Nde1 Promotes Lis1 Binding to Full-Length Autoinhibited Human Dynein-1

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11 Abstract

12 Cytoplasmic dynein-1 (dynein) is the primary motor for the retrograde transport of 13 intracellular cargoes along microtubules. The activation of the dynein transport machinery 14 requires the opening of its autoinhibited Phi conformation by Lis1 and Nde1/Ndel1, but 15 the underlying mechanism remains unclear. Using biochemical reconstitution and cryo-16 electron microscopy, we show that Nde1 significantly enhances Lis1 binding to 17 autoinhibited dynein and facilitates the opening of Phi. We discover a key intermediate step in the dynein activation pathway where a single Lis1 dimer binds between the Phi-18 19 like (Phi^L) motor rings of dynein. In this "Phi^L-Lis1", Lis1 interacts with one of the motor 20 domains through its canonical interaction sites at the AAA+ ring and stalk and binds to 21 the newly identified AAA5, AAA6, and linker regions of the other motor domain. 22 Mutagenesis and motility assays confirm the critical role of the Phi^L-Lis1 interface. This 23 intermediate state is instantly and efficiently formed in the presence of Nde1, but Nde1 is 24 not part of the Phi^L-Lis1. These findings provide key insights into the mechanism of how 25 Nde1 promotes the Lis1-mediated opening of Phi dynein.

26 Introduction

Cytoplasmic dynein is a motor protein essential for transporting nearly all intracellular
cargoes toward the minus end of microtubules (MTs) in most eukaryotes. Dynein's
cargoes include organelles, proteins, RNA, viruses, and vesicles¹. Additionally, dynein
plays key roles in other cellular processes such as mitosis and organelle positioning^{1,2}.
Mutations affecting dynein or its regulatory proteins have been linked to
neurodevelopmental and neurodegenerative diseases, including spinal muscular atrophy
(SMA), ALS, Huntington's disease, lissencephaly, and microcephaly³⁻⁹.

34 The 1.4 MDa dynein complex contains pairs of six subunits. The largest subunit is the 35 dynein heavy chain (DHC), which contains the N-terminal tail domain and C-terminal 36 motor domain. The tail domain facilitates dimer formation, recruits the dimers of 37 intermediate chain (DIC), light intermediate chain (DLIC), and interacts with dynactin, cargo adaptors and other regulatory proteins¹⁰⁻¹³. The N-terminus of DIC (DIC-N) recruits 38 39 three pairs of light chains, Robl, LC8, and Tctex^{1,14,15}. DIC segments binding to LC8 and 40 Tctex, known as IC-LC tower, play a crucial role in assembling the dynein-dynactin-41 adaptor (DDA) complex¹¹. The dynein motor domain belongs to AAA+ (ATPases 42 Associated with diverse cellular Activities) protein families and consists of six AAA subdomains (AAA1-6)^{16,17}. AAA1 is the primary subdomain that hydrolyzes ATP to power 43 44 dynein motility along MTs. The dynein motor domain attaches to MTs via the coiled-coil 45 stalk and the MT-binding domain (MTBD) and connects to the tail through the linker domain located at the surface of the AAA+ ring¹⁷⁻²⁰. 46

47 Dynein alternates between two key conformational states: the autoinhibited Phi (similar to the Greek letter " ϕ ") and the open conformations^{14,21}. Phi dynein adopts a compact 48 49 structure that limits its interactions with MTs and dynactin, which serves to minimize 50 unnecessary ATP hydrolysis when motor protein is not engaged in active transport^{14,22}. 51 Opening of the Phi enables dynein to assemble with dynactin and the cargo adaptor. allowing processive movement along MTs^{10,14}, a process supported by dynein regulators 52 53 Nde1/Ndel1 and Lis1^{3,15,23-30}. However, the underlying mechanism promoting Phi to open 54 transition is unclear.

Lis1, the first gene identified in relation to a neuronal migration disease, plays a crucial 55 role in dynein-related function^{3,11,31-39}. Lis1 possesses an N-terminal LisH domain that 56 57 facilitates dimerization and a C-terminal WD-40 β-propeller domain that binds to 58 Nde1/Nde11, dynein, and other proteins³. Both domains are important at different stages 59 of dynein activation⁴⁰⁻⁴². The LisH domain interacts with dynactin p150 and DIC-N, thus 60 promoting the recruitment of dynactin and adaptors at the later stage of dynein 61 activation¹¹. WD-40 domains of Lis1 can directly bind to the dynein motor domain at the AAA3-AAA4 sites (site-ring) and stalk coiled-coil (site-stalk)^{11,38,43}. Functional studies 62 63 proposed that Lis1 facilitates the formation of highly processive DDA complex by favoring the release of Phi dynein and stabilizing open dynein^{34-37,44}. Lis1 binding is thought to be 64

65 incompatible with Phi dynein based on the steric clash when docking a Lis1 β-propeller to the motor domains of Phi dynein^{3,35,37,44}. However, there is no direct biochemical or 66 structural evidence of whether Lis1 can open Phi dynein and intermediary states that 67 facilitate opening of Phi dynein remain unclear. A recent structural study on tail-truncated 68 69 yeast dynein motor domains reported a "Chi" conformation, in which two Lis1 dimers are wedged between two AAA+ rings³⁷, suggesting that two Lis1 dimers between the motor 70 71 domains are required to crack open Phi dynein. Furthermore, it is well-known that the 72 AAA+ ring undergoes substantial conformational changes under different nucleotide-73 binding states, which can potentially regulate dynein-Lis1 binding. A more recent work 74 has demonstrated that specific nucleotide "codes" at the three variable nucleotide-binding 75 sites (AAA1, 3, and 4) govern the stoichiometry of dynein–Lis1 interactions by tuning their binding affinity at two distinct locations⁴⁵. However, the intermediate structure for full-76 77 length human dynein alone bound to Lis1 is still lacking and whether dynein forms a Chi 78 conformation at the initial state of its activation remains unclear.

Nde1 and its paralog Ndel1^{12,23}, are critical for all dynein-mediated functions in cell 79 division^{28,46}, cargo trafficking^{47,48} and neuronal migration^{49,50}. Nde1/Ndel1 is predicted to 80 81 be composed of the N-terminal coiled-coil domain and the C-terminal unstructured 82 region⁵¹⁻⁵³. The coiled-coil domain interacts with DIC-N, which overlaps with the DIC binding site of the p150 subunit of dynactin^{52,54,55}. It also interacts with the WD-40 83 84 domains of Lis1, and its binding site on WD-40 domain overlaps with the Lis1 binding site of dynein^{26,52}. The multi-protein interaction modes of Nde1/Ndel1, along with its 85 86 overlapping binding sites with other proteins, make the functional interpretation of Nde1/Ndel1 elusive. It has been proposed that Nde1 tethers Lis1 to dynein^{12,26,56-58} and 87 promotes Lis1-mediated activation of dynein^{11,26} (**Fig. 1a**). According to this model, Phi 88 dynein first adopts an 'open' conformation. Lis1 stabilizes the open dynein by preventing 89 it from transitioning back to Phi, thus favoring the assembly of the DDA complex^{11,34,35,44}. 90 Consistent with this model, overexpression of Lis1 can rescue the deletion of Nde1/Ndel1 91 in cells^{58,59} and Nde1/Ndel1-mediated recruitment of Lis1 to dynein enhances DDA 92 assembly in vitro²⁶. An alternative model suggests that Ndel1 negatively regulates dynein 93 94 activation by competing with p150 for DIC-N binding and by sequestering Lis1 away from 95 dynein⁵². Consequently, how Lis1 and Nde1/Ndel1 form a complex with dynein and 96 promote the opening of Phi dynein is not well understood.

97 To understand the mechanism by which Nde1/Ndel1 and Lis1 prime dynein for the DDA 98 assembly, we investigated how human Lis1 and Nde1 affect the conformational states of 99 full-length human dynein using biochemical reconstitution and electron microscopy. We 100 showed that Nde1 promotes the formation of the dynein-Lis1 complex. Using negative-101 stain EM, we provide direct evidence that Lis1 or Nde1 alone has little effect on the 102 equilibrium between the Phi and open conformations of dynein, but Lis1 and Nde1 103 together significantly bias the equilibrium toward the open conformation, demonstrating 104 that Nde1 acts like a molecular chaperone to promote Lis1-mediated opening of dynein. 105 Cryo-EM imaging of dynein-Lis1 in the presence or absence of Nde1 captures a new 106 intermediate and rate-limiting state during the dynein activation process, characterized by 107 a single Lis1 dimer binding between the motor domains of Phi dynein. Lis1 binding to Phi 108 dynein causes the rotation between two motor domains to relieve the steric clash, forming 109 a Phi-like dynein and Lis1 complex, named "Phi^L-Lis1". While Lis1 binds to one of the 110 motor domains through its canonical interaction sites, it also forms new interfaces 111 between the other motor domain at the AAA5, AAA6, and linker regions. Mutagenesis at 112 the novel interfaces together with single molecule motility assays supported the critical 113 role of this Phi^L-Lis1 during the dynein activation process. Collectively, our findings shed 114 light on the roles of Nde1 and Lis1 in the dynein activation pathway.

115 Results

116 Nde1 promotes Lis1 binding to Phi dynein and cooperatively releases dynein 117 autoinhibition

118 To determine whether Nde1 promotes Lis1 binding to dynein, we performed mass 119 photometry (MP) assays to assess the Lis1 and Nde1 binding to full-length human dynein 120 under different conditions. In the absence of Nde1, Lis1 alone exhibited an increased 121 binding to dynein in a time-dependent manner, with the formation of a 37% 1:1 122 dynein:Lis1 (DL) complex within 60 minutes. The inclusion of Nde1 significantly enhanced 123 the dynein-Lis1 binding (69% of complex formation) in less than a minute, consistent with 124 Nde1/Ndel1-mediated tethering of Lis1 to dynein^{12,26,57} (Fig. 1b). To further test if Nde1 125 preferentially recruits Lis1 to Phi or open conformation, we performed these assays using 126 the Phi mutant of dynein¹⁴ that only forms open conformation. Lis1 was readily bound to 127 open dynein, but the addition of Nde1 did not further enhance Lis1 binding to dynein 128 (Extended Data Fig.1), suggesting that Nde1 is required for Lis1 recruitment to Phi 129 dynein. These results are also consistent with the previous observation that the Nde1 130 addition does not further enhance Lis1-mediated activation of DDA complexes assembled with the Phi dynein mutant²⁶. Interestingly, the formation of the dynein-Lis1-Nde1 tripartite 131 132 complex was not observed (Fig. 1b and Extended Data Fig.1), regardless of different 133 nucleotide conditions (Extended Data Fig. 2). Although previous reports indicated that Nde1 can interact with DIC-N in single molecule imaging^{12,26} and pull-down assays^{26,52}, 134 135 we also did not detect the dynein-Nde1 complex (Fig. 1b, and Extended Data Fig. 2), 136 suggesting that Nde1 rapidly dissociates from the complex after handing off Lis1 to 137 dynein¹².

To determine whether Nde1-mediated Lis1 recruitment to dynein shifts the equilibrium between Phi and open conformations, we used the negative stain EM imaging¹⁴ to quantify the ratio of Phi dynein in the presence and absence of Lis1 and Nde1. Specifically, we used freshly prepared dynein with ~50% of motors forming the Phi and then incubated 142 dynein with Lis1 and Nde1 (Fig. 1c and Extended Data Fig. 3) in the presence of ATP.

143 We found that Lis1 or Nde1 alone does not change the Phi ratio compared with the control

144 (Fig. 1d-e). However, the Phi ratio decreased 44% when we incubated dynein with both

Lis1 and Nde1 at a ratio of 1:2:2 (Fig. 1d-e). Collectively, our results demonstrate that

146 Nde1 specifically promotes Lis1 binding to Phi dynein and facilitates opening of this

147 autoinhibited conformation.

148A novel Phi^L-Lis1 structure

149 Our MP results suggest that there is a rate-limiting step of Lis1 binding to Phi dynein, and 150 this step can be significantly accelerated by the addition of Nde1 (Fig. 1b). We used cryo-151 EM to capture the potential intermediate states to reveal the structural basis of this 152 process. We focused on the particles that form the autoinhibited dynein (Fig. 2a, b and 153 **Extended Data Fig. 4, 5**). In the absence of Nde1, we unexpectedly observed a novel 154 structure in which a Lis1 dimer is wedged between the two stacked motor rings of Phi dynein. The dynein in this complex shows a compact conformation, similar to but not the 155 156 same as the previously reported Phi structure¹⁴ and we referred this complex as the "Phi^L-157 Lis1" (Fig. 2a, c and Supplementary Video 1). Despite excess Lis1, nearly half of Phi 158 dynein does not bind to Lis1 (42.7% Phi vs. 57.3% Phi^L-Lis1) (Fig. 2a). Notably, we did 159 not observe the "Chi" (two Lis1s bound to dynein), suggesting that the "Chi" conformation 160 may be specific to yeast dynein or may form when truncated dynein containing only the 161 motor domains is used instead of full-length motor³⁷.

In the Phi^L-Lis1 structure, Lis1 appears on the same side as the IC-LC tower of dynein (front side), independent of the presence of Nde1 (**Fig. 2a, b**). Based on this structural observation, along with previous evidence indicating that DIC-N can bind to both Lis1¹¹ and Nde1¹², we speculate that the transient dynein-Lis1-Nde1 forms only on the IC-LC tower side. Although we do not observe clear cryo-EM densities for the DIC-N, it is possible that the DIC-N can weakly interact with Lis1, thus recruiting Lis1 to the front side of dynein.

169 Previous results suggest Lis1 can affect dynein's mechanochemical cycle and nucleotide state^{11,60,61}. The high-resolution structure of the motor domain enabled us to identify and 170 171 compare the nucleotide states of the motor domains in Phi and Phi^L-Lis1 structures. 172 Specifically, we found that AAA1 pockets of both motor domains exhibit the same ADP-173 Mg²⁺ density (Extended Data Fig.6a, b), accompanied by a flexible sensor-I loop, 174 indicating the intermediate state of Pi releasing (Extended Data Fig. 6c, d). Similarly, 175 AAA3 pockets show clear ADP binding (Extended Data Fig.6a, b), indicating that Lis1 176 binding does not influence the nucleotide states of the motor domains of autoinhibited 177 dynein.

We next premixed equimolar Nde1 and Lis1, then added dynein to achieve a final 1:2:2 ratio of dynein:Lis1:Nde1 (**Fig. 2b**). Strikingly, we obtained a similar Phi^L-Lis1 structure, but no longer observed the Phi dynein alone in the presence of Nde1 (Fig. 2a, b and
Supplementary Video 1). This is consistent with the MP analysis showing that Nde1
instantly promotes Lis1 binding to dynein (Fig. 1b) and forming Phi^L-Lis1. These results
suggest that Phi^L-Lis1 formation is a rate-limiting intermediate state before dynein
opening. Remarkably, none of the 3D classes shows clear Nde1 density in Phi^L-Lis1,
suggesting that Nde1 is not part of this complex.

We also analyzed individual motor domains of open dynein and did not observe an increase in the propensity of the dynein-Lis1 complex compared to open dynein alone in the presence of Nde1 (**Extended Data Fig. 7**). Consistent with MP of the Phi dynein mutant¹⁴ (**Extended Data Fig. 1**), these results show that Nde1 promotes Lis1 binding to Phi dynein to form Phi^L-Lis1, whereas Lis1 can readily bind to open dynein and does not require Nde1.

192 Lis1 induces a relative rotation in the Phi conformation to accommodate its binding

Consistent with previous reports^{35,37,44}, our structural analysis reveals a severe steric 193 194 clash between Lis1 and Phi motor domain A (Fig. 3a) and this needs to be relieved to 195 accommodate Lis1's binding in Phi^L-Lis1 (Fig. 3b). Comparing Phi to Phi^L-Lis1 reveals a 196 relative rotation between the two motor domains, which results in the groove on the front 197 side of dynein becoming larger than the corresponding groove on the back side (Fig. 3c, 198 d). The enlarged groove on the front side allows Lis1 to fit between Phi^L motor domains. 199 However, docking of Lis1 to the back side with a smaller groove shows severe clashes, 200 explaining why there is only one Lis1 present in Phi^L-Lis1 (**Fig. 3e**). The rotation between 201 the motor domains in Phi^L-Lis1 also causes a slight anticlockwise twist in the neck region 202 (Fig. 3d). This twist likely promotes the unwinding of the tail, generating a trend toward 203 an open conformation of dynein. We concluded that Lis1 binding induces a rotation of Phi 204 dynein motor domains to avoid steric clash with Lis1 (Fig. 3 and Supplementary Video 205 1).

206 Novel interactions identified in Phi^L-Lis1

207 In Phi^L-Lis1, WD-40 domains of Lis1 interact with both motor domains of dynein (MD-A 208 and MD-B). While Lis1 interacts with MD-B through its canonical interaction sites at the AAA3, AAA4, and AAA5 regions^{38,43,62}, we observed previously uncharacterized 209 210 interaction sites of Lis1 with the linker, AAA6, and AAA5 regions of MD-A (Fig. 4a, 211 **Extended Data Fig. 8 and Supplementary Video 1).** Sequence alignment of Lis1 212 homologs shows that MD-A and Lis1 interface is highly conserved among higher eukaryotes but less conserved in yeast (Extended Data Fig. 9), suggesting different 213 214 regulatory roles of Lis1 between higher eukaryotes and yeast. Structural comparison 215 indicates that two motor domains in Phi^L-Lis1 adopt an almost identical conformation. The 216 root-mean-squared-displacement of alpha carbon atoms (C α -RMSD) of the two motor 217 domains was 0.513 Å (Fig. 4b). Additionally, the MD-B bound with Lis1_{ring} and Lis1_{stalk} in

our results also shows no significant difference from the structure of Lis1 bound to the human dynein motor domain³⁸ (C α -RMSD: 0.868 Å) (**Fig. 4c**), suggesting Lis1 binding

human dynein motor domain³⁸ (Cα-RMSD: 0.868 Å) (Fig. 4c), suggestin
 does not induce structural changes within an individual motor domain.

The interactions between MD-A and Lis1_{ring} are notably compact (**Fig. 4a and Extended Data Fig. 8**). The WD-40 domain of Lis1 interacts with dynein MD-A at regions distributed across the linker, AAA6, and AAA5 region (**Fig. 4d-f**). Specifically, at the linker-Lis1 binding site, the side chains of M329 and E300 of Lis1 engage in hydrophobic and polar interactions with the side chains of V1563, P1562 and H1559 of dynein. Additionally, K303 and S304, located on the flexible loop of the Lis1 WD-40 surface, interact with R1621, D1556, and E1622 of linker region (**Fig. 4d, Extended Data Fig. 8a**).

Within the AAA6 interaction region, the interface is characterized by polar interactions involving N203, D205, and Q222 of the Lis1 WD-40 domain and K4089, N4085, Q4117, and S4115 of MD-A. The side chain of K4089 forms a salt bridge with the side chain of D205 and establishes a hydrogen bond with the oxygen atom in the main chain of N203. Q222 and D205 of Lis1 also form polar interactions with S4115 and Q4117 and N4085 residues of dynein (**Fig. 4e, Extended Data Fig. 8b**).

234 The AAA5-Lis1_{ring} WD-40 interface shows a more compact interaction (Fig. 4f). This 235 interface is mainly composed of residues Q3636, S3613, T3612, D3616, D3617, A3618, 236 and K3621 of dynein MD-A and Y225, C226, G190, H191, M172, and D192 of the Lis1 237 WD-40 domain (Fig. 4f and Extended Data Fig. 8b). Notably, the side chain of Y225 of 238 Lis1 forms a polar interaction with the side chain of Q3636. Additionally, the side chain of 239 D3616 and the main chain of D3617 form hydrogen bonds with the main chain of C226. 240 The side chain of A3618 forms hydrophobic interaction with the main chain of G190, while 241 residues M172 and H191 of Lis1 form hydrophilic interaction with the side chain of K3621. 242 Additionally, novel interactions are formed between MD-A and MD-B in the Phi^L-Lis1, 243 compared with canonical Phi (Extended Data Fig. 10), suggesting that the dynein Phi^L-244 Lis1 is a stable conformation.

244 LIST IS a Stable conformation.

245 The Phi^L-Lis1 interface regulates Nde1-dependent dynein activation

246 To evaluate whether the new interaction sites we detected between Lis1 and dynein MD-247 A in Phi^L-Lis1 are critical for activation of dynein, we introduced three sets of Lis1 248 mutations targeting the interfaces that interact with the linker, AAA6, and AAA5 of dynein. 249 Key residues of Lis1_{ring} at each interface were mutated to charged residues or alanine to disrupt these interactions (Lis1^{linker}: E300K, K303E, S304R and M329A; Lis1^{AAA6}: N203K, 250 D205K and Q222A; Lis1^{AAA5}: M172K, D192K, Y225A, C226D). Similar to wild-type Lis1 251 252 (WT Lis1), these Lis1 mutants formed homodimers and interacted with Nde1 (Fig. 4g). 253 However, the binding efficiency of Nde1 was reduced for the Lis1^{AAA5} mutant (**Fig. 4g**), 254 suggesting that the AAA5-Lis1ring interface may share a region involved in Nde1 binding 255 to Lis1 (Extended Data Fig. 11). The Lis1 mutants also bound to dynein, and the dynein binding efficiency of Lis1 was increased with Nde1 (Fig. 4h). Notably, we detected a mass
population corresponding to dynein bound to one Nde1 and two Lis1^{AAA6} mutants,
indicating that mutations to the AAA6 interaction site of Lis1 prevent dissociation of Nde1
from the dynein-Lis1 complex (Fig. 4h).

260 To determine how these mutations affect activation of dynein motility, we assayed single 261 molecule motility of complexes assembled with wild type dynein, dynactin, and the 262 BicDR1 adaptor (DDR) on surface-immobilized MTs in vitro in the presence and absence of Lis1 and Nde1. Consistent with our previous observations²⁶, Lis1 enhanced the run 263 264 frequency of DDR about 3-fold, and Nde1 and Lis1 together increased the run frequency 265 15-fold (Fig. 4i, j and Supplementary Video 2). In the absence of Nde1, Lis1 mutants 266 activated dynein motility at similar levels of WT Lis1. In the presence of Nde1, Lis1^{linker} 267 and Lis1^{AAA5} triggered activation of DDR motility similar to WT Lis1 (Fig. 4i, j). In 268 comparison, Nde1 failed to enhance Lis1^{AAA6}-mediated dynein motility, suggesting that 269 Lis1^{AAA6} cannot form the stable Phi^L-Lis1 complex and open the Phi conformation (Fig. **4i**, **j**). Together with MP results, we demonstrate that mutations in the Lis1^{AAA6} interface 270 271 disrupt Nde1-mediated opening of the Phi conformation by Lis1, highlighting the 272 importance of the Phi^L-Lis1 structure in the dynein activation pathway.

273 Discussion

274 In this study, we investigated the structure and mechanism of how Lis1 and Nde1 rescue 275 dynein from autoinhibition prior to the assembly of active dynein transport machinery. 276 Using negative stain EM imaging, we directly showed that Nde1 and Lis1 cooperatively 277 promote the opening of Phi dynein, whereas neither Nde1 nor Lis1 alone exhibited a 278 significant effect on Phi opening (Fig. 1c-e). Lis1 can readily bind to open dynein 279 (Extended Data Fig. 1) and facilitate the assembly of DDA complexes and Nde1 addition does not further enhance DDA motility²⁶. Despite that Lis1 alone can bind to the Phi 280 281 dynein and potentially open this autoinhibited conformation⁴⁵, we show that Nde1 282 facilitates Lis1 to dock onto Phi dynein more efficiently and promote its switch it to the 283 open state before DDA assembly in this work. Similar to molecular chaperones, Nde1 284 rapidly dissociates from dynein after handing off Lis1, promoting dynein-Lis1 complex 285 formation but not existing in the final complex. Because Nde1 has an overlapping binding 286 site with the p150 subunit of dynactin on DIC-N¹², dissociation of Nde1 from DIC-N after 287 tethering Lis1 to dynein may enable efficient recruitment of dynactin to dynein-Lis1 288 complexes.

Our structural and functional studies of dynein-Lis1 complexes revealed a key intermediate step on Lis1 and Nde1 mediated opening of dynein. Although a dynein dimer contains Lis1 binding sites on each motor domain, our MP assays showed that Phi dynein recruits a single Lis1. Using cryo-EM imaging, we revealed a Phi^L-Lis1 structure in which the two AAA+ rings of Phi dynein rotate slightly backward to accommodate Lis1 binding to the front side. This rotational motion reduces the spacing between the AAA+ rings,
thereby preventing Lis1 binding to the back side. Preferential binding of Lis1 to the front
side may be due to the IC-LC tower, which is located at the front side of the Phi motor.

The Phi^L-Lis1 is fundamentally distinct from the previously reported Chi of yeast dynein 297 298 monomers³⁷. The Chi is stabilized by two Lis1, one on each side, and adopts a more open 299 and extended conformation compared to the Phi^L-Lis1 motor domains (Extended Data 300 Fig. 12). In comparison, our study utilized full-length, wild-type human dynein and we 301 could not detect Chi dynein even when we used this conformation as a reference during 302 cryo-EM image processing. It is possible that isolated dynein motor domains may prefer 303 to recruit two Lis1s and form more extended Chi. In comparison, full-length dynein readily 304 forms the compact Phi and structural constraints imposed by the tail domain may restrict 305 the relative movement of the Phi motor domains. Lis1 binding to the front side of Phi^L 306 dynein reduces the spacing on the back side, thereby preventing the formation of Chi. 307 Most interaction sites located at Lis1ring surface of Chi-Lis1 are also present in that of Phi^L-308 Lis1 (Extended Data Fig. 9). In comparison, Phi^L-Lis1 exhibits more compact interactions 309 (Fig. 4 and Extended Data Fig. 8,10). Consistent with Phi^L-Lis1, we did not detect complexes with one dynein and two Lis1s in MP, suggesting that Phi^L-Lis1, rather than 310 311 Chi. is the stable intermediate of full-length human dynein.

312 The mutagenesis of the interactions between Lis1 and the AAA6 subdomain of MD-A in 313 Phi^L-Lis1 disrupts Nde1's ability to promote Lis1-mediated dynein activation, confirming 314 that the Phi^L-Lis1 is a key intermediate in the dynein activation pathway. However, these 315 mutations did not disrupt the mutant Lis1's ability to increase the run frequency of dynein 316 several-fold on its own. This is because WT dynein can be either in open or Phi 317 conformations with near equal probability in our conditions. Our model predicts that 318 mutant Lis1 can still bind and enhance DDA assembly of open dynein without Nde1. 319 However, it cannot further enhance dynein motility synergistically with Nde1 because this 320 mutant is deficient in forming Phi^L-Lis1.

321 Based on our results and previous observations, we propose a mechanism underlying 322 dynein activation by Lis1 and Nde1. In the absence of Nde1, Lis1 alone can bind to both 323 DIC-N and dynein motor domains¹¹, inducing a conformational change from canonical Phi 324 to Phi^L. However, the efficiency of this process is low and Lis1 cannot open Phi^L dynein 325 on its own (Fig.1c-e, 2a). In the presence of Nde1, Lis1-Nde1 is initially recruited to DIC-326 N positioned at the front side of Phi dynein, facilitating more efficient binding of Lis1 to the 327 front side of Phi. The local enhancement of Lis1 near the Phi motor by Nde1 facilitates 328 more efficient binding of Lis1 to Phi dynein (Fig. 5 step-i, -ii). Nde1 and Lis1 form a 329 transient Phi^L-Lis1-Nde1 complex (Fig. 5 step-i). Nde1 dissociates spontaneously, 330 leading to Phi^L-Lis1 formation (Fig. 5 step-ii). Lis1 binding induces a slight backward 331 rotation of the two motor rings in Phi^L-Lis1 (Fig. 3c), suggesting an intermediate state 332 prior to an open state. Additionally, a slight twist in the neck region, caused by motors

rotation and likely inducing an unwinding trend in the tail, may also contribute to dynein opening (**Fig. 3d**). Subsequently, Phi^L-Lis1 transitions to open dynein-Lis1 with the assistance of Nde1 (**Fig. 5 step-iii**). The binding of Lis1 to dynein facilitates DDA assembly and activates dynein motility by recruiting the p150 subunit of dynactin to dynein through its LisH domain¹¹ (**Fig. 5, step-iv**). Future studies are required to understand how Nde1 hands off Lis1 to dynein and why it rapidly dissociates from dynein. In addition to its tethering role and facilitating the formation of Phi^L-Lis1, it remains to be determined

340 whether Nde1 has additional roles in helping Lis1 convert Phi^L-Lis1 to the open dynein.

341 Methods

342 Cloning and expression

The plasmid encoding full-length human dynein⁶³ was generously provided by Andrew 343 344 Carter (His-ZZ-TEV-SNAPf DHC1 IC2C LIC2 Tctex1 Robl1 LC8, Addgene plasmid #111903). The His-ZZ-TEV-SNAPf tag is fused to the N-terminus of the dynein heavy 345 346 chain. Human Lis1 and Nde1 (residues N-terminal 1-190 residues, which functions 347 similarly to full-length Nde1 in the single motility assay²⁶), and the mouse BIDCR1 gene, 348 were each cloned individually into the pOmniBac backbone. The constructs featured a 349 ZZ-TEV tag at the N-terminus and a SNAPf tag at the C-terminus. Lis1 mutants containing 350 point mutations were generated using purchased DNA fragments (IDT) containing the 351 mutations and inserted into the plasmid backbone. The mutations were verified by Oxford 352 Nanopore full-plasmid sequencing. The constructs used in this study are listed in 353 Supplementary Table 1.

354 These proteins were all expressed in insect sf9 cells, as describe previously^{14,19,26} with 355 slight modifications. Briefly, Bacmid DNA isolated from the from DH10MultiBac competent 356 cells (Geneva Biotech) were transfected into the in sf9 insect cells with the Cellfectin® II 357 (Gibco) reagent. Protein expression in sf9 cells was accomplished by infecting them with 358 P2 virus at a cell density of 2.5 million cells/mL. For dynein expression, 28 mL of P2 virus 359 was added into a 1.4 L culture of sf9 cells. For Lis1, Nde1, and BicDR1, 7 mL P2 virus 360 was used to infect the 0.7 L sf9 cells. Cells were harvested after 75 hours by centrifugation 361 at 1000 rcf for 15 minutes at 4°C. The cell pellets were flash-frozen in liquid nitrogen and 362 stored at -80°C.

363 **Protein purification**

364 Purification for full-length human dynein was previous described¹⁹. Briefly, the cell pellets 365 from a 1.4 L cell culture were resuspended in 100 mL lysis buffer (50 mM HEPES pH 7.2, 366 100 mM NaCl, 1 mM DTT, 0.1 mM ATP, 10% glycerol) containing 2 tablets of Complete 367 EDTA-free protease inhibitor (Roche) and 2 mM PMSF. The suspension was 368 homogenized using a Dounce with a tight plunger for 15~25 strokes, followed by 369 clarification through centrifugation at 65,000 rpm with a Ti70 rotor (Beckman) for 1 hour 370 at 4°C. The supernatant was then incubated with 3 mL IgG Sepharose 6 fast flow resin 371 (Cytiva) for 3~4 hours on a roller at 4°C, followed by washed with 200 mL lysis buffer and 372 200 mL TEV buffer (50 mM Tris-HCl pH 7.4, 150 mM K-acetate, 2 mM Mg-acetate, 1 mM 373 EGTA, 10% glycerol, 0.1 mM ATP, 1 mM DTT). Afterward, the resins were incubated with 374 TEV buffer supplemented with 400 ug TEV protease overnight at 4°C. The supernatant 375 was collected and concentrated with a 100 kDa MWCO Amicon concentrator, then loaded 376 into a TSKgel G4000 column pre-equilibrated with the GF150 buffer (25 mM HEPES pH 377 7.2, 150 mM KCl, 1 mM MgCl₂, 5 mM DTT, 0.1 mM ATP). Peak fractions were collected

and concentrated to 2~3 mg/mL for Cryo-EM grid preparation. The quality of the sample
 was evaluated with the SDS-PAGE gels and the negative-stain EM.

The purification of Lis1, Nde1, and BicDR1 from a 0.7 L cell culture followed a similar protocol to that of dynein, with a few modifications. Specifically, 50 mL of lysis buffer was used to resuspend the cell pellets, and ATP was omitted from the GF150 buffer. And Superose 6 Column (Cytiva) was used for size exclusion chromatography. The concentrated proteins were aliquoted, flash-frozen in liquid nitrogen, and stored at -80°C. The quality of the proteins was assessed using SDS-PAGE gels.

386 Dynactin was isolated from pig brains through a series of purification steps, including SP 387 Sepharose Fast Flow and MonoQ ion exchange chromatography (Cytiva), followed by 388 size exclusion chromatography using a TSKgel G4000SWXL column (Tosoh), as 389 described by previous protocol⁶⁴.

390 MT reconstitution

391 MTs were reconstituted using porcine tubulin, which was either purchased from 392 Cytoskeleton or purified in-house in MT buffer (25 mM MES, 70 mM NaCl, 1 mM MgCl2, 393 1 mM EGTA, and 1 mM DTT, pH 6.5). The tubulin was concentrated to 10 mg/mL at 4°C, 394 then flash-frozen and stored at -80°C. To polymerize the MTs, the tubulin was diluted to 395 5 mg/mL in MT buffer supplemented with 3 mM GTP. The tubulin mixture was incubated 396 on ice for 5 minutes and then transferred to a 37°C incubator for 1 hour. After incubation, 397 the MTs were pelleted at 20,000 rcf for 8 minutes at room temperature and resuspended 398 in MT buffer supplemented with 5 µM paclitaxel before being stored at room temperature.

399 MP assay

400 High-precision coverslips (Azer Scientific) were cleaned by alternating washes with 401 isopropanol and water three times in a bath sonicator, then air-dried. The gasket was 402 cleaned similarly, without sonication, and air-dried before being placed onto a clean 403 coverslip. A total of 14 µL of filtered mass photometry buffer (30 mM HEPES pH 7.4, 5 404 mM MgSO₄, 1 mM EGTA, and 10% glycerol) was added to a well for autofocus. The 405 protein sample was then applied to the well and diluted to a concentration of 5-20 nM in 406 the buffer. Protein contrast data were collected using a TwoMP mass photometer (Refeyn 407 2) with two technical replicates. The instrument was calibrated with a standard mix of 408 conalbumin, aldolase, and thyroglobulin. MP profiles were analyzed by fitting to multiple 409 Gaussian peaks, with the mean, standard deviation, and percentages calculated using 410 DiscoverMP software (Refeyn). The data for parameters of a multi-Gaussian fit of MP 411 measurements is summarized in Supplementary Table 2.

412 **Negative-stain EM and data quantification**

413 Freshly purified dynein was diluted in GF150 buffer to a final concentration of 14.3 nM in 414 the presence of 0.1 mM ATP and subsequently evaluated using negative-stain electron 415 microscopy (EM). A 4 µL aliquot of the sample was applied to glow-discharged carbon 416 film grids (Electron Microscopy Sciences) and stained with 2% uranyl acetate. The grids 417 were then imaged using a 120 kV Talos L120C electron microscope. Micrographs were 418 manually acquired at a magnification of 45,000x. More than 40 micrographs were 419 collected per experiment for each condition to obtain sufficient particles for statistical 420 analysis.

- 421 Samples for statistical analysis of the Phi ratio were prepared with the following molar 422 ratios: dynein: Lis1(dimer) at 1:0, 1:1, and 1:2; dynein: Nde1(dimer) at 1:0, 1:1 and 1:2; 423 dynein : Lis1 : Nde1 at 1:0:0, 1:1:1 and 1:2:2. The mixtures were incubated on ice for 90 424 minutes and subsequently subjected to negative staining. Each experimental group was 425 accompanied by its respective control. The assay was repeated independently three or 426 more times, using different batches of freshly purified protein.
- 427 Micrographs were processed using cryoSPARC, including blob picking, micrograph 428 extraction, and 2D classification. Briefly, micrographs from each experimental group were 429 merged, and particle picking was performed using templates of Phi and open dynein. The 430 particle diameter was set to 750 Å, and the distance cutoff for dynein particles was 400 Å 431 to optimize particle selection. All particles were extracted, followed by three rounds of 2D 432 classification. Phi and open dynein particles were identified from the 2D classification and 433 traced back to the corresponding micrographs for each condition, where Phi and open 434 particles were quantified. To calculate the normalized fraction of Phi, the total number of 435 Phi and paired open dynein particles was determined for micrographs under a given 436 condition. The ratio of Phi particles in the dynein-alone condition (N(Phi) / N(Phi + open)) 437 was defined as proportion A, serving as the control for each group. The ratio of Phi 438 particles in each experimental condition, excluding the dynein-alone control, was defined 439 as proportion B (N(Phi) / N(Phi + open)). The normalized fraction of Phi was calculated 440 as B/A, and GraphPad Prism was used to plot the normalized fraction of Phi.

441 Cryo-EM sample preparation

For the dynein, Lis1, and Nde1 sample, Lis1 and Nde1 were incubated at a 1:1 molar ratio for 30 minutes on ice. Freshly purified dynein, at a concentration of 2 mg/mL, was then added to the Lis1-Nde1 complex at a 1:2:2 molar ratio and incubated on ice for 5 minutes, with 5 mM ATP added immediately prior to freezing. For the dynein and Lis1 complex, dynein (2 mg/mL) was incubated with Lis1 for 90 minutes on ice, and 5 mM ATP was added just before vitrification.

For vitrification, 3.5 µL of the prepared sample was applied to glow-discharged Quantifoil
holey carbon grids (R2/1, 300 mesh gold), which were treated for 45 seconds at 25 mA
using a GloQube Glow Discharge system (Quorum Technologies). The grids were blotted

451 for 2.5 to 4.5 seconds at 4°C and 100% humidity, then vitrified by plunging into liquid 452 ethane using a Vitrobot Mark IV (Thermo Fisher Scientific).

453 Cryo-EM data collection

454 Data were collected at the Yale ScienceHill Cryo-EM facility using a Glacios microscope 455 (Thermo Fisher Scientific) operated at 300 keV and equipped with a K3 detector. Data 456 collection was facilitated by SerialEM software, targeting a defocus range of -1.2 μ m to -457 2.6 μ m. Four exposures per hole were recorded as movies, comprising 40 frames each, 458 with a total electron dose of 40 e⁻/Å². A total of 7,128 movies were collected for the 459 Dynein-Lis1-Nde1 condition, while 16,558 movies were collected for the Dynein-Lis1 460 condition.

461 Cryo-EM data processing

462 Cryo-EM movies were pre-processed using CryoSPARC Live, which included patch
463 motion correction and patch CTF estimation. The processing workflows are illustrated in
464 Extended Data Fig. 4 and 5. The statistics are summarized in Table 1.

465 For the dynein-Lis1 condition dataset, particles were picked using the blob picker, 466 extracted with a box size of 512 pixels, and downscaled to 128 pixels with a pixel size of 467 3.456 Å. In total, 7,254,515 particles were extracted. The iterative 2D classification was 468 performed to filter the particles, resulting in the selection of 204,005 high-quality particles 469 for ab initio reconstruction. Initial maps for the dynein single motor domain and the Phi^L-470 Lis1 motor domains were identified. The map of the Phi^L-Lis1 motor domains was 471 subsequently used for heterogeneous refinement of all original particles. The original 472 particles were divided into four subsets, each subjected to heterogeneous refinement (4 473 classes). Three rounds of heterogeneous refinement were performed, updating the 474 reference each time, ultimately identifying the Phi and Phi^L-Lis1 motor domains. The Phi 475 and Phi^L-Lis1 motor domains were merged separately and extracted from micrographs 476 using a box size of 512 pixels, which was then binned to 384 pixels, resulting in a pixel 477 size of 1.1573 Å. Two rounds of heterogeneous refinement were conducted to exclude junk particles. High-quality subsets were selected for homogeneous refinement, followed 478 479 by two rounds of CTF refinement and local refinement. The Phi^L-Lis1 motor domains 480 achieved a resolution of 2.86 Å, and the Phi motor domains reached a resolution of 2.71 481 Å, exhibiting C2 symmetry.

To reconstitute the tails of the Phi^L-Lis1 and Phi motor domains, the particles were recentered at the tail and then extracted from micrographs using a box size of 512 pixels, binned to 256 pixels, yielding a pixel size of 1.736 Å/px. Following this, heterogeneous refinement was applied to filter the particles, and high-quality subsets were selected for homogeneous refinement. The overall tail resolutions reached 4.21 Å for the Phi tail and 4.05 Å for the Phi^L-Lis1 tail. Four masks were devised to cover the tail segments, which were divided into the NDD, left, right, and neck regions. Maps of local refinement using
these masks were integrated, referring to the consensus map of the tail. The full-length
map was assembled by stitching together the tail and motor domains in ChimeraX,
corresponding to Phi and Phi^L-Lis1, respectively.

492 For the Lis1-Nde1-dynein condition dataset, the process closely mirrored that of the 493 dynein-Lis1 condition described above. Briefly, the blob picker identified 2,393,279 494 particles. Following extraction from the micrographs and iterative 2D classification, 495 204,005 particles were selected for initial map generation through ab initio reconstruction. 496 An initial map for the Phi^L-Lis1 motor domains were obtained, which were then subjected 497 to iterative heterogeneous refinement using all particles with the Phi^L-Lis1 initial map as 498 a reference. However, the Phi motor domains did not appear in the heterogeneous 499 refinement, even when the verified Phi map from this study was utilized as a reference. 500 Subsequently, the Phi^L-Lis1 domain map was re-extracted from the micrographs and 501 underwent two rounds each of heterogeneous refinement, CTF refinement, and local 502 refinement, achieving a resolution of 2.88 Å. The consensus map of the tail reached a 503 resolution of 6.22 Å after recentering and re-extracting the tail region. Masks and local 504 refinement were employed to enhance the local resolution of the tail. The full-length Phi^L-505 Lis1 structure was reconstituted by integrating the composite tail map and motor map in 506 ChimeraX.

507 Model building and refinement

For model building, previously reported structures 9BLY¹⁹, 9BLZ¹⁹, and 8FDT³⁸ were 508 509 utilized as the initial models for the full-length and motor domains of Phi and Phi^L-Lis1. 510 The individual domains, including the tail, single motor and Lis1 dimer, were extracted 511 from the 9BLY, 9BLZ, and 8FDT, and rigid-body fitting into the Cryo-EM maps were 512 performed using UCSF ChimeraX. The models were then manually constructed in 513 $COOT^{65,66}$, and followed by real-space refinement in Phenix⁶⁷. The quality of the refined 514 models was assessed using the MolProbity integrated into Phenix, with the statistics 515 reported in Table 1.

516 Single-molecular motility assay

517 Fluorescent imaging was conducted using a custom-built, multicolor objective-type TIRF 518 microscope based on a Nikon Ti-E microscope body. It was equipped with a 100X 519 magnification, 1.49 N.A. apochromatic oil-immersion objective (Nikon) and a Perfect 520 Focus System. Fluorescence signals were captured by an electron-multiplied charge-521 coupled device camera (Andor, Ixon EM+, 512 × 512 pixels), with an effective pixel size 522 of 160 nm after magnification. Probes such as Alexa488/GFP/mNeonGreen, LD555, and 523 LD655 were excited by 488 nm, 561 nm, and 633 nm laser beams (Coherent), coupled 524 to a single-mode fiber (Oz Optics), and their emissions were filtered using 525/40, 585/40,

and 697/75 bandpass filters (Semrock), respectively. The entire system was controlledvia MicroManager 1.4 software.

527 Biotin-PEG treated flow chambers were treated with 5 mg/ml streptavidin for 2 minutes, 528 followed by washing with MB buffer (30 mM HEPES pH 7.0, 5 mM MgSO₄, 1 mM EGTA, 529 1 mg/ml casein, 0.5% pluronic acid, 0.5 mM DTT, and 1 µM Taxol). Biotinylated MTs were 530 then added to the chamber for 2 minutes and washed again with MB buffer. Proteins were 531 prepared by diluting them to the desired concentrations in MB buffer. For DDRNL complex 532 assembly, a mixture of 10 nM dynein, 150 nM dynactin, 50 nM BicDR1, 200nM Lis1 and 533 10nM Nde1 was incubated on ice for 15 minutes, then diluted tenfold into imaging buffer 534 (MB buffer containing 0.1 mg/ml glucose oxidase, 0.02 mg/ml catalase, 0.8% D-glucose, 535 and 2 mM ATP) and introduced to the flow chamber. Motility was observed and recorded

536 for 40 seconds.

537 Data analysis for single-molecular motility assay

538 Single-molecule motility of the DDR complex was captured for 200 frames per imaging

area and analyzed as kymographs made in FIJI. Run frequency was determined by

540 counting the number of processive BicDR1 molecules on each MT, then dividing this

number by the MT length and the total data collection time, with a custom MATLAB script.

542 The p-values for the two-tailed Student's t-test were determined in Excel.

543 Data availability

544 Cryo-EM Density maps and models have been deposited in the Electron Microscopy Data 545 Bank and Protein Data Bank as follows: In the dynein and Lis1 condition: PDB-546 9E12/EMD-47381 for full length Phi and PDB-9E10/EMD-47379 for motor domains of Phi; 547 PDB-9E13/EMD-47382 for full length Phi^L-Lis1 and PDB-9E11/EMD-47380 for motor 548 domains of Phi^L-Lis1. In the dynein, Lis1 and Nde1 condition: PDB-9E14/EMD-47383 for 549 full-length Phi^L-Lis1 and 9E0Z/EMD-47378 for motor domains of Phi^L-Lis1.

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559 Author contributions

K.Z. and A.Z. designed the study. J.Y. expressed and purified dynein, Lis1 and Nde1
proteins for EM. J.Y. and P.C. prepared the cryo-EM samples, collected and processed
the data, and built the PDB models. P.C. and J.Y. processed negative stain EM data and
quantified the particle numbers. Y.Z. performed Lis1 mutagenesis, protein preparation,
TIRF and mass photometry assays. J.Y., P.C., Y.Z., K.Z., and A.Y. analyzed the data and
prepared figures. J.Y., Y.Z., P.C., K.Z., and A.Y. wrote the manuscript with input from all
authors.

567 Competing Interests

568 The authors declare no competing interests.

569 Figures



Fig. 1. Nde1 promotes Lis1 binding to Phi dynein and cooperatively releases dynein
autoinhibition. a, Schematic of dynein activation. Nde1 tethers Lis1 to dynein and
promotes Lis1-mediated formation of the active dynein-dynactin-adaptor-Lis1 (DDAL)
complex. Intermediate states between the association of Lis1-Nde1 and opening of
dynein are unknown. b, MP shows that Lis1 alone slowly binds to dynein in tens of

576 minutes, whereas Nde1 promotes more rapid and efficient binding of Lis1 to dynein. 577 Dynein, Lis1 and Nde1 were included at a 1:2:2 ratio. Only one Lis1 dimer is tethered to 578 one dynein. Solid curves represent a fit to multiple Gaussians to predict the average mass 579 and percentage of each population. c, A representative image of dynein motors captured 580 using negative stain electron microscopy. Without Lis1 and Nde1, dynein is distributed 581 almost equally between Phi and open conformations. The percentage of Phi was 582 quantified after incubating dynein with Lis1, Nde1, or both proteins for 90 min. The 583 percentage of Phi (d) and the relative change of Phi (e) in the presence and absence of Lis1 and Nde1 (mean ± s.d.; from left to right, n=14, 6, 6, 3, 4, 4, 4 from three or more 584 independent experiments). P values are calculated from a two-tailed *t* test. The control 585 586 (Ctrl) of panel e represents dynein alone from the three groups of panel d (Ctrl) normalized 587 to 100%.



589 Fig. 2. The structure of Phi^L-Lis1 complex. a, (Top) Full length human dynein was 590 incubated with Lis1 with a molar ratio of 1:2 on ice for 90 min before flash frozen. (Bottom) 591 Cryo-EM maps of Phi and Phi^L-Lis1 complex structures in front view and percentages of particles with these two conformations (open dynein excluded). b, (Top) Dynein was 592 593 incubated with Lis1 and Nde1 with a molar ratio of 1:2:2 on ice for 5 min before flash 594 frozen. (Bottom) The structure of Phi^L-Lis1 obtained under this condition is shown in front 595 view and side view. All particles were classified into the Phi^L-Lis1 conformation and Phi 596 conformation could not be detected. c, Model of the Phi^L-Lis1 motor domains is shown in 597 three views. One Lis1 dimer is clamped between MD-A and MD-B of Phi^L-Lis1 on the 598 front side, while the back side of the motor domains is vacant, with no Lis1 bound. The 599 top view reveals that the linker regions (purple) of MD-A and -B interact with each other.

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600

Fig. 3. Lis1 induces a transition in the Phi conformation to accommodate its binding. 601 602 a, Docking of a Lis1 monomer between dynein motor rings shows a clash with the rigid 603 canonical Phi MD-A. b, A Lis1 dimer bound to dynein motors on the front side is compatible with the coordinated Phi^{L} structure. **c**. (Top) The superimposition of the Phi 604 605 and Phi^L-Lis1 motor domains. Phi and Phi^L-Lis1 are colored with grey and rainbow 606 respectively. (Bottom) Comparison of the motor domains between Phi^L-Lis1 and 607 canonical Phi. Lis1 is hidden for clarity. Vectors present interatomic distance of pairwise 608 $C\alpha$ atoms between the Phi to Phi^L-Lis1 structures. Lis1 binding induces a slight rotation 609 of the dynein motor domains from the Lis1-bound (front) side toward the back, resulting 610 in a larger groove at the front side compared to the smaller groove at the back. d, Overlay 611 of the motor domain maps of Phi and Phi^L-Lis1 with MD-A aligned at a lower contour level. 612 The rotation of Phi^L-Lis1 motor domain induces rotation in the neck region, showing a 613 slight unwinding trend, which contributes to the formation of a smaller groove on the back 614 side. e, Docked Lis1 on the back side of Phi^L-Lis1 causes a significant clash with MD-B.





617 Fig. 4. Novel interactions identified between Lis1_{ring} and MD-A in Phi^L-Lis1. a, Cryo-618 EM density map highlighting the subdomains involved in the interface between Lis1 and 619 the dynein motor. Lis1 and subdomains of MD-A and MD-B are colored separately. Novel 620 interfaces between MD-A and Lis1ring are marked by dash rectangle and enlarged in 621 panels d, e, and f. b, Comparison of MD-A (grey) and -B (rainbow) of Phi^L-Lis1. c, 622 Comparison of MD-B (rainbow) with the Lis1-bound structure of the motor domain of 623 human dynein (PDB: 8FDT, grey). Representative interactions located at linker-Lis1ring 624 (d), AAA6-Lis1_{ring} (e), and AAA5-Lis1_{ring} (f) interface are shown with stick mode, colored

according to their respective subdomains. g, MP profiles illustrate the interaction of Nde1 625 (N) with WT Lis1 (L) and Lis1 mutants (Lis1^{AAA5}, Lis1^{AAA6}, and Lis1^{linker}). Nde1 interacts 626 with one (NL) or two (NL₂) Lis1 dimers. Lis1^{AAA5} exhibits a reduced binding percentage to 627 Nde1. h. MP shows the binding of Lis1 mutants to dynein with or without Nde1. Dynein, 628 629 Lis1 and Nde1 were incubated for 2 minutes at 1:2:2 ratio (D: dynein only, DL: one dynein 630 and one Lis1, DL₂N: one dynein, two Lis1, and one Nde1). i, Representative kymographs 631 show the motility of WT DDR complexes with or without Nde1 and Lis1. **i**, Run frequency 632 of WT DDR with or without Nde1 and Lis1 (mean \pm s.d.; n = 20 MTs for each condition; 633 statistics from two independent experiments). Results were normalized to the -Lis1, -634 Nde1 condition.



636

637 Fig. 5. Model for the initial stage of dynein activation by Nde1 and Lis1. Step-i, Nde1 638 tethers Lis1 to Phi dynein and forms a transient Phi^L-Lis1-Nde1 complex. Step-ii, Nde1 639 dissociates spontaneously from dynein, leading to the formation of Phi^L-Lis1. Without 640 Nde1, Lis1 binding to Phi dynein and formation of Phi^L-Lis1 becomes a less efficient and 641 slow process. The binding of Lis1 induces a slight backward rotation of the two motor 642 rings and a twist in the neck region, likely leading an unwinding trend in the tail and 643 contributing to dynein opening. Step-iii, Phi^L-Lis1 transitions to open dynein-Lis1 assisted 644 by Nde1. Step-iv, Lis1 promotes processive DDA complex assembly by interacting with 645 p150 and DIC-N through its LisH domain. The dashed line indicates the possibility that Lis1 in the Phi^L-Lis1 complex promotes dynactin recruitment and DDA assembly without 646 647 switching to the open conformation.

In dynein and Lis1 condition				In Nde1, Lis1 and dynein condition				
Description	Composite map of full-length Phi	Composite map of full-length Phi ^L -Lis1	Phi motor	Phi ^L -Lis1 motor	Composite map of full-length Phi ^L -Lis1	Phi ^L -Lis1 motor		
PDB code	9E12	9E13	9E10	9E11	9E14	9E0Z		
Data Collection and Processing								
Facility		Yale ScienceHill-Cr	o-EM facility		Yale ScienceHill-Cryo-EM facility			
Microscope		Glacios			Glacios			
Voltage (kV)		200			200			
Camera	КЗ				Ka	КЗ		
Magnification		45k			45k			
Pixel Size (Å)	0.432 (super resolution)				0.432 (super resolution)			
Total Electron Exposure (e-/A ²)		40			40			
Defocus Range (µm)	1.5-2.7			1.5-2.7				
Symmetry Imposed	C1	C1	C2	C1	C1	C1		
Num of mics		16,558			7,128			
Initial Particles		7,254,51	5		2,393,279			
Final Particles	103,097	127,963	160,539	215,049	61,684	127,611		
Refinement								
Initial models		9BLY		0FDT	9BLY	AFRT		
	ab initio	8FDT ab initio	ab initio	ab initio	8FDT ab initio	ab initio		
Map pixel size	1.7360	1.7360	1.1573	1.1573	1.736	1.157		
Map Resolution (Å) (FSC 0.143)	4.50	4.50	2.71	2.86	5.0	2.88		
Map sharpening B-factor (Å ²)	/	/	72.3	69.6	/	51.7		
Model Composition								
Non-hydrogen atoms	89,388	94,497	47418	21,370	94,497	52,508		
Protein residues	11,061	11,704	5784	6514	11,704	6,515		
Ligands	MG:4	MG:4	MG:4	MG:4	MG:4	MG:4		
	ATP:2	ATP:2 ADP:6	ATP:2 ADP:6	ATP:2	ATP:2	ATP:2 ADP:6		
Model vs. Data		712110	7,5110	7.51.10		7.01.10		
FSC Map to Model (Å) (FSC 0.5)	3.5	3.8	3.0	3.1	3.5	3.4		
Correlation coefficient (mask)	0.60	0.59	0.86	0.87	0.51	0.84		
B factors (A ²)								
Protein	117.05	112.65	14.13	38.69	260.48	52.33		
Ligand	75.90	79.64	1.76	14.44	69.27	32.47		
R.m.s deviation				1				
Bond length (Å)	0.006	0.004	0.003	0.003	0.002	0.002		
Bond angles (°)	0.683	0.693	0.548	0.527	0.544	0.548		
Validation				1				
Molprobity score	1.87	1.88	1.40	1.41	1.75	1.45		
Clashscore	13.26	13.07	7.31	7.33	10.62	8.06		
Rotamer outliers (%)	0.00	0.00	0.00	0.00	0.00	0.02		
Ramachandran plot								
Outliers (%)	0.09	0.07	0.00	0.00	0.05	0.00		
Allowed (%)	3.52	3.74	1.50	2.05	3.23	2.05		
Favored (%)	96.39	96.20	98.50	97.95	96.72	97.95		
Rama-Z (whole)	0.62	0.53	1.60	1.72	0.92	1.86		

648 Table 1. Cryo-EM data collection, refinement, and validation statistics.

Construct	Vector	Source	Figures	
SNAPf-DYNH1C1-	pOmniBac	Schlager et	F1, 2, 3, 4,	
IC2C-LIC2-Robl1-	-pIDC	al., 2014	EDF2, 3, 4, 5, 6, 8, 9, 12	
Tctex1-LC8				
SNAPf-	pOmniBac	Zhang et al.,	EDF1, 7, 13	
DYNH1C1 ^{R1567E, K1610E} -	-pIDC	2017		
IC2CLIC2-Robl1-				
Tctex1-LC8				
BicDR1-SNAPf	pOmniBac	Urnavicius et	F4,	
		al. 2018	EDF13	
Lis1-SNAPf	pOmniBac	Elshenawy et	F1, 2, 3, 4,	
		al., 2020	EDF1, 2, 3, 4, 5, 6, 7, 8, 9, 11,	
			12, 13	
Lis1 ^{E300K, K303E, S304R,}	pOmniBac	This study	F4,	
M329A-SNAPf			EDF11	
Lis1 ^{N203K, D205K, Q222A} -	pOmniBac	This study	F4,	
SNAPf			EDF11	
Lis1 ^{M172K, D192K, Y225A,}	pOmniBac	This study	F4,	
C226D-SNAPf			EDF11	
Nde1 ¹⁻¹⁹⁰ -SNAPf	pOmniBac	Zhao et al.,	F1, 4,	
		2023	EDF1, 2, 7, 11, 13	

649

Supplementary Table 1. The list of protein constructs used in this study. Dynein chains were codon-optimized for *Spodoptera frugiperda* (Sf9) expression and inserted into the pOmniBac backbone. Nde1, Lis1, and BicDR1 constructs were cloned into the pOmniBac backbone. Constructs were tagged with an N-terminal 6xHis-ZZ-TEV site for affinity purification and TEV protease cleavage during protein purification. The SNAPf tag was inserted for labeling the proteins with fluorescent dyes (F: Figure, EDF: Extended Data Figure).

Fig.	Sample	Complex	Expected	Measured	%
	-	-	(kDa)	(kDa)	
F 1b	WT Dyn	WT Dyn	1376	1326 ± 49	100*
	WT Dyn + Lis1 0 min	WT Dyn	1376	1350 ± 65	100*
	WT Dyn + Lis1 10 min	WT Dyn	1376	1344 ± 55	80*
		WT Dyn + Lis1	1546	1521 ± 129	20*
	WT Dyn + Lis1 30 min	WT Dyn	1376	1364 ± 60	65*
		WT Dyn + Lis1	1546	1544 ± 68	35*
	WT Dyn + Lis1 60 min	WT Dyn	1376	1359 ± 48	63 [*]
		WT Dyn + Lis1	1546	1528 ± 40	37*
	WT Dyn + Lis1 + Nde1	WT Dyn	1376	1356 ± 64	31*
	0 min	WT Dyn + Lis1	1546	1525 ± 70	69 [*]
	WT Dyn + Nde1	WT Dyn	1376	1330 ± 83	100*
F 4g	WT Lis1	WT Lis1	133	137 ± 17	70
	Lis1 ^{AAA5}	Lis1 ^{AAA5}	133	141 ± 26	87
	Lis1 ^{AAA6}	Lis1 ^{AAA6}	133	141 ± 26	87
	Lis1 ^{Linker}	Lis1 ^{Linker}	133	142 ± 31	90
	Nde1 ¹⁻¹⁹⁰ + WT Lis1	Nde1 ¹⁻¹⁹⁰	84	85 ± 18	31
		WT Lis1	133	142 ± 15	29
		1 Nde1 ¹⁻¹⁹⁰ + 1 WT Lis1	217	231 ± 20	28
		1 Nde1 ¹⁻¹⁹⁰ + 2 WT Lis1	340	371 ± 44	12
	Nde1 ¹⁻¹⁹⁰ + Lis1 ^{AAA5}	Nde1 ¹⁻¹⁹⁰	84	90 ± 17	37
		Lis1 ^{AAA5}	133	142 ± 13	53
		1 Nde1 ¹⁻¹⁹⁰ + 1 Lis1 ^{AAA5}	217	229 ± 48	10
	Nde1 ¹⁻¹⁹⁰ + Lis1 AAA6	Nde1 ¹⁻¹⁹⁰	84	82 ± 17	25
		Lis1 ^{AAA6}	133	141 ± 15	30
		1 Nde1 ¹⁻¹⁹⁰ + 1 Lis1 ^{AAA6}	217	233 ± 30	29
		1 Nde1 ¹⁻¹⁹⁰ + 2 Lis1 ^{AAA6}	340	373 ± 16	9
	Nde1 ¹⁻¹⁹⁰ + Lis1 ^{Linker}	Nde1 ¹⁻¹⁹⁰	84	88 ± 16	24
		Lis1 ^{Linker}	133	143 ± 14	26
		1 Nde1 ¹⁻¹⁹⁰ + 1 Lis1 ^{Linker}	217	229 ± 18	30
		1 Nde1 ¹⁻¹⁹⁰ + 2 Lis1 ^{Linker}	340	372 ± 15	12
F 4h	WT Dyn	WT Dyn	1376	1367 ± 93	100*
	WT Dyn + WT Lis1	WT Dyn	1376	1387 ± 52	82 [*]
		WT Dyn + WT Lis1	1509	1530 ± 39	18*
	WT Dyn + WT Lis1 +	WT Dyn + WT Lis1	1509	1514 ± 67	100*
	Nde1 ¹⁻¹⁹⁰				
	WT Dyn + Lis1 ^{AAA5}	WT Dyn	1376	1367 ± 94	100*
	WT Dyn + Lis1 ^{AAA5} +	WT Dyn	1376	1361 ± 57	57*
	Nde1 ¹⁻¹⁹⁰	WT Dyn + Lis1 ^{AAA5}	1509	1581 ± 76	43*
	WT Dyn + Lis1 AAA6	WT Dyn	1376	1356 ± 60	100*
		WT Dyn	1376	1356 ± 75	77*

	WT Dyn + Lis1 AAA6 +	WT Dyn+2*Lis1 ^{AAA6} +	1716	1680 ± 77	23*
	Nde1 ¹⁻¹⁹⁰	Nde1 ¹⁻¹⁹⁰			
	WT Dyn + Lis1 ^{Linker}	WT Dyn	1376	1349 ± 56	100*
	WT Dyn + Lis1 ^{Linker} +	WT Dyn + Lis1 ^{Linker}	1509	1464 ± 75	100*
	Nde1 ¹⁻¹⁹⁰				
EDF	mtDyn	mtDyn	1376	1362 ± 46	100*
1	mtDyn + Lis1 + Nde1	mtDyn	1376	1386 ± 77	32*
		mtDyn + Lis1	1546	1525 ± 74	68 [*]
	mtDyn + Lis1	mtDyn	1376	1411 ± 73	50 [*]
		mtDyn + Lis1	1546	1540 ± 57	50 [*]
EDF	WT Dyn	WT Dyn	1376	1396 ± 241	100*
2a	WT Dyn + Lis1	WT Dyn	1376	1414 ± 85	100*
	WT Dyn + Nde1	WT Dyn	1376	1371 ± 68	100*
	WT Dyn + Lis1 + Nde1	WT Dyn	1376	1415 ± 85	30*
		WT Dyn + Lis1	1546	1580 ± 63	70*
EDF	WT Dyn	WT Dyn	1376	1347 ± 278	100*
2b	WT Dyn + Lis1	WT Dyn	1376	1395 ± 167	100*
	WT Dyn + Nde1	WT Dyn	1376	1376 ± 78	100*
	WT Dyn + Lis1 + Nde1	WT Dyn	1376	1391 ± 65	48*
		WT Dyn + Lis1	1546	1585 ± 133	52 [*]
EDF	WT Dyn	WT Dyn	1376	1381 ± 61	100*
2c	WT Dyn + Lis1	WT Dyn	1376	1371 ± 136	100*
	WT Dyn + Nde1	WT Dyn	1376	1377 ± 113	100*
	WT Dyn + Lis1 + Nde1	WT Dyn	1376	1409 ± 75	45*
		WT Dyn + Lis1	1546	1584 ± 47	55*
EDF	WT Dyn	WT Dyn	1376	1411 ± 108	100*
2d	WT Dyn + Lis1	WT Dyn	1376	1377 ± 124	100*
	WT Dyn + Nde1	WT Dyn	1376	1366 ± 130	100*
	WT Dyn + Lis1 + Nde1	WT Dyn	1376	1383 ± 56	28*
		WT Dyn + Lis1	1546	1561 ± 51	72*
EDF	WT Dyn	WT Dyn	1376	1396 ± 304	100*
2e	WT Dyn + Lis1	WT Dyn	1376	1403 ± 76	100*
	WT Dyn + Nde1	WT Dyn	1376	1379 ± 54	100*
	WT Dyn + Lis1 + Nde1	WT Dyn	1376	1402 ± 53	27*
		WT Dyn + Lis1	1546	1581 ± 46	73*

657 Supplementary Table 2. The parameters of a multi-Gaussian fit of MP 658 measurements. Dynein (Dyn), Lis1, and Nde1 were mixed at 1:2:2 ratio if present and 659 diluted into 5-20 nM. Measured mass and percentage represent the center (mean \pm s.d.) 660 and the percent area of the corresponding Gaussian peak (*percentages are normalized 661 so that the sum of all dynein included peaks are 100%). Expected mass corresponds to 662 the dimeric forms of Lis1, Nde1, and Dyn (F: Figure, EDF: Extended Data Figure). Lis1 663 used in F 1b, and EDF 1 contains a ZZ tag, and its expected MW is 170 kDa.

Supplementary Video Legends 664

- Supplementary Video 1. Full-length human dynein in Phi^L conformation, bound to a Lis1 665 dimer and displaying the newly identified interface with Lis1. 666
- 667 Supplementary Video 2. Single-molecule motility recordings of WT DDR complexes, in
- the presence or absence of Nde1, WT Lis1, and Lis1 mutants. The fluorescence signal 668
- originates from BicDR1-mNeonGreen. 669

670 Extended Data Figures



671

672 Extended Data Fig. 1. MP analysis of Nde1's effect on Lis1 binding to open dynein.

673 MP shows that Nde1 does not promote increased Lis1 binding to open dynein compared

to the Lis1-alone condition, indicating that Lis1 can efficiently bind to open dynein and

675 Nde1 does not enhance Lis1's interaction with open dynein.

+WT Lis1

+WT Nde1

+WT Lis1

+WT Nde1

2.0







679 Extended Data Fig. 2. MP analysis of nucleotide conditions on Nde1 and Lis1 binding to WT dynein. MP shows that under apo buffer (a), 0.1 mM ATP (b), ADP (c), 680 ATP.vi (d), and AMPPNP (e) conditions, Nde1 promotes Lis1 binding to dynein, forming 681 682 a 1:1 dynein-Lis1 (DL) complex. The nucleotide condition does not affect Nde1's ability to 683 tether Lis1 to dynein. Importantly, the formation of dynein-Nde1, dynein-Lis1-Nde1 and

- 684 1:2 dynein-Lis1 complexes were not observed. In the Lis1-alone condition, no significant
- 685 DL complex was formed immediately.





đ 0 0 % 8-0 8-ct 8 Ū, 1 00 . .9 0 • 5 à ø 6 3rd (final) round

Count particle numbers from each group (Single motor is considered as open dynein with particle number divided by 2)

687 Extended Data Fig. 3. Workflow for negative-stain EM data processing. a, 688 Representative micrographs for dynein alone (42 micrographs), dynein-Lis1 at 1:1 (41 689 micrographs), and dynein-Lis1 at 1:2 (47 micrographs) molar ratios from batch #1. b, 690 Particle picking from representative micrographs in each dataset using a template 691 matching approach based on Phi and open dynein models (particle diameter: 750 Å, 692 distance cutoff: 400 Å). c, Three rounds of 2D classification were performed after extracting all particles (box size: 960 Å × 960 Å), yielding class averages of Phi dynein, 693 open dynein motors, single motors, and junk particles. d, Final classified 2D averages 694 showing Phi dynein, two-motor open dynein, and single-motor open dynein. The particle 695 696 numbers for each group were counted, and single motors were considered as open 697 dynein by dividing the total number of particles by two.



699 Extended Data Fig. 4. Cryo-EM data processing for the dynein-Lis1 dataset. a, A

representative cryo-EM micrograph and the flowchart of cryo-EM data processing. **b**,

- 701 Fourier Shell Correlation (FSC) curves showing the final resolution estimates for the
- motor domains of the Phi (2.71 Å) and Phi^L-Lis1(2.86 Å) datasets.



704 Extended Data Fig. 5. Cryo-EM data processing for the Nde1-Lis1-dynein dataset.

- **a**, A representative cryo-EM micrograph and the flowchart of cryo-EM data processing. **b**,
- Fourier Shell Correlation (FSC) curve showing the final resolution estimate for the motor
- 707 domains of the Phi^L-Lis1 (2.88 Å) dataset.



710 Extended Data Fig. 6. Comparison of local resolution, nucleotide binding in AAA1,

- 711 AAA3, AAA4, and sensor-I loop conformation in MD-A of the Phi and Phi^L-Lis1.
- Local resolution, and nucleotide binding states in MD-A at AAA1, AAA3, and AAA4 of the
- 713 Phi (a) and Phi^L-Lis1 (b). MD-A and -B share the same nucleotide binding in AAA1, AAA3,
- and AAA4 across both the Phi and Phi^L-Lis1. The sensor-I loop adopts almost the same
- conformation in MD-A (or -B) of both Phi (c) and Phi^L-Lis1 (d), indicating that Lis1 binding
- does not affect phosphate release. The color scheme is the same with Fig. 4.



717

718 Extended Data Fig. 7. Comparison of Nde1's effect on Lis1 binding to the open

719 dynein. The particle numbers for open dynein-Lis1 and open dynein alone were

quantified in both the dynein-Lis1 and dynein-Lis1-Nde1 datasets. These results indicate

that Nde1 does not promote Lis1 binding to the open dynein motor.

> E 1622 E 1622 D 1556 H 1559 B 162 B 163 B 162 B 163 B 165 B 165 B 165 B 165 B 155 B 165 B 155 B 155

b AAA5-Lis1_{ring} AAA6-Lis1_{ring}



722

723 Extended Data Fig. 8. Density quality at the dynein MD-A and Lis1 interface in Phi^L-

Lis1. a, Flexible density at the linker-Lis1_{ring} interface, indicating dynamic interactions in this region. b, Well-defined density at the AAA5-Lis1_{ring} and AAA6-Lis1_{ring} regions, showing compact and stable interactions. The color scheme for the motor domains is consistent with Fig. 4.

a Linker-Lis1



Extended Data Fig. 9. Sequence alignment of Lis1 homologs among multiple species. Sequence alignment of Lis1 proteins among Homo sapiens (Hs Lis1), Xenopus 730 laevis (XI Lis1), Gallus gallus (Gg Lis1), Danio rerio (Dr Lis1), and Saccharomyces 731 cerevisiae (Sc Lis1). The secondary structure elements are placed on the top of the 732 733 alignment. Strictly conserved residues are highlighted in shaded red boxes, while less-

conserved residues are shown in open red boxes. In these open red boxes, red font
indicates residues with similar polarity and high conservation, whereas black font
represents residues with low similarity. The colored triangles represent key residues
involved in interactions at the linker-Lis1_{ring} (purple), AAA5-Lis1_{ring} (green), and AAA6Lis1_{ring} (orange) interfaces in Phi^L-Lis1. The black triangle represents reported
interactions at MD-A and Lis1 interface of modeled human Chi-Lis1 based on the yeast

740 Chi-Lis1³⁷. S. cerevisiae Lis1 shows more variation compared to the vertebrate species,

suggesting greater evolutionary divergence.



742

743 Extended Data Fig. 10. Comparison of motor domain A-B interfaces in Phi and Phi^L-

Lis1. a. Representative residues located on MD-A that are involved in the MD-A and MD-

- B interface of Phi, showing interactions at the linker-linker, linker-AAA4, AAA4-linker,
- AAA4-AAA4, AAA5-AAA5, and stalk-stalk interfaces. **b**, Representative residues located

- 747 on MD-A of the Phi^L-Lis1, involved in the MD-A and Lis1_{ring} interface (including linker-
- Lis1_{ring}, AAA6-Lis1_{ring}, and AAA5-Lis1_{ring}), and the motor domain A-B interface (including
- 749 linker-linker, AAA3-linker, linker-AAA4, AAA5-AAA5, and stalk-stalk interfaces). Residues
- in both panels (a) and (b) are displayed in sphere mode. **c**, Detailed view of the motor
- 751 domain A-B interface in Phi^L-Lis1, showing key residues involved in interactions at the
- 752 linker-linker, AAA3-linker, AAA5-AAA5, and linker-AAA4 interfaces. d, Detailed view of
- motor domain A-B interface of Phi, showing key residues at the linker-linker, linker-AAA4,
- AAA4-linker, AAA4-AAA4, and AAA5-AAA5 interfaces.



Find the structure of the Nde1-Lis1 interface predicted by Alphafold. a, Predicted structure of the Nde1-Lis1 complex. Interactions involved in the Phi^L MD-A and Lis1 interface are shown on the Lis1 surface. b, D192 and M172 involved in AAA5-Lis1 interface also show contact with R115 of Nde1. c, D192K and M172K mutation of Lis1^{AAA5} show charge repulsion with R115 of Nde1. The prediction supports an overlap of the interfaces between the AAA5-Lis1 and Nde1-Lis1.



763

Extended Data Fig. 12. Comparison of yeast Chi-Lis1 and human Phi^L-Lis1 motor 764 765 domains. a, The structure of yeast Chi-Lis1 (PDB:8DZZ)³⁷, showing two tail-truncated 766 yeast dynein motor domains (grey) bound to two Lis1 dimers (colored, Chi-Lis1 1:2). b, 767 Residues of MD-A that interact with Lis1ring are located in AAA6-Lis1ring and AAA5-Lis1ring 768 regions and highlighted with dashed rectangle. Representative residues of MD-A involved 769 in the canonical Lis1ring binding sites are located in Lis1ring-AAA3, Lis1ring-AAA4, Lis1ring-AAA5 and Lis1ring-stalk region. Interactions between MD-A and MD-B are in stalk-stalk 770 771 region. Residues are displayed in sphere mode and are colored according to the 772 subdomains in Fig. 4. c, Superimposition of the human Phi^L-Lis1 and yeast Chi-Lis1 773 structures, showing that Chi-Lis1 adopts a more expanded conformation, with larger 774 grooves on both the front and back sides compared to the more compact Phi^L-Lis1

- 575 structure. Lis1 is hidden for clarity. Vectors represent interatomic distances of pairwise
- 776 C α atoms between the Phi^L-Lis1 and Chi-Lis1 structures.

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