Dynactin integrity depends upon direct binding of dynamitin to Arp1

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Abstract Dynactin is a multiprotein complex that works with cytoplasmic dynein and other motors to support a wide range of cell functions. It serves as an adaptor that binds both dynein and cargoes and enhances single-motor processivity. The dynactin subunit dynamitin (also known as p50) is believed to be integral to dynactin structure because free dynamitin displaces the dynein-binding p150Glued subunit from the cargo-binding Arp1 filament. We show here that the intrinsically disordered dynamitin N-terminus binds to Arp1 directly. When expressed in cells, dynamitin amino acids (AA) 1–87 causes complete release of endogenous dynamitin, p150, and p24 from dynactin, leaving behind Arp1 filaments carrying the remaining dynactin subunits (CapZ, p62, Arp11, p27, and p25). Tandem-affinity purification–tagged dynamitin AA 1–87 binds the Arp filament specifically, and binding studies with purified native Arp1 reveal that this fragment binds Arp1 directly. Neither CapZ nor the p27/p25 dimer contributes to interactions between dynamitin and the Arp filament. This work demonstrates for the first time that Arp1 can directly bind any protein besides another Arp and provides important new insight into the underpinnings of dynactin structure.

Introduction First identified as an activity required for dynein to move membrane vesicles on microtubules in vitro (Schroer and Sheetz, 1991), dynactin has emerged as an essential component of the cytoplasmic dynein motor complex. Dynactin in most species contains 11 different polypeptide components in stoichiometries ranging from 1 to ≥5. Dynactin’s largest structural domain is a 37 × 5 nm copolymer of Arp1, actin, and Arp11 capped with other subunits (Schroer, 2004). Its conspicuous 24-nm-long projecting arm (p150Glued amino acids [AA] 1 to –350), which can bind microtubules at the distal tip, extends from a V-shaped “shoulder” structure that is docked at one end of the Arp filament (Imai, Narita, Maeda, and Schroer, unpublished data). The shoulder contains the remainder of p150Glued (AA = 350–1280), including the dynein-binding site (Sigglin et al., 2013), plus dynamitin and p24 (reviewed in Schroer, 2004; Cheong and Schroer, 2011). For dynein to be able to transport cargo, p150Glued must be anchored to the cargo-binding Arp filament domain, but the structural features that underlie this critical interface remain undefined. It has been proposed that p150Glued binds Arp1 directly (Waterman-Storer et al., 1995), and in yeast, the dynamitin homologue (Jnm1p) was reported to interact with Arp1 in a two-hybrid analysis (Clark and Rose, 2005), suggesting that multiple protein–protein interactions may be involved.

The importance of dynamitin to dynactin’s structural integrity was first revealed through protein overexpression experiments, and, as a result, dynamitin overexpression has become a widely used tool for interfering with dynein-based events in vivo. The name “dynamitin” was coined based on the “explosion” of mitotic spindle poles, Golgi complex, and dyactin molecules seen in overexpressing cells (Echeverri et al., 1995; Burkhardt et al., 1997). Dynamitin also integrates dynactin structure when the purified proteins are mixed in vitro (Eckley et al., 1999; Wittman and Hyman, 1999; Melkonian et al., 2007). Dynamitin’s remarkable ability to separate the dynactin molecule into its cargo- and dynein-binding components suggests that its associations with other dynactin subunits is of fundamental importance.
importance to dynactin structure. Purified dynamitin can form complexes with itself and the shoulder components p24 (Maier et al., 2008), but it has never been shown to bind directly to any other dynactin subunit in a biochemical assay. An intriguing and obvious possibility is that dynamitin also binds directly to Arp1 to anchor the shoulder/arm complex to the Arp filament. In support of this notion, an N-terminal fragment of dynamitin tagged with glutathione S-transferase was seen to pull down the p62 and Arp1 components of dynactin from cultured cell lysates (Jacquot et al., 2010). Whether dynamitin was binding Arp1 directly or whether another dynactin subunit or extrinsic protein was required was not determined, leaving open the structural details of dynamitin–shoulder anchoring.

We previously proposed that dynamitin’s ability to disassemble dynactin depends upon binding of exogenous protomers to endogenous protomers via an interaction involving a series of self-association motifs (Melkonian et al., 2007; Maier et al., 2008). However, fragments lacking some or all of these motifs are sufficient for disruption (Maier et al., 2008; Jacquot et al., 2010), indicating that dynamitin may be able to trigger dynactin disassembly via more than one mechanism. In the present study, we used direct protein–protein binding assays to characterize interactions between dynamitin’s N-portion and the self-associating C-terminal portion with other dynactin subunits. Tandem-affinity purification (TAP) revealed that dynamitin AA 1–87 binds directly to the Arp1 filament but not components of the shoulder and projecting arm. Biochemical and RNA interference (RNAi)–based assays were used to demonstrate that the dynamitin N-terminus binds directly and specifically to Arp1. A complementary dynamitin fragment (AA 100–403) that does not disrupt dynactin was found to bind p150Glued and p24 but not Arp filament components. Our findings indicate that dynamitin is a divalent scaffold that tethers dynactin’s dynein and microtubule-binding subunit to its cargo-binding domain.

### RESULTS

**In silico and structural analysis of dynamitin AA 1–87**

The dynamitin N-terminus is well conserved from human to *Caenorhabditis elegans* (Figure 1A). It is enriched in charged amino acids and has an overall acidic pI (4.2–4.4) but also contains a number of basic residues. It is highly susceptible to phosphorylation in vitro (Cheong, 2010) and can be phosphorylated in vivo at multiple sites (Figure 1A; PhosphoSitePlus, www.phosphosite.org), suggesting that it may be a regulatory site. The remainder of dynamitin (AA 100 to end) is predicted to fold into a series of α-helices with coiled-coil and multicoil propensity that support oligomerization (Maier et al., 2008), in which one of the hydrophobic heptad repeat residues is replaced with proline (Maier et al., 2008), exhibiting a spectrum similar to wild type (wt) AA 1–210, with a slightly lower α-helix content (11.7%), consistent with the expected effect of the mutant. By contrast, the spectra obtained for dynamitin AA 1–78 and AA 1–87 were consistent with minimal secondary structure, with 2.8 and 1.8% α-helix content, respectively. This analysis verifies that the dynamitin N-terminus is intrinsically disordered. It further suggests that the reason dynamitin AA 1–78 does not disrupt dynactin structure in vivo (Valetti et al., 1999) or in vitro (Maier et al., 2008), it is that it adopts an aberrant structure that is not exhibited by the longer 1–87 fragment. To learn more about the activities of dynamitin’s N- and C-terminal domains, we used AA 1–87 and a complementary fragment, AA 100–403 (Figure 1B), in overexpression and biochemical studies.

**Dynamitin (AA 1–87) is sufficient to cause dynactin disruption**

Overexpression of chicken dynamitin AA 1–87 in Cos-7 cells had effects on cell architecture that were indistinguishable from those of full length dynamitin (Figure 2, A–C, and Supplemental Figure S1). We observed scattered Golgi complexes (Figure 2A) and endocytic recycling compartment/trans-Golgi network components (stained for TGN46; Supplemental Figure S1). Cells overexpressing dynamitin AA 1–87 also contained disorganized, unfocused mitotic spindles (Figure 2, B and C, and Supplemental Figure S1) and
 showed an increased mitotic index (control, 4.6 ± 0.3%; full-length dynamitin, 10.3 ± 0.6%; N-terminus, 9.9 ± 0.2%; n = 1000 cells per condition per experiment in three independent experiments), with most cells arrested in pseudoprometaphase. Because another N-terminal fragment, AA 1–78, could interfere with dynactin function in vivo without triggering p150Glued release (Valetti et al., 1999), we did not expect AA 1–87 to affect dynactin structure. To our surprise, sucrase gradient sedimentation revealed that AA 1–87 did, in fact, cause p150Glued and endogenous dynamitin to be released from the Arp1 filament (Figure 2, D and E). A comparable fragment of human dynamitin (AA 1–90) has a similar effect on dynactin integrity when overexpressed (Jacquot et al., 2010). We conclude that chicken dynamitin AA 1–87, which lacks the oligomerization motifs we showed previously to correlate with dynamitin–dynamitin binding and subunit release, can indeed bind and destabilize dynamitin. This suggests a different mechanism of dynamitin-mediated disruption.

The dynamitin N-terminus binds the Arp filament but does not interact with shoulder-arm subunits

To better understand the mechanism by which the dynamitin N-terminus triggers dynactin disassembly, we used TAP to identify binding partners. Cos-7 cells were transfected with TAP-tagged AA 1–87 or a control protein and allowed to express the proteins for 48 h, and then cytosols were prepared and subjected to affinity purification on streptavidin and calmodulin beads (see Materials and Methods). Components of the Arp1 filament (p62, Arp1/actin, Arp11, CapZ α and β, p27, and p25) were the predominant species on silver-stained SDS gels of the TAP pull down (Figure 3A). The identity of these proteins was confirmed by immunoblotting (Figure 3C, lane 3) and mass spectrometry (Supplemental Table S1). The TAP-tagged dynamitin N-terminus showed no evidence of interaction with p150Glued, dynamitin, or p24, verifying that it binds the Arp filament directly and not through the shoulder/arm complex. Similar results were obtained in an experiment in which we immobilized purified hexahistidine (6X-His)-tagged dynamitin AA 1–87 or a control protein on beads and then added purified bovine dynactin. Once again, the dynamitin N-terminal fragment could pull down Arp1 and its end-binding proteins (CapZ, Arp11, p62, p27, and p25) but not other dynactin subunits (Figure 3B).

The results were obtained using myc-tagged proteins, but similar results were obtained using mCherry-tagged dynamitin species. Myc-tagged dynactin p62 AA 370–467 and monomeric red fluorescent protein (mRFP) were used as controls in A, and CMV-β gal was used as a control in B. (C) Left, representative image of a cell expressing myc-tagged AA 1–87, stained for tubulin. The nonexpressing cell at the upper right is in a different focal plane, and the inset shows this cell’s spindle in focus. (See Supplemental Figure S1 for a merged image showing myc staining.) Right, A control cell expressing CMV-β gal, stained for tubulin. Scale bar, 5 μm. (D) cDNA encoding TAP-tagged AA 1–87 (or buffer as a control) was electroporated into Cos-7 cells. After 48 h, detergent lysates were subjected to velocity sedimentation into a 5–20% sucrose gradient. Gradient fractions were analyzed by immunoblotting to detect the dynamitin subunit p150Glued, Arp1, or dynamitin (DM). Dynamitin AA 1–87 was detected using an antibody to TAP. Similar results were obtained with myc- or mCherry-tagged AA 1–87. (E) Purified bovine dynactin (10 μg) was mixed with 100× molar excess of recombinant (6X-His) dynamitin AA 1–87 and subjected to velocity sedimentation as in D. AA 1–87 was detected using an antibody to the Xpress tag.
Dynamitin AA 1–87 binds the shoulder/arm complex but does not cause dynactin disruption

In our previous structure–function analysis of dynamitin, we focused on three α-helical coiled-coil motifs in AA 100–403 that are necessary for dynamitin self-association, p24 binding, and stable association with the Arp filament (Maier et al., 2008) and proposed that one or more was necessary for disruption. A later report indicated that AA 100–403, a fragment that contains all these motifs, did not cause organelle or spindle derangement when overexpressed (Jacquot et al., 2010). We, too, found that overexpression of TAP-tagged dynamitin AA 100–403 did not trigger release of p150Glued, indicating that it was active for binding, but it also pulled down a minor amount of Arp1, making it difficult to rigorously exclude the possibility of Arp-filament binding. We realized that a trace amount of overexpressed TAP AA 100–403 might have been incorporated into the dynamitin shoulder/arm complex during biosynthesis, which would lead to a small amount of Arp1 being present in the TAP pull down. To rigorously demonstrate that binding of p150Glued to AA 100–403 was independent of Arp1, we added purified AA 1–87 to the cytosol to release the shoulder/arm complex from the Arp filament. When the resulting sample was incubated with immobilized TAP-AA 100–403, p150Glued was once again pulled down, but Arp1 was not (Figure 3C, lane 5, AA 100–403*).

Dynamitin AA 1–87 binds directly to Arp1 but not to other dynactin components

These affinity purification experiments clearly showed that dynamitin AA 1–87 binds dynactin’s cargo-binding domain but did not identify which subunit (i.e., Arp1, CapZ, p27, p25, Arp11, or p62) was involved. Each dynactin molecule contains four dynamin protomers (Eckley et al., 1999), two of which remain tightly associated with the Arp filament after dynamitin-mediated disruption (Melkonian et al., 2007). Arp1 is the only component of the filament that is present in two or more copies, making it an obvious candidate for a binding partner. Unfortunately, binding studies with purified Arp1 are extraordinarily difficult. Arp1 is completely insoluble when expressed in bacteria or insect cells, and so a recombinant form is not available, and cultured animal cells do not contain any free Arp1 protomers even when the protein is overexpressed. To get around these problems, we used native Arp1 isolated from purified bovine dynactin (as in Bingham and Schroer, 1999; see Materials and Methods) for binding assays. Purified Arp1 was found to bind purified 6X-His-AA 1–87 that had been immobilized on beads, but it did not bind a 6X-His-tagged control protein, indicating that binding was specific (Figure 4A). Conventional G-actin processed in parallel showed no evidence of binding to AA 1–87 (Figure 4B). Purified 6X-His-AA 1–87 did not copellet with polymerized F-actin (Figure 4C), further verifying the specificity of its interaction with Arp1. Together these findings demonstrate that AA 1–87 interacts directly and specifically with Arp1.

Dynactin’s Arp filament is “capped” at both ends by protein complexes that prevent subunit addition and loss. The near end of the shoulder-arm complex is located at the conventional actin-capping protein, CapZ/CP (Schafer et al., 1994), whereas the opposite end terminates in the C-terminal fragments of Arp11, p62, p27, and p25 (Eckley et al., 1999). Although dynamitin AA 1–87 binds Arp1 directly, these proteins might participate in binding as well. To determine whether other Arp-filament components contributed to interactions with dynamitin AA 1–87, we embarked on a series of binding studies using cell extracts that had been depleted of dynactin components via RNAi.

The actin capping protein, CapZ, is located at the “barbed” end of the Arp filament in the immediate vicinity of the dynamitin shoulder (Schafer et al., 1994; Imai, Narita, Maeda, and Schroer, unpublished data), making it an obvious potential binding partner for dynamitin. Previous work indicated that CapZ can be depleted from cells using RNAi (Meijllano et al., 2004), and we, too, found that cells transfected with CapZ small interfering RNAs (siRNAs) for 48 h
largely lacked CapZ. However, when we analyzed the depleted cell lysates more carefully using sucrose gradient sedimentation, we detected a small pool of CapZ that sedimented at 20S (Supplemental Figure S3). This represents CapZ that is incorporated into dynactin and thus apparently resistant to depletion owing to dynactin’s long half-life (Brown et al., 2005). Although Arp1 cannot be depleted from the dynactin-associated pool prevented us from using RNAi to eliminate CapZ as a candidate. Instead, we examined binding of free cytosolic CapZ to 6X-His-AA 1–87. Free CapZ dimers were separated from the dynactin-associated, 20S pool using sucrose gradient sedimentation, and then the CapZ-containing fractions were incubated with 6X-His-AA 1–87 adsorbed to beads. CapZ showed no evidence of interaction with AA 1–87 (Figure 5B), but it did bind a known binding partner (mCAH3; Fujiwara et al., 2010; Supplemental Figure S4). We conclude that dynamitin AA 1–87 does not bind CapZ.

We showed previously that the dynamitin components p27 and p25 can be selectively and coordinately depleted from cells using RNAi (Yeh et al., 2012; Figure 5A). p150

\( ^{\text{glx}} \) dynamitin and Arp1 still cosegregated as a 20S particle that contains all the remaining dynamitin subunits. Similarly, Arp1 can still be pulled down with p150

\( ^{\text{glx}} \) when p25 is deleted from Aspergillus (Zhang et al., 2011). These data are strong evidence that the interaction of dynamitin with Arp1 binding does not require p27 or p25. To verify that loss of p27 and p25 had no effect on the ability of AA 1–87 to bind the Arp filament, lysates prepared from cells depleted of p27 and p25 were mixed with 6X-His-AA 1–87, and then AA 1–87 and any associated proteins were pulled out using beads. The Arp filaments in the depleted lysate bound the resin equally well as undepleted controls (Figure 5A), providing clear demonstration that p27 and p25 are not required for dynamitin binding to the Arp1 filament.

Yeast dynamitin (Jnm1p) has been shown to interact with Arp11 (Arp10p; Clark and Rose, 2005) in a two-hybrid assay, but this interaction has not been reported in other species. No free pool of Arp11 or its binding partner p62 can be detected in cultured cells, and both proteins are insoluble when expressed in bacteria, so we could not perform direct binding assays on either using purified proteins. Unfortunately, it is not possible to use RNAi in experiments like those used to test the role of p27/p25 because depletion of either Arp11 or p62 also causes loss of Arp1 (Yeh et al., 2012). Despite these challenges, we were able to obtain anecdotal evidence indicating that p62 and Arp11 do not bind the dynamitin N-terminus. The purified Arp1 used in the direct binding experiment (Figure 4A) contains trace amounts of Arp11 and p62, CapZ, and p27/p25 (Bingham and Schroer, 1999; Eckley et al., 1999). Although Arp1

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**FIGURE 4:** Analysis of dynamitin AA 1–87 binding to Arp1 and actin. (A, B) Arp1 isolated from bovine dynactin by KI treatment and gel filtration (A; Bingham and Schroer, 1999) or G-actin (B) was dialyzed into G-buffer for 1 h and then mixed with Talon beads bearing purified 6X-His-AA 1–87 or a control (6X-His-Fis1TM). Samples were then analyzed by immunoblotting to detect Arp1 (2% of input and 2.5% of beads) or actin (2% of input and beads). Bait proteins were detected on the PVDF membrane by Ponceau S staining. (C) F-actin (see Materials and Methods) was incubated with buffer alone, α-actinin, BSA, or purified 6X-His-AA 1-87 (DM AA 1–87) for 30 min at room temperature. F-actin and bound proteins were then pelleted, and equal proportions of the supernatants (S) and pellets (P) were evaluated by SDS–PAGE, followed by Coomassie blue staining.

**FIGURE 5:** Analysis of the contributions of p27/p25 and CapZ to dynamitin–Arp1 binding. (A) Lysates of Cos-7 cells electroporated with p27 or control (ctr) siRNAs were incubated with 6X-His-AA 1–87 or a control (Fis1TM) before addition of Talon beads. Dynactin subunits in the eluates were detected by immunoblotting (1% of lysate and 5% of eluates). Bait proteins were detected by Ponceau S staining of the PVDF membrane. (B) Pools of cytosolic proteins (“lysate” lane) containing dynamitin (top: 20S fraction) or free CapZ (bottom; 4–5S fraction) were separated by sucrose gradient sedimentation and then mixed with beads bearing 6X-His-AA 1–87 or a control (6X His-TrbB). The 4–5S fractions (bottom left) and the bead eluates (bottom middle and right) were immunoblotted for CapZ.
reproducibly bound dynamitin N-terminus immobilized on beads in these experiments, none of the other proteins (p62, Arp11, CapZ, p27, or p25) did and were instead always found in the column flowthrough (unpublished data).

**DISCUSSION**

The organization of subunits in the dynactin molecule has been generally defined (Schafer et al., 1994; Eckley et al., 1999; Imai et al., 2006; Imai, Narita, Maeda, and Schroer, unpublished data), but a high-resolution structure is not available. We show here that dynamitin occupies a key position by serving as a bipartite binding platform that anchors the dynemin-binding p150Glued subunit to the Arp1 filament. The present biochemical demonstration that dynamitin binds Arp1 directly and specifically corroborates previous reports of close associations between these proteins in yeast (Clark and Rose, 2005; Amaro et al., 2008) and solves the long-standing question of how the shoulder-sidearm is tethered to the Arp1 filament.

Several lines of evidence suggest that the N-terminal portion of dynamitin is structurally distinct from the rest of the polypeptide. Sequence analysis predicts AA 1 to 100 to be largely unstructured except for a short α-helix (AA 55–62; predicted by DomPred; Marsden et al., 2002). In keeping with this prediction, the dynamitin N-terminus is highly labile to proteolysis (Tanimoto, Maeda, Imai, and Maeda, unpublished data; Maier et al., 2008). The rest of the sequence, by contrast, is predicted to be folded into a series of α-helices that we showed support homo- and hetero-oligomerization (Maier et al., 2008). The N-terminus is required for orderly homo-oligomerization in vitro (Maier et al., 2008) but does not contribute to interactions with the shoulder component, p24 (Tanimoto, Maeda, Imai, Maeda, Ketcham, and Schroer, unpublished observations). The present work reveals an important new role: Arp1 binding.

Many questions remain regarding the dynamitin/Arp1 interaction. It must be of sufficient affinity and stability to allow p150Glued to remain bound to the Arp1 filament even under tension, but soluble N-terminus can readily displace the entire shoulder/arm complex. The dynamitin N-terminus has also been reported to bind calmodulin (Yue et al., 2000). The ability of the same peptide to participate in multiple interactions may reflect its intrinsic disorder.

The dynamitin N-terminus contains a number of conserved charged residues, most of which are acidic, and this intrinsic negative charge may be accentuated by phosphorylation. The Arp1 primary sequence predicts a basic surface, suggesting that the interaction with dynamitin is electrostatic. Alanine-scanning mutagenesis of yeast Arp1 identified a cluster of charged amino acids near the C-terminus (K369, D371, E374, and D375; Clark and Rose, 2005) that appeared to be important for dynamitin and p150Glued binding. However, similar residues are present in actin, which we show does not bind dynamitin, suggesting that this may not be the binding site for dynamitin on Arp1.

We previously proposed that free dynamitin triggers subunit release by destabilizing contacts among dynamitin, p24, and p150Glued (Melkonian et al., 2007; Maier et al., 2008). The propensity of full-length dynamitin to oligomerize allows two exogenous protomers to be exchanged for two endogenous protomers (cartoon in Figure 6). Dynamitin AA 1–210, which lacks two of the three known oligomerization motifs, caused subunit release without remaining associated with the Arp filament (Maier et al., 2008), indicating that the subunit release does not require stable dynamitin–dynamitin binding. Our present findings verify that disruption does not depend upon dynamitin–dynamitin binding. The dynamitin N-terminus can trigger release of all four endogenous dynamitin protomers when it is overexpressed in vivo (Figure 2D; cartoon in Figure 6), whereas full-length dynamitin and AA 1–210 release only two protomers when mixed with purified dynactin in vitro (Melkonian et al., 2007; Maier et al., 2008). The simplest proposal that reconciles these findings is that disruption can occur via two different mechanisms. One (our earlier model) involves dynamitin–dynamitin binding and shoulder remodeling mediated by motifs in AA 100–403. The other (as defined here) involves direct binding of the dynamitin N-terminus to Arp1, leading to displacement of the entire shoulder/arm complex.

Previous work on dynamitin focused on its role in dynactin disintegration. The discovery that it binds Arp1 directly suggests that it may also play a role in dynactin assembly. Dynactin’s Arp filament is highly uniform in length, yet the mechanism by which length is specified is unknown. Polymerization of purified Arp1 yields filaments ~50 nm in length (Bingham and Schroer, 1999), which are longer than the ~35-nm filaments found in dynactin, so an extrinsic mechanism is assumed to exist to limit assembly. The unequal stoichiometries of dynamitin and Arp1 in dynactin (4 vs. 25; Imai, Narita, Maeda, and Schroer, unpublished data) precludes a simple one-to-one assembly model in which dynamitin templates a four-protomer Arp1 polymer. The fact that full-length dynamitin can displace only two of the four dynamitin protomers and leaves the other two bound to the Arp filament suggests that the two pairs of dynamitin protomers experience different environments, with the N-termini of two being closely bound to Arp1, and two being exposed on the periphery. Engagement of the N-termini of only one pair of dynamitins with Arp1 may allow for shoulder anchoring while leaving the remaining two dynamitin N-termini free to associate with extrinsic proteins such as calmodulin (Yue et al., 2000). We propose that the shoulder/arm complex binds
an Arp1 oligomer (a dimer or trimer, the two stable forms of isolated Arp1; Bingham and Schroer, 1999) via direct interactions with two dynamitin protomers. This assembly intermediate must also incorporate CapZ (and possibly via an undefined mechanism. A second “half-dynamitin” may form via association of Arp oligomer(s) with the pointed-end subunits Arp11 and p62, both of which are also able to bind Arp1 (Garces et al., 1999; Karki et al., 2000; Eckley and Schroer, 2003). These two half-molecules could join by Arp1/Arp1 annealing (Bingham and Schroer, 1999), leading to formation of the holocomplex, a structure whose overall integrity may involve interactions between shoulder/arm and pointed-end components.

MATERIALS AND METHODS

Constructs

Full-length chicken dynamitin–green fluorescent protein (GFP; as described in Quintyne et al., 1999) was used as the template for PCR-based subcloning of the constructs described in this article (see Table 1). Commercial vectors pRSET-A (Invitrogen, Carlsbad, CA) and pNTAP-C (Interplay Mammalian TAP system; Stratagene, La Jolla, CA) were used for making His-DM AA 1–87 and the TAP-tagged constructs respectively, as listed in Table 1. Commercial vectors pRSET-A (Invitrogen); Myc epitope (ab9106; AbCam, Cambridge, UK). Billerica, MA); Xpress epitope in 6X-His–tagged proteins (R91025, Invitrogen); Myc epitope (ab9106; AbCam, Cambridge, UK).

Recombinant protein purification

Plasmids encoding 6X-His–tagged recombinant proteins were transformed into BL21-CodonPlus (DE3)-RIL cells, and colonies were grown overnight in a 5-ml culture. One liter of Luria broth supplemented with 100 μg/ml ampicillin and 35 μg/ml chloramphenicol was inoculated with the overnight culture, grown to OD

Antibodies

p150Gluad and dynamitin: monoclonal antibody (mAb) product numbers 610473 and 611002, respectively, from BD Transduction Laboratories, San Jose, CA. Arp1 and p62: mAb 45A and 62B (Schafer et al., 1994). CapZ α: mAb 5B12 (Schafer et al., 1996). Arp11: affinity-purified rabbit antibody raised against the C-terminal of Arp11 (a gift from M. Way, Cancer Research United Kingdom, London, UK). p27: mAb 27A (Melkonian et al., 2007). p25: DCTNS polyclonal antibody (10182-1-AP; ProteinTech Group, Chicago, IL). p24: affinity-purified rabbit polyclonal antibody R5700 (Pfister et al., 1998). Giantin: G1/133 (Listed and Hauri, 1993). TGN46 (AbD Serotec). Tubulin: mAb DM1A (T9026; Sigma-Aldrich, St. Louis, MO) or rat mAb (YL1/2; AbD Serotec, Oxford, UK). TAP-tagged proteins: anti-calmodulin-binding protein epitope tag (07-482; Millipore, Billerica, MA); Xpress epitope in 6X-His–tagged proteins (R91025, Invitrogen); Myc epitope (ab9106; AbCam, Cambridge, UK).

The vectors, restriction sites, and primers used in making each construct are shown. RE, restriction sites.

| Construct   | Vector               | Up RE | Down RE | Primers                                                                 |
|-------------|----------------------|-------|---------|-------------------------------------------------------------------------|
| His-DM AA 1–87 | pRSET A            | BamHI | EcoRI  | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse ATAATCGAATTCTCAGCAGGACTCATAG                                    |
| Myc-DM      | pCMV5 myc2          | EcoRI | BamHI  | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse TATAGGGATCTCAGTCGCGGACCTTGGATG                                |
| Myc-DM AA 1–87 | pCMV5 myc2          | EcoRI | BamHI  | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse TATAGGGATCTCAGTCGCGGACCTTGGATG                                |
| Myc-DM AA 1–78 | pCMV5 myc2          | EcoRI | BamHI  | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse TATAGGGATCTCAGTCGCGGACCTTGGATG                                |
| Myc-p62 AA 370–467 | pCMV5 myc2          | EcoRI | BamHI  | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse TATAGGGATCTCAGTCGCGGACCTTGGATG                                |
| TAP DM      | pNTAP-C             | EcoRI | Xhol   | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse TATAGGGATCTCAGTCGCGGACCTTGGATG                                |
| TAP DM AA 1–87 | pNTAP-C             | EcoRI | Xhol   | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse TATAGGGATCTCAGTCGCGGACCTTGGATG                                |
| TAP DM AA 1–78 | pNTAP-C             | EcoRI | Xhol   | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse TATAGGGATCTCAGTCGCGGACCTTGGATG                                |
| TAP DM AA 1–210 | pNTAP-C             | EcoRI | Xhol   | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse TATAGGGATCTCAGTCGCGGACCTTGGATG                                |
| TAP DM AA 100–403 | pNTAP-C             | EcoRI | Xhol   | Forward GATTATGGATCGCCGACCCAAATAC                                      |
|             |                      |       |         | Reverse TATAGGGATCTCAGTCGCGGACCTTGGATG                                |

TABLE 1: Constructs used.
purified as in King et al. (2003). Dynamitin AA 1–210 (wt and L118P variant) and AA 1–81 were purified as in Maier et al. (2008).

**Circular dichroism spectroscopy**

All proteins were in 50 mM sodium phosphate, 150 mM NaCl, pH 7.0. CC1 was 0.13 mg/ml, AA 1–87 was 0.28 mg/ml, AA 1–78 was 0.18 mg/ml, AA 1–210 was 0.072 mg/ml, and AA 1–210 L118P was 0.342 mg/ml. Spectra were recorded in a 0.1-cm quartz cuvette at 25°C with a Jasco J-710 spectropolarimeter. The scans were collected from 195 to 260 nm using four consecutive accumulations, the average of which gave the final CD spectra. The continuing scans were done at scanning speed 50 nm/min with 4 s per point response time under standard sensitivity settings. The data pitch was 0.2 nm, and the bandwidth for samples was 1 nm. Each spectrum was subtracted by a buffer spectrum and further normalized by amino acid number and concentration. Eventually, measurements were reported in deg cm² dmol⁻¹.

The α-helical content was estimated from \( f_\alpha = (\theta_{222} + 2340)/(\text{[deg]} 	imes 30,300) \) (Chen and Yang, 1971), where \( \theta_{222} \) represents the mean residue ellipticity at 222 nm and \( f_\alpha \) is the fraction of α-helical content.

**Cell culture, transfection, and RNAi**

Cos-7 cells were grown in high-glucose DMEM (Life Technologies, Carlsbad, CA) supplemented with 10% fetal bovine serum (Atlas Biologicals, Fort Collins, CO), 1% l-glutamine, and penicillin/streptomycin (Life Technologies) and 1% MEM nonessential amino acids (Life Technologies) at 37°C with 5% CO₂. For most experiments using transient transfection, cells were grown to 70–90% confluency where

\[
a = \frac{\text{number of transfected cells}}{\text{total number of plated cells}}
\]

was scored per condition per experiment in three independent experiments.

For disruption experiments using purified proteins, 10 μg (9 × 10⁻⁶ mole) of purified bovine dynactin was mixed with a 25x molar excess (9 × 10⁻⁴ mole; see Maier et al., 2008) of the different recombinant dynamin constructs in a total volume of 0.1–0.2 ml of sedimentation buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM dithiothreitol (DTT)) and then incubated on ice for 30 min. In some cases, 9 × 10⁻⁴ mole of the purified recombinant dynamin fragments was added to 500 μg of Cos-7 detergent lysates (prepared as described later). After incubation, the samples were sedimented into a 5–20% sucrose gradient in a Beckman SW55 Ti rotor (Beckman Coulter, Brea, CA) at 30,500 rpm for 16 h at 4°C. Fractions (0.5 ml) were collected, and dynactin subunits were detected using SDS–PAGE, followed by immunoblotting. For analysis of dynactin integrity in cells overexpressing different dynamin constructs, cells were harvested 18–48 h posttransfection, lysed with lysis buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, and 0.1% Triton X-100) for 30 min at 4°C, and then centrifuged at 44,700 rpm in a Beckman SW55Ti rotor for 30 min. The supernatants were collected, and 500 μg to 1 mg of protein in a volume of 0.2 ml was sedimented into a 5–20% sucrose gradient and analyzed as described.

**Immunofluorescence**

Cells were rinsed with phosphate-buffered saline (PBS) and then fixed in −20°C methanol for 5 min or 4% formaldehyde in PBS at 37°C for 15 min. Formaldehyde-fixed cells were quenched with 50 mM ammonium chloride in PBS for 5 min and then permeabilized with 0.1% Triton X-100 in PBS/1% bovine serum albumin (BSA) for 15 min. All coverslips were incubated in blocking buffer (1% BSA in PBS) for 15 min before antibody addition. Primary antibodies were diluted in blocking buffer and added to cells for 1 h at room temperature in a humidified chamber, rinsed three times with blocking buffer, and then incubated with secondary antibody for 30 min. Before mounting, coverslips were rinsed three final times with blocking buffer. 4’,6-Diamidino-2-phenylindole (1 μg/ml) was included in the penultimate rinse. Cover slips were briefly rinsed in Milli-Q water (Millipore) and mounted using Fluoromount (Sigma-Aldrich) and dried overnight in the dark before imaging.

Immunofluorescence microscopy was performed on an Axiovert 100 LM microscope (Carl Zeiss, Jena, Germany). Images were captured using a CoolSnap digital monochrome camera and processed using SlideBook and Photoshop (Adobe). To quantify Golgi disruption, a minimum of 100–400 transfected cells were scored per condition per experiment in three independent experiments. To measure mitotic index, at least 1000 transfected mitotic cells were scored per condition per experiment in three independent experiments.

**Dynactin disruption and sucrose gradient analysis**

For disruption experiments using purified proteins, 10 μg (9 × 10⁻⁶ mole) of purified bovine dynactin was mixed with a 25x molar excess (9 × 10⁻⁴ mole; see Maier et al., 2008) of the different recombinant dynamin constructs in a total volume of 0.1–0.2 ml of sedimentation buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM dithiothreitol (DTT)) and then incubated on ice for 30 min. In some cases, 9 × 10⁻⁴ mole of the purified recombinant dynamin fragments was added to 500 μg of Cos-7 detergent lysates (prepared as described later). After incubation, the samples were sedimented into a 5–20% sucrose gradient in a Beckman SW55 Ti rotor (Beckman Coulter, Brea, CA) at 30,500 rpm for 16 h at 4°C. Fractions (0.5 ml) were collected, and dynactin subunits were detected using SDS–PAGE, followed by immunoblotting. For analysis of dynactin integrity in cells overexpressing different dynamin constructs, cells were harvested 18–48 h posttransfection, lysed with lysis buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, and 0.1% Triton X-100) for 30 min at 4°C, and then centrifuged at 44,700 rpm in a Beckman SW55Ti rotor for 30 min. The supernatants were collected, and 500 μg to 1 mg of protein in a volume of 0.2 ml was sedimented into a 5–20% sucrose gradient and analyzed as described.
frozen more than once. Each binding reaction contained 7.3 nmol of protein (e.g., 100 μg of 6X-His-AA 1–87) and 12.5 μl of beads. 6X-His-TrbB (a conjugal transfer protein in Escherichia coli; Hemmis et al., 2011) or 6X-His-Fis1ΔTM (Picton et al., 2009) proteins were used as negative control. To assay binding to dynactin, 7.3 nmol (100 μg) of 6X-His-AA 1–87 in binding buffer was bound to 12.5 μl of Talon beads for 30 min and then 10 μg purified bovine dynactin (Quintyne et al., 1999) in binding buffer was added and the mixture rotated for 30 min at 4°C. Beads were washed three times with binding buffer and proteins eluted with elution buffer (150 mM imidazole, 300 mM NaCl, 50 mM sodium phosphate, pH 7.0). For experiments involving siRNA-treated cells, 100 μg of 6X-His-AA 1–87 was incubated for 30 min at 4°C with 500 μg of cell lysis (prepared using lysis buffer; see earlier description). Talon beads were added, the mixture incubated another 30 min, and the beads were washed and eluted as described. For CapZ binding, proteins in a detergent lysate made from untreated Cos-7 cells were sedimented into a 5–20% sucrose gradient, and fractions containing free CapZ (4–5S) or dynactin (19–20S) were used.

Arp1 isolation and binding assays
Arp1 was isolated from bovine dynactin using gel filtration in the presence of 0.7 M KI as described previously (Bingham and Schroer, 1999; Eckley et al., 1999) and then cryostored in 50% sucrose. Samples were thawed, passed through a 0.22-μm syringe-tip polyvinylidene fluoride (PVDF) filter, and subjected to a second round of chromatography on a Superose12 column in column buffer (2 mM Tris-Cl, pH 8.0, 0.2 mM ATP, 0.2 mM CaCl2, 0.5 mM DTT) containing 0.7 M KI to remove sucrose and any oligomers. Fractions containing Arp1 monomers (and trace amounts of p62, Arp11, actin, CapZ α/β, p27, and p25; Bingham and Schroer, 1999) were pooled, dialyzed into column buffer without KI for 1 h, and used immediately in binding assays.

Actin copelleting assay
Human platelet actin (Cytoskeleton, Denver, CO) was resuspended in G-buffer (5 mM Tris-HCl, pH 8.0 and 0.2 mM CaCl2) at 1 mg/ml, preclarified by centrifuging at 150,000 × g in a SW55 Ti rotor for 90 min at 4°C, and then polymerized by addition of F-buffer (50 mM KCl, 2 mM MgCl2 and 1 mM ATP), followed by incubation for 4 h at room temperature. 6X-His-tagged-AA 1–87 was dialyzed into 10 mM Tris-Cl, pH 7.5, and 20 mM NaCl. A 40-μg amount of 6X-His-AA 1–87 (15.1 μM), α-actinin (2 μM), or BSA (2 μM) was added to 160 μg of human platelet actin and incubated for 30 min at room temperature. The mixtures were centrifuged at 150,000 × g in a SW55Ti rotor for 90 min at room temperature, and the supernatant was carefully separated from the pellet. Equal proportions of the supernatants and pellets were analyzed by SDS-PAGE, followed by Coomassie blue staining.

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