Mutually Exclusive Cytoplasmic Dynein Regulation by NudE-Lis1 and Dynactin

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Background: Cytoplasmic dynein performs a great variety of cellular functions using a diversity of regulators.

Results: NudE and dynactin compete for a common site within the dynein complex.

Conclusion: This mechanism prevents dual regulation by dynactin and LIS1 and suggests a major new mode of regulatory control.

Significance: This is the first insight into coordination of cytoplasmic dynein regulators.

Cytoplasmic dynein is responsible for a wide range of cellular roles. How this single motor protein performs so many functions has remained a major outstanding question for many years. Part of the answer is thought to lie in the diversity of dynein regulators, but how the effects of these factors are coordinated in vivo remains unexplored. We previously found NudE to bind dynein through its light chain 8 (LC8) and intermediate chain (IC) subunits (1), the latter of which also mediates the dynein-dynactin interaction (2). We report here that NudE and dynactin bind to a common region within the IC, and compete for this site. We find LC8 to bind to a novel sequence within NudE, without detectably affecting the dynein-NudE interaction. We further find that commonly used dynein inhibitory reagents have broad effects on the interaction of dynein with its regulatory factors. Together these results reveal an unanticipated mechanism for preventing dual regulation of individual dynein molecules, and identify the IC as a nexus for regulatory interactions within the dynein complex.

Cytoplasmic dynein is a 1.2 MDa protein complex that functions as the predominant microtubule minus end-directed molecular motor in most cell types. It is involved in a very wide range of cellular roles, but the underlying basis for its great functional diversity is poorly understood. A variety of regulatory factors appear to tailor the motor protein for specific cellular roles. These factors are responsible for recruitment of dynein to appropriate sites within the cell, proper temporal activation of motor activity, and modulation of mechanochemical behavior to accommodate different cellular tasks.

The most well-studied dynein regulatory factors are dynactin (3–7) and the LIS1-NudE/L complex (8–15). Each of these systems is involved both in dynein cargo recruitment and mechanochemical regulation, though via different mechanisms. Dynactin is itself a ~1 MDa multi-subunit complex. It was initially found to be required for dynein vesicular transport in vitro (4, 16), and to recruit the motor to mitotic kinetochores, and vesicular organelles (5, 17). Dynactin has also been found to increase dynein processivity by up to 2-fold in single molecule in vitro assays (7, 18, 19). The mechanism responsible for this effect is incompletely understood. Processivity of mammalian dynein is stimulated in both the plus- and minus-end directions along microtubules (20, 21), though yeast dynein with or without dynactin is primarily unidirectional (19). Although the microtubule binding CAP-Gly domain of the dynactin p150 glued subunit had been assumed to contribute to the enhancement of dynein processivity, recent studies showed no effect after its removal. Nonetheless, it was still required for complete dynactin function in vivo (6, 19, 22, 23).

LIS1 and its binding partners NudE and NudEL form a tripartite complex with dynein (15). LIS1 and NudE/L play critical roles in a subset of dynein functions, many of which appear to involve high-load dynein mediated transport. LIS1 is required for nuclear migration in neural progenitors and post mitotic neurons in vertebrates, and for nucleokinesis in several organisms (24–27). LIS1 and its interactors have also been implicated in translocation or reorientation of the entire microtubule cytoskeleton during mitosis and cell migration, as well as in centrosome and kinetochore dynamics, (1, 25, 28–32). The range of cellular functions involving LIS1 and NudE/L and their extent of overlap with dynactin-requiring functions remains incompletely resolved. Aspects of vesicular transport that involve dynactin were found not to require LIS1 (32, 33), though general (34–38) or conditional (39) roles for LIS1, NudE, and NudEL have been reported in other studies.

NudE and NudEL have been implicated in recruiting cytoplasmic dynein to cargo (1, 30, 40–42) as well as in recruiting LIS1 to dynein (15). We recently identified effects of LIS1 and NudE/L on dynein motor activity, and found them to be complex and distinct from those reported for dynactin (15). LIS1 stabilized the dynein-MT interaction during the transition state of the cross-bridge cycle, resulting in persistent force production under load. NudE alone inhibited the dynein-MT interaction. Strikingly, the tripartite complex of LIS1, NudE, and dynein transformed the motor to a persistent force-pro-
ducings state and enhanced multiple motor transport under load (15). This behavior is likely to be important in cellular scenarios requiring dynein to produce force against large opposing loads, such as nuclear migration (25).

Dynactin, NudE, and NudEL each interact with the tail region of the dynein complex. Dynactin binds via the central region of its p150Glued subunit to the N terminus of the dynein intermediate chain (IC)3 (2, 43, 44). NudE and NudEL have been found to bind to both the dynein IC and LC8 subunits (1, 15). NudE and NudEL were initially reported to contain a C-terminal dynein-interaction site (12), but a separate N-terminal site has also recently been reported as well (45, 46).

The current study was initiated to define the nature of the NudE-dynein interaction in greater detail. We find the primary binding site for NudE to lie within the dynein IC N terminus, the same region implicated in dynactin binding (2, 43). We observe clear competition between NudE and dynactin for dynein, identifying a novel mechanism for coordinating dynein regulators. The common interaction site is also a target for frequently used inhibitory probes, and our results, therefore, have important implications for phenotypic analysis of dynein function in vivo.

EXPERIMENTAL PROCEDURES

DNA Cloning and Protein Purification—Full-length mouse NudE was cloned into pGEX6P-1 (GE Biosciences) with N-terminal HA- and C-terminal His tags. NudE fragments were also cloned into this vector. p150Glued fragments were cloned from a full-length rat construct into pGEX6P-1 with an N-terminal FLAG-tag and human LC8 (accession number NM_003746) was also cloned into this vector. Dynein IC fragments from rat were also cloned into pGEX6P-1 with a Myc tag at the C terminus, or into pCDNA 3.1 (IC2C 1–260 and 123–280) or pEGFP (IC2C 1–100) for mammalian cell expression. For expression in bacteria, constructs were transformed into BL21-CodonPlus RIPL competent cells (Agilent Technologies) and expressed in Luria broth or Terrific broth. Protein production was induced by addition of 0.1–0.5 mM IPTG, and the culture was moved to 18 °C overnight. Bacteria were broken by sonication, and proteins were purified by batch incubation of a high speed supernatant with glutathione resin (GE Biosciences) in lysis buffer (PBS, 1 mM DTT, protease inhibitor mixture (Sigma), 1% Triton X-100) for 1–2 h at 4 °C. The beads were collected and washed extensively with lysis buffer in a column. The beads were then washed into PMEG buffer (100 mM PIPES, 5 mM EGTA, 4 mM MgCl, 0.1 mM EDTA, 0.9 mM glycerol, 1 mM DTT, pH 7.0) for freezing, or washed into PreScission protease cleavage buffer (50 mM Tris-HCl pH 7.0, 150 mM NaCl, 1 mM EDTA), and incubated with PreScission protease (GE Biosciences) overnight to cleave off the GST moiety according to the manufacturer’s instructions. The cleaved proteins were collected, concentrated using Amicon concentrators (Amicon) and flash frozen in liquid nitrogen for storage. For full-length NudE, the protein was then incubated with Talon resin for 1 h at 4 °C, the beads were washed extensively and eluted in PMEG containing 350 mM imidazole. Protein containing fractions were pooled, concentrated, and flash frozen in liquid nitrogen. Cytoplasmic dynein was purified from rat brain tissue as described (47) and frozen in liquid nitrogen. This dynein preparation is essentially free of dynactin (15). Baculovirus expressed LIS1 was purified as described (15).

Protein Biochemistry—Protein interaction experiments were performed in buffer A: 50 mM HEPES (pH 7.4), 150 mM NaCl, 2 mM DTT, 0.2 mg/ml BSA, 0.1% Nonidet P-40. For GST pull-downs, proteins were bound to glutathione-agarose and incubated with interactors in 350 μl total volume for 1 h at 4 °C. For HA or FLAG pull-downs, tagged proteins were first incubated with an equal molar amount of monoclonal anti-HA or anti-FLAG antibodies for 1 h on ice. The protein-antibody complexes were then bound to protein A beads (Invitrogen) for 1 h at 4 °C, washed twice to remove unbound protein, and then used for pull-downs as above. Beads were washed four times with 350 μl of buffer A before being processed for SDS-PAGE analysis. For pull-downs from brain lysate, a high speed supernatant (47) of rat brain was used. In competition experiments, beads coated with dynein interactors were incubated with ~4 nm purified brain dynein for 1 h at 4 °C, followed by three buffer washes of 350 μl each. The beads were then resuspended to volume, and the indicated amount of competitor protein was added for an additional hour, followed by four 350 μl washes. Beads were resuspended in 50 μl and processed for gel analysis. Western blots were processed on an Odyssey IR scanner (LI-COR Biosciences). Densitometry was performed using ImageJ software (NIH).

Antibodies—Antibodies used in this study were: monoclonal anti-dynein intermediate chain clone 74.1 (a gift from Dr. Kevin Pfister), or clone 70.1 (Sigma). Monoclonal and polyclonal anti-FLAG, anti-HA, anti-Myc, anti-GFP, and anti-LC8 (Abcam) and monoclonal anti-LIS1 (Sigma). Polyclonal anti-p150Glued (D’art) (48) and polyclonal anti-NudE/NudEL (1).

Pepsan—A membrane array spotted with overlapping dodecapeptides was generated based on mouse NudE (GenBank accession number Q9CZA6) C-terminal residues 192–344 (JPT Peptide Technologies, Berlin). Each spot contains ~5 nmol of a 12-amino acid long peptide that is covalently linked to a cellulose-β-alanine membrane. The sequence of peptides in adjacent spots are shifted C-terminally by two residues such that two neighboring spots overlap by ten residues (Fig. 2B). Before use, the membrane was reconstituted at room temperature in methanol for 5 min, followed by three 10-min washes with TBS. Blocking was performed for 1 h with 5% milk in TBS-T followed by a 1-h incubation simultaneously with primary monoclonal LC8 antibody at a dilution of 1:2500 and secondary anti-rabbit Alexa Fluor 680 (Invitrogen, A10043) at a dilution of 1:10,000 in 5% milk in TBS-T at room temperature. The membrane was then scanned using Odyssey Imaging System (LI-COR) to assess nonspecific interactions of the antibodies. 200 nm of recombinant purified human LC8 in 5% milk in TBS-T was incubated with the membrane overnight at 4 °C and for 1 h at room temperature the following day, followed by three 10-min washes with TBS-T, sequential probing with primary and secondary antibodies, and scanning as before. Scans of the membrane before and after incubation with LC8 were compared to...
identify the residues in the C terminus of NudE that are involved in binding LC8.

**Inhibition of CC1/NudE-Dynein Interaction with Antibodies**—Monoclonal anti-dynein intermediate chain antibodies 74.1 and 70.1 were incubated with rat brain purified cytoplasmic dynein at 10-fold molar excess for 60 min at 4 °C with gentle rotating. Antibody-dynein complexes or dynein alone were subsequently incubated with bacterially expressed GST-CC1 or GST-NudE on glutathione beads or beads alone for 90 min at 4 °C with gentle rotating in buffer A. In a similar experiment, GST-CC1 or GST-NudE either alone or after preincubation with a polyclonal anti-NudE/L antibody for 60 min at 4 °C were incubated with rat brain purified cytoplasmic dynein for 90 min at 4 °C with gentle rotating in buffer A. After the incubation periods, unbound dynein was separated from the beads by centrifugation. The beads were washed three times with buffer (15-fold bead volume) and resuspended in protein sample buffer. Coomassie-stained gels were scanned using the Odyssey IR system.

**RESULTS**

*NudE Binds to the Dynein Intermediate Chain N Terminus*—In a previous study we screened an array of dynein and dynactin subunits for NudE binding, and identified interactions with the dynein IC and LC8 subunits (1, 15). To gain further insight into the nature of these interactions we first determined where NudE bound within the dynein IC. This subunit consists of a short N-terminal α-helical coiled-coil, followed by binding sites for dynein’s three LC classes, a dimerization domain, and finally by a WD40 domain responsible for dynein heavy chain (HC) binding (49, 50) (Fig. 1B). Expression of the WD40 domain was toxic in HeLa cells, as previously reported (43), but the N-terminal fragments expressed well. Pull-downs with GST-NudE localized IC binding activity to the first 100 a.a. of rat IC2C (Fig. 1A). GST-NudE also interacted with the endogenous dynein complex as evidenced by the presence of the full-length dynein IC in the pulldowns (1). To test for a direct NudE-IC interaction, we expressed a series of GST-tagged N-terminal IC constructs in *Escherichia coli* (Fig. 1B). All constructs that contained the N-terminal coiled-coil domain were capable of interacting with native dynactin complex in rat brain lysates (Fig. 1C) indicating the proteins were correctly folded. The IC constructs were screened for interactions with purified HA-NudE, which mapped NudE binding to IC amino acids 1–70 (Fig. 1, B–D). NudE bound to each of six alternatively spliced variants of rat IC1 and IC2 (supplemental Fig. S1B), which all contain a common N-terminal predicted coiled-coil sequence, but diverge immediately downstream (2, 49) (supplemental Fig. S1A), arguing that IC isoform composition is unlikely to affect the IC-NudE interaction.

*Dynne LC8 Binds Directly to NudE*—Although LC8 interacts with both NudE and dynein, the specific role of the LC in the
NudE-dynein interaction is unknown. LC8 interacts with full-length NudE ((1); this study), but not with the truncation mutants NudE10–165, NudE10–191, and NudE1–191 (Fig. 2A and C; data not shown), consistent with binding of LC8 only to full-length, but not an N-terminal NudEL fragment (45). To map the site of LC8 binding more precisely we probed an array of overlapping dodecapeptides covering the C-terminal region of NudE (Fig. 2A, amino acids 192–344) using purified bacterially expressed LC8 as judged by Western blotting (Fig. 3A). We then used CC1 to map its interaction site within the C-terminal region of NudE (Fig. 2B). In support of this result we found LC8 to bind NudE1–218, but not NudE1–191 (Fig. 2C). The LC8 binding sequence does not contain the canonical LC8 binding motif identified in many other LC8 interactors (53), but whether this indicates a unique mode of LC8 binding is uncertain (Refs. 54, 55). We also found the NudE-dynein interaction to be unaffected by exposure of NudE to LC8 before, during, or after dynein-NudE binding (Figs. 2, D and F, supplemental Figs. S2, A and B). These results contrasted with the ability of excess ICs to disrupt the dynein-NudE interaction (Fig. 2, E and F). These data reveal that LC8 can bind directly and apparently independently to the IC and NudE polypeptides. Furthermore, the data indicate that dynein interacts with NudE predominantly via the ICs, rather than LC8.

*Overlap between Dynactin and NudE Binding Sites*—Dynactin binds to dynein through a direct p150Glued-IC interaction (2, 43, 44). The CC1 coiled-coil region (3, 43, 56) (Fig. 3A) of p150Glued and a downstream region (amino acids 600–811) (57) have each been implicated in dynein binding. To determine which region of p150Glued interacts with the native dynein complex, we performed pull-downs from rat brain lysate using a series of bacterially expressed GST-tagged p150Glued constructs (Fig. 3A). The CC1 region as well as a C-terminal subdomain, CC1B, each clearly pulled down brain cytoplasmic dynein, whereas an N-terminal CC1 subfragment, CC1A, as well as the p150Glued 600–811 fragment did not (Fig. 3B). None of the fragments pulled down dynactin, indicating that CC1 cannot exchange with full-length p150Glued in the dynactin complex. These results are consistent with previous work (19, 43, 56) and demonstrate that the CC1 region can bind to the native dynein complex in cytosol.

We then used CC1 to map its interaction site within the dynein ICs. CC1 binding specifically required the presence of the IC N-terminal 70 amino acids (Fig. 3C). These results identify a common IC region for NudE and dynactin binding, and a common range of interacting IC isoforms. As a further test of this conclusion we utilized two well characterized and widely used function-blocking anti-IC monoclonal antibodies, 74.1 and 70.1 (58–60). Each antibody recognized IC2C amino acids 1–70 as judged by Western blotting (Fig. 3D), localizing the epitopes within this region. Purified rat brain cytoplasmic dynein was incubated with an excess of each antibody and then exposed to beads coated with GST-NudE or GST-CC1. Alternatively, the GST-NudE and GST-CC1 beads were pre-exposed to a polyclonal anti-NudE antibody (1), followed by addition of...
dynein or an interaction between NudE and CC1. As further arguing against simultaneous binding of the two factors to and CC1 showed no evidence of specific binding to the beads, by NudE from FLAG-CC1 (Fig. 4, displacement of up to 53% of dynein (47% dynein bound to GST-NudE (Fig. 4, for displacement of dynein by the reciprocal binding partner prebound dynein to GST-NudE or FLAG-CC1, and then tested a higher affinity for dynein than NudE in this assay. We also conserved N-terminal coiled-coil domain of the IC.

NudE bind to overlapping, but not identical sites within the incubated with a preformed dynein-CC1 or dynein-NudE complex.

Interaction. Similar results were obtained if the antibodies were expected the NudE antibody specifically disrupted the binding of dynein to NudE, (1), but had no effect on the dynein-CC1 interaction. Similar results were obtained if the antibodies were incubated with a preformed dynein-CC1 or dynein-NudE complex (data not shown). These results reveal that dynactin and NudE bind to overlapping, but not identical sites within the conserved N-terminal coiled-coil domain of the IC.

Dynactin and NudE Compete for Binding to Dynein—In view of these results we tested relative abilities of NudE and dynactin to bind dynein. When simultaneously mixed with equal concentrations of NudE and CC1, purified brain dynein was pulled down predominantly by the latter (Fig. 4A), indicating CC1 has a higher affinity for dynein than NudE in this assay. We also prebound dynein to GST-NudE or FLAG-CC1, and then tested for displacement of dynein by the reciprocal binding partner (Fig. 4B). FLAG-CC1 caused a clear concentration-dependent displacement of up to 53 ± 22% (n = 3 experiments) of the dynein bound to GST-NudE (Fig. 4, B and D), and a similar amount of dynein (47 ± 11%, n = 3 experiments) was displaced by NudE from FLAG-CC1 (Fig. 4, B and D). The added NudE and CC1 showed no evidence of specific binding to the beads, arguing against simultaneous binding of the two factors to dynein or an interaction between NudE and CC1. As further evidence for the specificity of this assay, purified actin had no effect on dynein binding to GST-NudE (Fig. 4, B and D). We also tested the ability of NudE to interfere with the interaction between dynein and the complete dynactin complex pulled down from rat brain lysate using beads coated with GST-IC 1–250 (Fig. 4E). Preincubation of the IC beads with HA-NudE largely abrogated their ability to pull-down dynactin (Fig. 4E), confirming that NudE competes with endogenous dynactin for binding to IC. Preincubation of the beads with FLAG-CC1 or the 74.1 anti-IC antibody also blocked the interaction with dynactin as expected, while preincubation with actin had no effect (Fig. 4E). Together, these data demonstrate that NudE and dynactin compete for a common binding site at the N terminus of the dynactin IC.

We previously proposed that NudE acts as a scaffold to mediate the interaction of LIS1 with dynein (1, 15). We therefore tested whether LIS1 affects the competition between NudE and dynactin for dynein. Purified dynactin and LIS1 were bound to GST-NudE coated beads, followed by incubation with increasing amounts of FLAG-CC1. Unexpectedly, inclusion of LIS1 in the competition assay facilitated the release of dynein from the GST-NudE beads by FLAG-CC1 (Fig. 4, C and D).

**DISCUSSION**

How cytoplasmic dynein can contribute to an extremely diverse array of intracellular functions has remained a long-
standing question. The discovery of dynein regulatory complexes has shed light on how the motor might be targeted to and otherwise adapted for various duties, but it is currently unclear how the regulatory complexes might work together to coordinate dynein function. Our finding that dynactin and NudE compete for binding to dynein provides the first insight into how the activities of distinct dynein regulators may be coordinated.

Our results also identify the IC as an important nexus for dynein regulation. In contrast we found the dynein LC8 subunit not to contribute to the NudE-dynein interaction, but, rather, that NudE interacts with LC8 independently of dynein. This is consistent with a growing body of literature supporting a dynein-independent role for LC8 (61).

Our results indicate most dramatically that dynein molecules cannot be simultaneously occupied by dynactin and NudE. This arrangement is novel to the motor protein field, and further implies separate functional pools of dynein motors within the cell. Dynactin or NudE-LIS1 regulated dyneins appear to have distinct mechanochemical outputs (15, 18–20), and the mutually exclusive interactions of the regulators with the dynein ICs revealed in the current study strongly suggest that individual dynein molecules can be tailored to fit specific transport roles (Fig. 5). In view of its stimulatory effect on dynein processivity, dynactin might be specifically required for fast, long-range transport of smaller vesicular and macromolecular cargoes. Conversely, NudE-LIS1 prolongs the dynein force producing state, resulting in increased forces under multi-motor conditions (15). This mode of regulation seems certainly to be required for high load forms of dynein transport, such as nuclear migration (Fig. 5) (25, 26).

Whether dynactin and NudE-LIS1 are segregated to different forms of cargo for different modes of transport remains to be seen. Equally interesting is the possibility that NudE-LIS1-dynein...
nein and dynactin-dynein coexist on common cellular cargoes, adapting transport to distinct subcellular environments (Fig. 5). Indeed, recent evidence suggests that teams of multiple dyneins move membranous cargoes in vivo (62–64). Finally, it is also possible that the balance of a dynein regulatory factors may be subject to regulation in vivo, an issue of considerable further interest. Indeed, phosphorylation of the dynein IC has been reported to affect the affinity of this subunit for dynactin (65), and might conceivably contribute to switching between dynein regulatory factors. Further experiments are needed to test this hypothesis.

Such a shift between regulatory modes may require new tools to assay properly. Dynactin and NudE/NudE-LIS1 each control aspects of dynein recruitment to subcellular cargo, as well as dynein mechanochemical activity. This dual role will make it necessary to quantify relative effects on the number of dynein molecules associated with cargo versus the nature of dynein regulation. Changes in the affinity of dynein for dynactin relative to NudE and NudEL could alter the number of cargo-associated dyneins or shuttle dyneins between high-force and long travel distance regulators, or both.

Surprisingly, the addition of LIS1 caused dynein to be released from NudE more easily in the presence of FLAG-CC1 (Fig. 4, C and D). This result is unexpected given NudE-established role in recruiting LIS1 to dynein (1, 15, 45, 46). These are the first results suggesting that LIS1 may affect the NudE-dynein interaction and suggest further complexity in the interaction between dynein and its regulators. Additional information on the structural nature of the various dynein complexes will be needed to clarify this issue.

Our study also reveals broader effects for commonly used dynein inhibitory probes than has been assumed. The dynactin CC1 fragment, as well as the 74.1 and 70.1 monoclonal antibodies, have been favored reagents for cytoplasmic dynein inhibition in vivo. Our data indicate that the first two of these should interfere with both dynactin and NudE-LIS1 binding, while the third interferes preferentially with NudE. However, some of the more readily assayed dynein functions require both types of regulatory factor. For this reason, physiological assays for the specificity of the dynein and dynactin inhibitory agents may require more quantitative in vivo assays for dynein behavior than are currently available (39). Although the effects of each reagent provide insight into dynein function, their implications for understanding dynein regulation now appear less clear. Further development of probes specific for cytoplasmic dynein and for its individual regulatory factors will be needed to address these issues.

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Competition between Dynein Regulatory Factors

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