (Fig. S5, D–F). In the microscopy assay in the absence of ATP, all Hook3 truncations did not bind microtubules alone (Fig. S5 G), but bound statically to microtubules in the presence of dynein–dynactin (Fig. 5, G and H). The results suggest that dynein–dynactin complexed with short Hook3 constructs can bind to microtubules in the absence of ATP (rigor microtubule binding with a low dissociation rate) but do not engage in a productive motility cycle in the presence of ATP (see Discussion). These findings indicate that the region of the coiled coil between aa 402 and 552 of Hook3 is required for robust activation of motility of the dynein–dynactin–Hook3 complex.

Discussion

In this study, we delineated the minimal binding regions for Hook3 that are required for two activities: (1) binding to the LIC1 C-terminal domain (Hook31–160) and (2) producing a dynein–dynactin complex that engages in robust processive motility (Hook31–552). Together, these results suggest a model for how cargo adaptors might regulate the minus end–directed motility of dynein–dynactin.

Our work provides structural insights into how Hook3 binds to dynein. We previously found that the C-terminal half of LIC1 is the docking site for several cargo adaptors (Schroeder...
**Figure 5.** Hook3 truncations that bind dynein–dynactin are not sufficient for motility. (A) Truncations of strepII-Hook3 were tested for binding to endogenous dynein–dynactin in porcine brain lysate as in Fig. 4 C. (B) The ratio of band intensity to the WT Hook31–552 signal in A; mean and SD from n = 3 independent experiments. The truncations not shown were measured to be the same as or less than the signal of the negative control. (C) C-terminal strepII-Hook3 truncations were tested for binding porcine brain dynein–dynactin as in A. The intermediate chain (IC) band in the lane for Hook31–239 is skewed because the IC and this Hook truncation run at the same molecular weight. (D) The ratio of band intensity to the WT Hook31–552 signal in C; mean and SD from n = 3 independent experiments. The following p-values are given for the truncations that differ statistically from Hook31–552: dynein signal-Hook31–160, P < 0.001; Hook31–160–239, P < 0.001; Hook31–348, P < 0.001; Hook31–402, P < 0.001; Hook31–440, P < 0.001; IC signal-Hook31–239, P > 0.03; and *Hook31–402, P < 0.05. (E) Hook3 truncations were tested for their ability to activate motility of the dynein–dynactin complex, indicating that the Hook3–LIC1 interaction is also required. Supporting this conclusion, single-point mutations in helix H might affect a conformational change that could regulate its interaction with the dynein LIC. Mutations in helix H might affect this conformational equilibrium. In addition, Hook31–239 may be able to bind LIC1 better than Hook31–160 because this longer construct might shift a conformational equilibrium of helix H toward its extended form. To test these ideas, further work will be needed to measure the conformational state of this helix.

Although the minimal Hook domain aa 1–160 binds the dynein LIC, it does not appear to be sufficient to recruit the dynactin complex. The first coiled coil of Hook3 (aa 160–239) enables dynactin binding, and the additional coiled coil sequence further enhances this interaction. A cryo-EM study revealed that the 270-residue coiled coil of another cargo adaptor, BicD2, interacts along the groove of the Arp1 filament of dynactin and also mediates an interaction with the dynein heavy chain (Urnauvicus et al., 2015). Similar to BicD2, Hook3’s coiled coils may sit in the groove of the Arp1 dynactin filament and promote an interaction between dynein and dynactin. Hook31–440, for example, may have ~270 residues of coiled coil. However, the coiled coil of Hook3 (aa 160–552) alone is insufficient for stabilizing the tripartite complex, indicating that the Hook3–LIC1 interaction is also required. Supporting this conclusion, single-point mutations in Hook3 (either Q147A or I154A) that abrogate LIC1 binding also completely block the ability of Hook3 to form a helix–helix interaction (Zhang et al., 2014), and we show here that Hook3 binds to this same region of the LIC. Helix H of Hook3, which extends from the CH domain, plays a key role in the LIC interaction, and our structure–function studies reveal two patches of residues in helix H (I136/I139/M142 and Q147/I154) that are involved in the interaction. These residues are highly conserved among all Hook isoforms, and thus it is likely that all Hook gene products bind LIC with a similar mechanism. Interestingly, residue I154 in human Hook3, which we find plays a key role in the LIC interaction, corresponds to L150 in Aspergillus nidulans, which is part of the A. nidulans double mutant (L150P/E151K) shown to disrupt early endosome transport and the HookA–dynein–dynactin interaction in vivo (Zhang et al., 2014). Our work corroborates this in vivo finding and provides structural insight into I154’s conserved role in binding dynein.
dynein–dynactin complex. Thus, multiple protein–protein interfaces of the adaptor Hook3 with the dynein heavy chain, LIC1, and dynactin appear to be required to form a stable tripartite motor complex.

We also show that the C-terminal region of our Hook3 construct is required for robust activation of dynein motility. Several possible models could explain how this additional coiled coil–containing region converts an inactive dynein–dynactin–adapter complex (e.g., one formed by Hook31–239) into an active processive motor (one formed by Hook31–552; Fig. 6). First, a certain length of Hook3 bound along the dynactin Arp1 filament may be required to induce an allosteric conformational change in the dynein heavy chains to release them from an inhibited state (Chowdhury et al., 2015; Urnavicius et al., 2015). For example, an autoinhibited state of dynein may exist in which the two motor domains are stacked, necessitating the separation and alignment in the same direction to become active (Torisawa et al., 2014). An alternative and not mutually exclusive model involves the allosteric regulation by Hook3 of the N-terminal CAP-Gly domain of dynactin’s p150 subunit. The p150 subunit regulates dynein motility (Kardon et al., 2009; McKenney et al., 2014; Tripathy et al., 2014), and p150’s CAP-Gly domain binds to the C terminus of tubulin, an interaction that greatly enhances an initial microtubule binding encounter of dynein–dynactin–BicD2 that leads to processive movements (McKenney et al., 2016). However, dynactin alone exhibits minimal binding to microtubules, suggesting that it is in an autoinhibited state (Kardon et al., 2009; McKenney et al., 2014). This finding agrees with a dynactin cryo-EM structure showing that the junction between CC1A and CC1B in p150 is positioned near the pointed end of the Arp1 filament; in this folded conformation, the CAP-Gly and CC1A domains are unlikely to be accessible to the microtubule (Urnavicius et al., 2015). In a lower-resolution structure of the dynein–dynactin–BicD2 complex, the C terminus of a 270-aa coiled coil of BicD2 is located at the pointed end of the dynactin Arp1 filament (Urnavicius et al., 2015). We speculate that the C-terminal end of our motility-inducing Hook3 construct (aa 400–552) may somehow act to dislodge CC1A-CC1B from the backbone of the Arp1 filament. The release of p150 may enable this subunit to extend fully into an active conformation, enabling access to the microtubule (Fig. 6).

Although our data reveal an important role of the LIC in Hook3-mediated dynein motility, several questions remain unanswered. First, it is unknown whether the LIC acts as a passive tether for linking the motor domain to cargo adaptors or whether it also undergoes a conformational change that plays an active role in eliciting dynein motility. Second, it remains to be determined whether other cargo adaptors that interact with LIC1 (e.g., FIP3, RILP, and BicD2) do so through mechanisms similar to or different from those of Hook proteins. Among these adaptors, the Hook domain appears to be unique. Third, we also do not know how many cargo adaptors interact with dynein chains other than the LIC. For example, BicD2 has been shown to interact with both the dynein intermediate chain (Hoogenraad et al., 2001, 2003) and the light intermediate chain (Schroeder et al., 2014). Differences in activation among the adaptors might allow for many ways of regulating dynein-based cargo transport. Many of these important questions can be addressed through structural and functional studies of multiple types of cargo adaptor proteins interacting with dynein and dynactin.

Materials and methods
Molecular biology
The cDNA of Hook3 was obtained from a human cDNA library made from mitotic RPE-1 cells, and all Hook3 constructs were cloned into vector pET28a with an N-terminal 6xHis-strepII-Superfolder GFP (stGFP) tag. All human LIC1 (RefSeq accession number NM_016141.3) constructs were cloned into pGEX6P1, which included an N-terminal GST tag and a C-terminal strepII tag. Truncations were based on both secondary structure prediction (Drozdetskiy et al., 2015) and coiled coils prediction (Lupas et al., 1991). To dimerize the Hook31–400 construct, a Gln4 sequence (Harbury et al., 1993) was added to the C terminus. The 29-aa sequence was VKQLEDKVEELLSDKNAHLEVARLKLV. Full-length human FIP3 (GenBank accession number AB383948) was cloned into pET28a with a strepII-SNAP tag and was used for the purification of the dynein–dynactin complex from porcine brain lysate.

Protein purification
All human Hook3 constructs were transformed into the Escherichia coli strain BL21 RIPL, and expression was induced with 0.5 mM IPTG at 37 °C for 3–6 h. Bacterial pellets were resuspended with lysis buffer (25 mM Hepes, pH 7.8, 150 mM NaCl, 2 mM TCEP, 1 mM PMSF, and a protease inhibitor cocktail [Roche], 1 tablet per 50 ml) and lysed using an Emulsiflex press (Avestin). The lysate was clarified by centrifugation at 40,000 g for 30 min, and Hook3 was purified using Strep-Tactin
Superflow Plus resin (QIAGEN). The agaroose was then washed with lysis buffer (excluding the Roche protease inhibitor cocktail) at ~20x
the resin volume, and the purified protein was eluted with 3 mM des-
thiobiotin. The protein was concentrated and flash frozen. Thawed
protein was then further purified by gel filtration with a Superose 6
10/300 GL or a Superdex 200 10/300 GL column (GE Healthcare).
The gel filtration buffer was 30 mM Hepes, pH 7.4, 150 mM NaCl,
2 mM MgCl₂, 5% glycerol, and 2 mM TCEP. The Hook3-containing
fractions were pooled, concentrated, and flash frozen. Strepll-SNP-
FIP3 was purified the same way as Hook3, except the lysis buffer in-
cluded 25 mM Tris, pH 8.5.

GST-LIC1-strepll constructs (full length and truncations) were
expressed as performed with the Hook3 constructs followed by lysis
with 50 mM Tris, pH 7.4, 150 mM NaCl, 5 mM TCEP, 1 mM PMSE,
and a protease inhibitor cocktail (Roche; 1 tablet per 50 ml).The pro-
tein was then purified using either glutathione agarose 4B (USB) or
Strep-Tactin Superflow Plus resin (QIAGEN). After extensive washing
and elution with either 10 mM reduced glutathione at pH 7.4 (for gluta-
Thione agarose) or 3 mM desthiobiotin (for Strep-Tactin resin), the pro-
tein was gel filtered using a HiPrep 16/60 Superdex S-200 HR column
(GE Healthcare) in 20 mM Tris, pH 7.4, 50 mM NaCl, and 2 mM TCEP.

Pull-downs
Clarified porcine brain lysate was used to test the binding of endoge-
 nous dynein–dynactin to Hook3 constructs and was prepared as pre-
viously described (McKenney et al., 2014). For each dynein–dynactin
pull-down, 500 µl porcine brain lysate in buffer A (30 mM Hepes, pH
7.4, 50 mM potassium acetate, 2 mM magnesium acetate, 1 mM EGTA,
and 10% glycerol) was combined with 60 µl of 50% Strep-Tactin
Sepharose slurry (GE Healthcare), 0.1% NP-40, 5 mM DTT, and
1 mM PMSE. SGFP-tagged Hook3 constructs were added at 200–400
µm to the brain lysate and resin and incubated for 1–2 h at 4°C. The
resin was pelleted and washed five times in 500 µl buffer A includ-
ing 0.1% NP-40 and 5 mM DTT. After the final wash, the resin was
resuspended in 50 µl loading buffer, and an equal volume of the sam-
ples was resolved on NuPAGE gels (Invitrogen). All dynein–dynactin
pull-downs were repeated at least three times on separate days starting
from frozen brain lysate.

To test the binding of human LIC1 to Hook3 constructs, 200 mM
GST-LIC1-strepll (full length or truncations) was incubated with 20 µl
glutathione resin in a 300-µl volume of buffer (50 mM Tris, pH 7.4,
100 mM NaCl, 5 mM TCEP, 0.1% Tween, and 2 mg/ml BSA). After
extensive washing of the resin, 300 µl of 200 nM SGFP-Hook3 con-
struct was added and incubated for 1 h at 4°C. The resin was washed
tensively and resuspended in 20 µl of 1x loading buffer. Samples
were resolved on NuPAGE gels. All pull-downs with purified LIC1 and
Hook3 proteins were repeated at least three times on separate days.

Western blot analysis
After samples were resolved by SDS-PAGE, they were transferred to
nitrocellulose membranes with the iBlot Gel Transfer Device (Invitro-
gen). Membranes were blocked with 5% milk in TBS and 0.1% Tween
(TBST) and probed at RT with primary antibody, which included rabbit
anti-GFP (1:1,000; Abcam), mouse anti-GST (1:1,000; Thermo Fisher
Scientific), mouse anti-dynein intermediate chain (clone 74.1, 1:1,000;
EMD Millipore), mouse anti-p150 (1:250; BD), and mouse anti-BicD2
(1:200; sc-393631; Santa Cruz Biotechnology, Inc.). Membranes were
then washed three times with TBST and incubated with anti-mouse–
800 or anti-rabbit–680 (1:10,000; Molecular Probes) for 45 min to 1 h
at RT. Blots were visualized with an Odyssey Clx Infrared Imaging
System (LI-COR Biosciences). Western blots were quantified using
ImageJ (National Institutes of Health). A box was drawn around each
band, and an equivalently sized box was drawn in the lane correpond-
ing to the negative control. The intensity of each box was plotted, and
the area under the subsequent curve was measured. The intensity of the
negative control was subtracted from the corresponding sample. Band
intensities of the prey in pull-downs were then normalized by the band
intensities of the bait used in the assay (sfGFP-Hook3 or GST-LIC1).
The prey’s normalized intensities (arbitrary units) or the ratio of normal-
ized intensities are presented. SDs were calculated for the intensities or
ratio of intensities from three independent experiments and displayed as
error bars. P values were calculated using an unpaired t test.

Cryocrystallization and structure determination
The LIC1 C-terminal half (LIC389-523) and GST-Hook31-160 were puri-
ified with glutathione agarose resin 4B (USB), cleaved from the resin
using purified GST-tagged human rhinovirus 3C protease, and incu-
batated overnight at 4°C. After the GST tag was cleaved, the two proteins
were combined at an equimolar ratio and incubated on ice for 30 min.
The proteins were gel filtered using a HiPrep 16/60 Superdex S-200
HR column (GE Healthcare) into the following buffer: 10 mM Tris, pH
7.4, 25 mM NaCl, and 2 mM TCEP. Fractions containing both proteins
were concentrated to ~20 mg/ml, and hanging drop vapor diffusion
experiments were set up using 96-well crystal screens (QIAGEN) at
RT. Native crystals grew from a reservoir solution containing 2 M so-
dium formate and 0.1 M sodium acetate, pH 4.6 (JCSG screen Core
III; QIAGEN). The crystals were cryoprotected with the addition of
35% glycerol to the crystallizing well solution and were flash cooled
by plunging in liquid nitrogen.

Native diffraction data were collected at beamline 8.3.1 at the
Advanced Light Source (Lawrence Berkeley National Laboratory), and
the dataset was indexed and integrated in P 2 1 2 1 using XDS (Kabsch,
2010). The structure was solved by molecular replacement using an
ensemble of 20 superimposed NMR models from PDB structure 1WXI
using Phaser (McCoy et al., 2007). The Phaser scores for the best solu-
tion were modest (RFZ = 4.8 and TFZ = 6.4), and the initial electron
density maps were noisy and discontinuous. Density modification and
chain tracing with SHELXE (Sheldrick, 2010) resulted in an easily in-
terpretabl map and a poly-alanine model that was further improved
using phenix.autobuild (Terwilliger et al., 2008). Multiple rounds of
model building and refinement were done using Coot (Emsley and
Cowtan, 2004) and phenix.refine (Adams et al., 2010). The data col-
clection and refinement states are presented in Table 1, and the PDB
accession number is 5J8E.

Purification of dynein–dynactin from human RPE-1 cells
RPE-1 cell lysate was prepared as previously described (McKenney
et al., 2014). The lysate was centrifuged at 266,000 g for 10 min at 4°C,
and final concentrations of 5 mM DTT, 0.1% NP-40, and 1 mM PMSE
were added before use. The lysate was incubated with purified strepll-
SNAP-FIP3 on Strep-Tactin Sepharose (GE Healthcare). After incuba-
tion at 4°C for 1–2 h, the resin was thoroughly washed with buffer A
(30 mM Hepes, pH 7.4, 50 mM potassium acetate, 2 mM magnesium ace-
tate, 1 mM EGTA, 10% glycerol, 5 mM DTT, 0.1% NP-40, and
1 mM PMSE) and resuspended in buffer A with 300 mM NaCl to release
dynein–dynactin from resin-bound FIP3. After incubating on ice for
10 min, the high-salt slurry was centrifuged through a 0.2-µm filter
to remove the resin. Then an equal volume of 50% Strep-Tactin Sepharose
slurry was added to the elution to bind any strepll-FIP3 that may have
released from the resin during the high-salt incubation. After incubating
on ice for 10 min, the slurry was once again filtered, and the final solu-
tion was diluted with buffer A to a final concentration of 200 mM NaCl.
Sucrose was also added at a final 6% concentration, and the affinity-
purified dynein–dynactin was flash frozen for single molecule imaging.
Single-molecule imaging
Preparation of microtubules. Tubulin was purified from porcine brain and labeled (fluorescently or with biotin) as previously described (Castoldi and Popov, 2003). To polymerize microtubules, unlabeled tubulin was combined with biotin-labeled tubulin and fluorescent tubulin (640 nm fluorescence) at a ratio of ∼10:2:1, respectively, in BRB80 (80 mM Pipes, 1 mM EGTA, and 1 mM MgCl₂) and 5 mM GTP. After incubating for 10 min at 37°C, taxol was added at a final concentration of 20 μM. To remove unpolymerized tubulin, the microtubules were layered over a 25% sucrose cushion and centrifuged at 65,000 g for 5 min at 22°C.

Preparation of dynein-dynactin-Hook3 complexes. A 30-μl reaction consisting of 10 nM sfGFP-tagged Hook3 and 5 μl of ∼0.15 mg/ml native dynein–dynactin purified from RPE-1 cells was incubated in 30 mM Hepes, pH 7.4, 50 mM potassium acetate, 2 mM magnesium acetate, 1 mM EGTA, 10% glycerol, 0.1 mg/ml BSA, 0.5% pluronic acid F-127, 0.2 mg/ml κ-casein, and a Trolox/PCA/PCD scavenging system (Dave et al., 2009).

TIRF microscopy. Flow chambers (volume ∼10 μl) were constructed using double-sided tape and acid-washed coverslips as described (Tienbaum et al., 2013). The chambers were prepared with immobilized fluorescent microtubules by coating the chamber in the following sequence of solutions: 10 μl of 5 mg/ml BSA-biotin (Thermo Fisher Scientific), 20 μl BC buffer (BRB80, 1 mg/ml BSA, 1 mg/ml casein, and 0.5% pluronic acid F-68, pH 6.8), 10 μl of 0.5 mg/ml streptavidin (Vector Laboratories), 20 μl BC buffer, and finally 10 μl of a 1:10 dilution of microtubules (prepared as described earlier). Microtubules were washed with the assay buffer, and a 1:10 dilution of the dynein–dynactin–Hook3 complex described earlier was added to the flow chamber in the presence of 1 mM ATP. Movies were acquired with an Eclipse TE200-E microscope (Nikon) equipped with an iXon EMCCD camera (Andor), a 100x 1.49-NA objective, and Micromanager software (Edelstein et al., 2010). A 491-nm laser (at 75% laser power) and a 640-nm laser (at half maximum laser power) were used to image sfGFP-Hook3 (100 ms exposure) and fluorescently labeled microtubules (50 ms exposure), respectively. Several 6-min movies (1- or 2-s intervals of image acquisition) were acquired at RT per flow chamber per construct. Molecules that moved >1 μm were scored as processive.Velocities were quantified by making kymographs in ImageJ (National Institutes of Health).

Online supplemental material
Fig. S1 shows the complete gels and relative inputs for pull-downs of Fig. 1, and it shows the co-gel filtration of LIC1389–523 and Hook31–239. Fig. S2 shows the alignment of human Hook3 and mouse Hook1 (PDB 394. http://dx.doi.org/10.1093/nar/gkv332

References


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