# Conformational transitions of the mitotic adaptor Spindly underlie its interaction with Dynein and Dynactin

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Cytoplasmic Dynein 1, or Dynein, is a microtubule minus-end directed motor. Dynein motility requires Dynactin and a family of activating adaptors that stabilize the Dynein-Dynactin complex and promote regulated interactions with cargo in space and time. How activating adaptors limit Dynein activation to specialized subcellular locales is unclear. Here, we reveal that Spindly, a mitotic Dynein adaptor at the kinetochore corona, exists natively in a closed conformation that occludes binding of Dynein-Dynactin to its CC1 box and Spindly motif. A structure-based analysis identified various mutations promoting an open conformation of Spindly that binds Dynein-Dynactin. A region of Spindly downstream from the Spindly motif and not required for cargo binding faces the CC1 box and stabilizes the intramolecular closed conformation. This region is also required for robust kinetochore localization of Spindly, suggesting that kinetochores promote Spindly activation to recruit Dynein. This mechanism may be paradigmatic for Dynein activation by other adaptors at various cellular locales.

#### Introduction

Eukaryotic cells maintain their internal order through the concerted action of a variety of functionally diverse energy-harnessing enzymes. Among these are molecular motors that convert the chemical energy of adenosine 5'-triphosphate (ATP) into mechanical work to dispatch various cargoes to different subcellular locations (Klinman and Holzbaur, 2018). Many molecular motors move along microtubules, polarized cellular tracks with plus- and minus-ends, the latter normally localized near microtubule-organizing centers, such as centrosomes. Motors use microtubules to transport cargoes of various size, ranging from individual protein complexes, to viruses, to organelles. Molecular motors also move chromosomes and transport, crosslink, and reciprocally slide microtubules to promote the assembly of the mitotic spindle in mitotic cells (Pavin and Tolic, 2021). The two main classes of intra-cellular molecular motors are the kinesins and Dynein. Kinesins populate a wide and diverse family of motors, prevalently with plus-end-directed polarity, with each family member specializing in the transport of distinct cargoes (Klinman and Holzbaur, 2018). Cytoplasmic Dynein-1 (Dynein), on the other hand, is a minus-end-directed multi-subunit assembly whose motor subunit, Dynein heavy chain (DHC), is encoded by a single gene. Its association with different cargoes relies therefore on various activating adaptors, each with a distinct cargo preference (Canty et al., 2021; Reck-Peterson et al., 2018; Roberts et al., 2013).

In humans, the 1.4 MDa Dynein complex consists of 12 subunits, with six different polypeptides all present in two copies, including the DHC, the intermediate chains (DIC), the light intermediate

chains (LIC), and the three Dynein light chains LC8, Roadblock, and Tctex (Reck-Peterson et al., 2018). The motility of Dynein requires the 1.1 MDa complex Dynactin (Carter et al., 2016; Reck-Peterson et al., 2018). Dynactin consists of 23 polypeptides and 11 individual subunits, organized in four main structural domains: 1) a central actin-like filament consisting of eight ARP1 subunits and one actin; 2) a four-subunit pointed-end capping complex including the subunits p25, p27, p62, Arp11; 3) a barbed-end capping complex containing CapZ $\alpha\beta$ ; and 4) a shoulder domain containing p24, p150<sup>glued</sup>, and p50/dynamitin (Chowdhury et al., 2015; Lau et al., 2021; Urnavicius et al., 2015).

The interaction of Dynein and Dynactin (DD) is weak but is strongly promoted by different classes of activating adaptors that also promote interactions with various cargoes (Hoogenraad and Akhmanova, 2016; Olenick and Holzbaur, 2019; Reck-Peterson et al., 2018). The prototypical activating adaptor, BICD2 (bicaudal D homologue 2), favors the incorporation of Dynein and Dynactin into a single complex with greatly increased motility and processivity in comparison to the isolated Dynein (McKenney et al., 2014; Schlager et al., 2014a; Schlager et al., 2014b; Splinter et al., 2012; Urnavicius et al., 2015). The DD-binding segment maps to the BICD2 N-terminal region, which forms an apparently uninterrupted dimeric coiled-coil of  $\approx$ 250 residues that binds alongside the Dynactin filament. The BICD2 N-terminal region makes contacts near the barbed end, while a more C-terminal region of BICD2 makes contacts near the pointed end. This arrangement promotes binding to Dynein through several contacts with the N-terminal tail domain of the DHC as well as with the C-terminal region of the LICs (Celestino et al., 2014; Schroeder and Vale, 2016; Urnavicius et al., 2018; Urnavicius et al., 2020; Schroeder et al., 2014; Schroeder and Vale, 2016; Urnavicius et al., 2018; Urnavicius et al., 2015; Yeh et al., 2012).

In addition to the paradigmatic BICD2 adaptor and its family, several other proteins are or are likely to be activating adaptors, including CCDC88B, FIP3, HAP1, HOOK1-3, JIP3, Ninein and Ninein-like (NIN and NINL), NuMa, RAB11FIP3, RILP, Spindly (SPDL1), and TRAK1 (Gama et al., 2017; Hueschen et al., 2017; McKenney et al., 2014; Olenick et al., 2016; Redwine et al., 2017; Renna et al., 2020; Schroeder and Vale, 2016). At least some of these adaptors promote the interaction of Dynactin with two Dynein dimers, which increases processivity, speed, force production, and unidirectional movement (Grotjahn et al., 2018; Urnavicius et al., 2018). Collectively, the interactions of Dynein with Dynactin and adaptors appear to induce large conformational changes that align the motor domains for concomitant binding to microtubules (Zhang et al., 2017), an effect that likely extends to both Dynein dimers, when present.

While diverse, adaptors also share certain structural and functional features. These include 1) a propensity to form long coiled-coils with a dimeric parallel organization (Olenick and Holzbaur, 2019); 2) the ability of the N-terminal region to interact specifically with a conserved helix of LIC isoforms (Celestino et al., 2019; Gama et al., 2017; Lee et al., 2020; Lee et al., 2018). LIC-interacting sequences on adaptors belong to at least three different subfamilies, containing either a CC1 box, a HOOK domain, or EF-hand pairs (Olenick and Holzbaur, 2019; Reck-Peterson et al., 2018), which bind the LIC in different manners (Lee et al., 2020; Lee et al., 2018). The CC1 box (Figure 1A and Figure 1 – Supplement 1A) encompasses a highly conserved AAXXG sequence, where X denotes any amino acid (Gama et al., 2017; Hoogenraad and Akhmanova, 2016; Lee et al., 2018; Schlager et al., 2014b); 3) a second conserved motif, adjacent to the CC1 box and possibly also implicated in Dynein binding (Sacristan et al., 2018) (Figure 1A and Figure 1 – Supplement 1A). Even if it lies within the CC1 coiled-coil, this motif has been named the CC2 box (Sacristan et al., 2018). Here, we refer to it as the CC1's second conserved motif (CC1 SCM); 4) a more C-terminal Spindly box motif (L $\Theta$ XE $\Theta$ , where  $\Theta$  indicates an aliphatic or aromatic side chain; Figure 1A and Figure 1 – Supplement 1B), conserved in many adaptors, that mediates binding to the four subunits of the Dynactin pointed end subcomplex (p25, p27, p62, Arp11) (Gama et al., 2017; Lau et al., 2021); 5) an extended stretch of coiled coil (corresponding to CC1), positioned between the CC1 box and the Spindly motif, that lodges between Dynein and the Dynactin mini-filament, strongly enhancing complex formation (Chowdhury et al., 2015; Urnavicius et al., 2018; Urnavicius et al., 2015); and 6) a C-terminal domain, positioned after the Dynein-Dynactin binding region, shown or hypothesized to bind to cargo (Akhmanova and Hoogenraad, 2015; Hoogenraad et al., 2003).

The exquisitely specific regulation of Dynein at different subcellular structures implies that the interaction with adaptors is tightly regulated, stimulated to occur only at the appropriate time and subcellular locale. Thus, in solution the majority of Dynein adopts the so-called phi-particle conformation, unable to interact with Dynactin and adaptors (Amos, 1989; Zhang et al., 2017). The Dynactin p150 subunit can dock onto the pointed end subcomplex, sterically preventing adaptor binding (Lau et al., 2021; Urnavicius et al., 2015). Similarly, the cargo-binding domains of several adaptors, and prominently of BICD2, have been shown to fold back onto the Dynein-binding domains, forcing an autoinhibited conformation that can be relieved either by cargo binding, or by removing the cargo-binding domain in recombinant protein (Hoogenraad et al., 2003; Liu et al., 2013; Splinter et al., 2012; Stuurman et al., 1999; Terawaki et al., 2015; Urnavicius et al., 2015).

Here, we have addressed the organization and regulation of the 605-residue adaptor Spindly (Figure 1A), a farnesylated mitotic regulator of DD. In early mitosis, Spindly promotes the recruitment of DD to kinetochores, the structures that connect chromosomes to spindle microtubules (Musacchio and Desai, 2017). This function of Spindly is enabled by the 800 kDa hexameric ROD-Zwilch-ZW10 (RZZ) cargo complex, to which Spindly binds directly through its farnesylated C-terminal region (Holland et al., 2015; Mosalaganti et al., 2017; Moudgil et al., 2015). At kinetochores, phosphorylation by the MPS1 kinase promotes the polymerization of the RZZ-Spindly (RZZS) complex into a mesh (Figure 1B), the kinetochore corona, which adopts a characteristic crescent shape and contributes to the initial phases of chromosome alignment that precede end-on microtubule attachment and chromosome bi-orientation (Kops and Gassmann, 2020; Magidson et al., 2015; Pereira et al., 2018; Raisch et al., 2021; Rodriguez-Rodriguez et al., 2018; Sacristan et al., 2018). Besides DD, the corona also recruits the MAD1:MAD2 complex (Barisic et al., 2010; Chan et al., 2009; Cheerambathur et al., 2013; Gassmann et al., 2008; Gassmann et al., 2010; Griffis et al., 2007; Rodriguez-Rodriguez et al., 2018; Starr et al., 1998; Yamamoto et al., 2008), a central component of the spindle assembly checkpoint (SAC), which synchronizes cell cycle progression with the completion of chromosome alignment (Kops and Gassmann, 2020; Musacchio, 2015). Upon achievement of end-on attachment, the DD-RZZS complex becomes activated and begins to travel away from the kinetochore and toward the spindle poles, in a process of corona disassembly referred to as "corona shedding" or "stripping" (Auckland et al., 2020; Basto et al., 2004; Howell et al., 2001; Mische et al., 2008; Sivaram et al., 2009; Varma et al., 2008; Williams et al., 1996; Wojcik et al., 2001). As the MAD1:MAD2 complex is transported away from kinetochores together with DD-RZZS, this also suppresses SAC signaling. Mutations in the Spindly motif abrogate kinetochore recruitment of DD, blocking concomitantly corona shedding and SAC silencing (Cheerambathur et al., 2013; Gassmann et al., 2010).

Spindly has all the sequence credentials of a *bona fide* DD activating adaptor (Gama et al., 2017) (Figure 1A); however, it activates DD only weakly in motility assays *in vitro* (McKenney et al., 2014). This suggests that Spindly may exist in autoinhibited and active forms, as its presence alone is not sufficient for DD activation. Previous studies have also highlighted interactions between the N-terminal and C-terminal domains on Spindly, supporting the idea of an autoinhibitory interaction that hinders DD binding (Mosalaganti et al., 2017; Sacristan et al., 2018). Here, we demonstrate that Spindly is subject to a complex regulation involving large conformational changes, and identify crucial intramolecular contacts that regulate Spindly autoinhibition and prevent Spindly interaction with DD. Our data suggest that relief of Spindly autoinhibition requires the interaction with the RZZ and an additional kinetochore trigger that can be bypassed mutationally. Our results describe

an in-depth characterization of Dynein adaptor autoinhibition, and have important general implications for the mechanism of DD activation.

## Results

Spindly adopts a complex dimeric structural organization We have previously shown through solution scattering studies and hydrodynamic analyses that Spindly is an elongated molecule, and could be a dimer in solution (Mosalaganti et al., 2017; Sacristan et al., 2018). To conclusively show the stoichiometry of Spindly in solution, we employed mass photometry, which identified recombinant full-length Spindly (indicated as Spindly<sup>FL</sup> or Spindly<sup>WT</sup>) as a dimer, with an excellent agreement between theoretical and observed molecular masses (Figure 1C). We then determined the crystal structure of human Spindly<sup>1-100</sup> (Table S1; only residues 2-97 were clearly resolved in the electron density), a fragment containing both the CC1 box and the CC1 SCM (Figure 1D-F). The structure demonstrated that Spindly forms a parallel dimeric coiled-coil, similar in its outline to that observed in structures of other adaptors captured in complex with DD (Lau et al., 2021; Urnavicius et al., 2018; Urnavicius et al., 2015). Thus, both Spindly<sup>FL</sup> and an N-terminal segment of Spindly exist as stable dimers. The structure of Spindly<sup>2-97</sup> is closely reminiscent of the structure of BICD2<sup>1-</sup> <sup>98</sup> in complex with a peptide encompassing the LIC1 helix (residues 433-458, PDB ID 6PSE (Lee et al., 2020); Figure 1 – Supplement 1C). In Spindly, Ala23 and Gly27 in the CC1 box occupy a and d positions within the heptad repeats of the coiled-coil, similarly to Ala43 and Gly47 in BicD2, a pattern also conserved in BICDL1 and BICD1 (Figure 1 - Supplement 1A). This unusual composition for a and d residues generates a cavity along the BICD2 coiled-coil axis that interacts with aromatic and hydrophobic LIC side chains (Lee et al., 2020). Its conservation in Spindly was confirmed by high-confidence prediction by AlphaFold2 (AF2) in the variants Colabfold and AF2-Multimer (Evans et al., 2021; Jumper et al., 2021; Mirdita et al., 2021), which only became available during the final phases of this study (Figure 1 – Supplement 1D). Indeed, Spindly and BICD2 interact with the LIC with similar affinity (Lee et al., 2020; Lee et al., 2018).

Structural work on the complex of DD with various adaptors has demonstrated the existence of an uninterrupted coiled-coil spanning the distance between the LIC-binding CC1 box in BICD2 and a coiled-coil break that immediately precedes the Spindly motif. The coiled-coil, referred to as coiled-coil 1 (CC1), is cradled between Dynein and Dynactin (Urnavicius et al., 2018; Urnavicius et al., 2015). In Spindly, the coiled-coil propensity between the CC1 box and the break immediately preceding the Spindly motif (around residue 256) is generally high, but there is a conserved 2- or 3-residue insertion around residue 155 that maps to an interruption of the register of CC1 not

expected in BICD1 and BICD2 (Figure 1 – Supplement 1E-F). The conservation of this feature in the Spindly family prompted us to investigate the possibility that the CC1 coiled-coil of Spindly splits into distinct segments (CC1a and CC1b). For this, we subjected full-length Spindly to crosslinking-mass spectrometry experiments with the bifunctional crosslinker DSBU (disuccinimidyl dibutyric urea) (Pan et al., 2018). This revealed multiple intramolecular contacts within Spindly, including "concentric" crosslinks of the putative CC1b region with the second half of CC1a, consistent with the idea that they may be arranged as an anti-parallel pair separated by a loop containing the conserved insertion around residues 154-155. In addition, we observed crosslinks of the first half of CC1a with CC2 and CC3, and of CC2 with CC3 (Figure 1G, Figure 1 – Supplement G and Table S2). Essentially identical crosslinks were observed in experiments with Spindly<sup>1-440</sup> (lacking the flexible C-terminal region that contributes to binding the RZZ complex (Mosalaganti et al., 2017)), with the expected exception of contacts involving the C-terminal disordered region (Figure 1 – Supplement 1H-I).

These observations suggested that Spindly, at least in isolation, may adopt a compact conformation relative to the extended conformations observed for adaptors bound to DD (Urnavicius et al., 2018; Urnavicius et al., 2015). To probe this, we harnessed amber codon suppression to introduce the UV-photoactivatable crosslinker *p*-Benzoyl-L-phenylalanine (BPA) (Ai et al., 2011; Davis and Chin, 2012) into selected positions of an mCherry (mCh)-tagged construct of Spindly (<sup>mCh</sup>Spindly). These included Tyr26 and Gln29 (Y26<sup>BPA</sup> and Q29<sup>BPA</sup>, respectively) in the CC1 box and Phe258 (F258<sup>BPA</sup>) in the Spindly box. After irradiation with UV light (Figure 1 – Supplement 1J-K), Y26<sup>BPA</sup> and Q29<sup>BPA</sup> generated largely equivalent crosslinking patterns, with a majority of targets near the center of the predicted CC2 coiled-coil, around residue 300 (e.g. K297, L298, Q299, I300, L303, and M306; Figure 1H, Table S2, and Figure 1 – Supplement 1L). F258<sup>BPA</sup>, on the other hand, crosslinked to residues E74 and L76 immediately after the CC1 SCM in the central half of CC1a, as well as to residues in CC3 and the tail (Figure 1H, Table S2).

Collectively, these results establish that Spindly is a dimer, and that it may fold as four interacting coiled-coil segments (CC1a, CC1b, CC2, and CC3), with extensive interactions of the first half of CC1a with CC2 and CC3, and of the second half of CC1a with CC1b. A high-confidence AF2 prediction rationalized these observations (Figure 1I and Figure 1 – Supplement 2). AF2 models depict Spindly as having a complex organization, where CC1 is almost invariably predicted to be interrupted around residues 154-155, giving rise to CC1a and CC1b coiled-coil segments. The  $\approx$ 23 nm CC1a coiled-coil, partly captured in our crystal structure (Figure 1D), is predicted to encompass the majority of the long-axis of the Spindly dimer ( $\approx$ 28 nm), in good agreement with values derived

from 2D class averages of negatively stained Spindly samples and small angle X-ray scattering (SAXS) experiments for both Spindly<sup>FL</sup> and Spindly<sup>1.440</sup> (Sacristan et al., 2018). CC1b packs against CC1a in such a way that the Spindly motif, located at the beginning of CC2, is positioned roughly halfway along the complex, facing the segment immediately C-terminal to the CC1 SCM, in excellent agreement with the BPA crosslinking experiments (Figure 1I and Figure 1 – Supplement 2). A loop around residue 360 separates CC2 and CC3, in such a way that both CC2 and CC3 are in contact with the first part of CC1. The CC1 box faces precisely the region centered on residue 300 identified by the crosslinking experiments with Y26<sup>BPA</sup> and Q29<sup>BPA</sup>. Importantly, the model predicts that both the CC1 box and the Spindly motif will be largely inaccessible to DD. While all parallel coiled-coils are roughly symmetric, residues in the two chains experience different environments due to the asymmetric intramolecular interactions of the coiled-coils. Thus, the folded structure of Spindly predicted by AF2 is inherently asymmetric.

Spindly is not accessible to Dynein-Dynactin In addition to structural work demonstrating that the adaptors showcase an uninterrupted CC1 to DD, a non-comprehensive AF2 analysis of a subset of adaptors (including BICD2, BICDL1, HOOK1, HOOK3, Spindly, and TRAK1) confirmed that they all have an N-terminal extended dimeric coiled-coil section of variable lengths. In these models, the CC1 continues more or less uninterrupted until a break of variable length (where CC propensity drops). This precedes CC2, which usually begins with, or is even preceded by, the Spindly motif (Figure 2 - Supplement 1A). This is true also of BICD2 (Figure 2A), even if a coiledcoil prediction algorithm suggested a  $\approx$  30-residue drop in coiled-coil propensity after the CC1 SCM (Figure 1 – Supplement 1E). The AF2 models show that the CC1 in all these adaptors has a rather regular length of 35-39 nm. In other adaptors, including JIP3 and RILP, CC1 is considerably shorter (approximately 20 nm) and there is no obvious Spindly motif (Reck-Peterson et al., 2018). As established in Figure 1 and Figure 1 – Supplement 2, Spindly can also adopt a closed compact conformation, but its constellation of DD binding motifs predicts that it opens as a canonical adaptor under appropriate conditions. Indeed, AF2 predicted a closed conformation for Spindly<sup>1-</sup> <sup>309</sup> (Figure 2B, *right* and Figure 1 – Supplement 2), while it did not predict convincing intramolecular interactions for Spindly<sup>1-275</sup> (Figure 2B, left and Figure 1 – Supplement 2). This suggests that residues 276-309, a region that according to our XL-MS analysis faces the CC1 box, contains determinants of a conformational transition from a closed to an open form.

In this context, a question of significant mechanistic relevance is whether binding of Spindly to its cargo is sufficient to relieve autoinhibition and trigger the formation of a complex with DD, as

postulated for other adaptors (Olenick and Holzbaur, 2019; Terawaki et al., 2015). To address this question, we used analytical size-exclusion chromatography (SEC) to monitor the formation of complexes of DD with either Spindly or BICD2<sup>1-400</sup>, which served as positive control (Schlager et al., 2014a). Because the Dynein phi-particle might prevent Dynein from engaging into complexes with Dynactin and adaptors, we used a Dynein tail construct (CDCH<sup>1-1455</sup>) that does not form the phi-particle (Urnavicius et al., 2015; Zhang et al., 2017). As expected, Dynein, Dynactin, and BICD2<sup>1-400</sup> interacted in a complex that eluted before any of the individual components, indicative of an increased Stokes' radius (Figure 2C). Conversely, there was no evidence of binding of Dynein and Dynactin to full length Spindly (Figure 2D). This result is consistent with the hypothesis that Spindly adopts an auto-inhibited conformation refractory to interact with DD in the absence of adequate triggers.

The RZZ complex, to which farnesylated Spindly (Spindly<sup>F</sup>) binds directly, mediates Spindly's kinetochore recruitment and can therefore be considered Spindly's cargo, or a connector of Spindly to its chromosome cargo. We asked therefore if Spindly interacted with DD in presence of the RZZ complex. As the RZZ complex and Spindly are both known to be phosphorylated, with phosphorylation being critical for corona function (Raisch et al., 2021; Rodriguez-Rodriguez et al., 2018), we tested binding to DD after dephosphorylation (RZZ<sup>D</sup> and Spindly<sup>D</sup>, Figure 2E), or after additional incubation with a mix of ATP and mitotic kinases, including CDK1/Cyclin B, MPS1, and Aurora B (RZZ<sup>P</sup> and Spindly<sup>P</sup>, Figure 2F and Figure 2 – Supplement 1B). In either case, no interaction of the RZZS<sup>F</sup> complex with DD was detected. Thus, the RZZ is not sufficient to relieve Spindly auto-inhibition and DD binding. Results presented below suggest that the kinetochore itself may play a role in the activation of Spindly required for DD binding.

*Relieving Spindly auto-inhibition* We next attempted to investigate what forces contribute to maintain Spindly's autoinhibited state. The pointed end (PE) subcomplex of Dynactin, which interacts with the conserved Spindly motif, has previously been shown to bind adaptors with limited but measurable binding affinity (Gama et al., 2017; Lau et al., 2021; Yeh et al., 2012). Thus, we developed a minimal recombinant adaptor-binding subcomplex of Dynactin, containing only the subunits of the pointed end-capping complex, p25, p27, p62 and Arp11 (Figure 3A). After purification to homogeneity, we tested whether this pointed end (PE) complex interacted with different fragments of Spindly. In SEC experiments, the PE subcomplex did not bind <sup>mCh</sup>Spindly<sup>FL</sup>, in agreement with the possibility that Spindly is auto-inhibited (Figure 3B). Conversely, Spindly<sup>1-275</sup>, a fragment that contains the Spindly box and that our AF2 predictions identified as having an

open, elongated coiled-coil (Figure 2B, *left*), bound to the PE complex, as indicated by a clear, albeit only partial shift in its elution volume (Figure 3C). The interaction of Spindly<sup>1-275</sup> with the PE complex required the Spindly motif, as Spindly<sup>1-250</sup>, a construct lacking it, was unable to bind the PE (Figure 3D). C-terminal deletions have been shown to relieve autoinhibition in BICD2, mimicking the effect of cargo binding (McKenney et al., 2014; Schlager et al., 2014b). Thus, we tested various Spindly C-terminal deletions for their ability to interact with the PE complex in the absence of cargo and other activators. Constructs lacking only the (disordered) C-terminal tail (Spindly<sup>1.440</sup>), or lacking the C-terminal tail and the CC3 (Spindly<sup>1.354</sup>) did not bind the PE complex in SEC experiments (Figure 3 - Supplement 1A), most likely because they adopt a closed conformation related to that predicted by AF2 for the Spindly<sup>1-309</sup> construct (Figure 2B, *right*). We reasoned therefore that a Spindly deletion mutant lacking determinants of autoinhibition in the CC2 region ought to show features of the open complex and bind the PE complex even in the context of full-length Spindly. Indeed, an <sup>mCh</sup>Spindly mutant lacking residues 276-306, a segment already identified for its interactions with the CC1 box, showed a clear level of binding to the PE complex (Figure 3E). Spindly<sup>1-275</sup> and <sup>mCh</sup>Spindly<sup>Δ276-306</sup> did not fully co-elute with the PE complex, possibly indicative of low binding affinity. These constructs may be only partially open, or their binding site for the PE may be partly disrupted. We have also found that both constructs populate a dimer-tetramer equilibrium and that the tetramer, especially for Spindly<sup>1-275</sup>, may prevail at the micromolar concentration used in these experiments, possibly counteracting the interaction with the PE (unpublished observations and see Figure 4C,E, discussed below).

We suspected that the N-terminal region of Spindly, which our experiments have suggested to be split in CC1a and CC1b segments, contributes to stabilize the autoinhibitory interaction that controls access of the PE complex to the Spindly motif. To test this idea, we asked if the <sup>mCh</sup>Spindly<sup>76-605</sup> construct, lacking the segment of CC1a where both the CC1 box and the CC1 SCM are located, bound the PE complex. Indeed, <sup>mCh</sup>Spindly<sup>76-605</sup> bound the Dynactin pointed end in SEC assays (Figure 3 – Supplement 1B), confirming that the N-terminal segment of Spindly contributes to maintain the autoinhibited state of Spindly. Finally, a mutant lacking the entire CC1a region and also truncated in the CC2 after the Spindly motif, Spindly<sup>156-275</sup>, interacted robustly with the PE complex (Figure 3F).

To identify the role of the CC1 box in this process, we designed two sequential N-terminal truncations. <sup>mCh</sup>Spindly<sup>22-605</sup>, which retains the CC1 box, did not bind the PE complex. Conversely, <sup>mCh</sup>Spindly<sup>33-605</sup>, which does not retain the CC1 box, bound the PE complex efficiently (Figure 3G). Two further short deletion mutants within the CC1, <sup>mCh</sup>Spindly<sup>Δ26-28</sup> and <sup>mCh</sup>Spindly<sup>Δ26-32</sup> respectively

did not bind and bound weakly to the PE complex (Figure 3 – Supplement 1C-D), indicating that shorter deletions elicit a less penetrant effect on Spindly auto-inhibition. Collectively, these observations indicate that the Spindly auto-inhibition mechanism involves a tight intramolecular interaction of the conserved, Dynein-binding CC1 box with a regulatory segment of the Spindly CC2 coiled-coil roughly comprised between residues 276 and 306. Our results also imply that this inhibitory control cannot be readily relieved by DD, not even in presence of cargo (RZZ), possibly implying that an additional trigger at the kinetochore catalyzes opening.

*Testing structural predictions* The interactions between CC1 and CC2 in the AF2 model of Spindly in Figure 1I, together with our extensive analysis in Figure 3, explain why Spindly<sup>1-250</sup> (CC1) and Spindly<sup>250-605</sup> (CC2 and CC3) interact with high affinity and coelute from a SEC column (Sacristan et al., 2018), a result that we could readily reproduce (Figure 4A). When tested in the same assay, however, Spindly<sup>51-250</sup>, where the N-terminal segment of CC1a predicted to bind CC2 has been removed, did not interact with Spindly<sup>250-605</sup>. Similarly, Spindly<sup>1-250</sup> seemed unable to bind Spindly<sup>250-</sup>

The disruption of interactions responsible for intramolecular folding may be expected to render the Spindly deletion mutants more elongated. To test this prediction, we first used mass photometry to verify that <sup>mCh</sup>Spindly<sup>76-605</sup> and <sup>mCh</sup>Spindly<sup>Δ276-306</sup> remained dimeric like wild type <sup>mCh</sup>Spindly. This was confirmed for both constructs (with <sup>mCh</sup>Spindly<sup>Δ276-306</sup> being predominantly dimeric but also showing a tendency to form tetramers already at very low concentration, Figure 4C,E). We then analyzed the sedimentation behavior of these constructs by analytical ultracentrifugation, and derived their Stokes' radii and frictional ratios (see Methods) (Figure 4D). For a given molecular mass, the product of the sedimentation coefficient and of the Stokes' radius is a constant (Siegel and Monty, 1966). Indeed, the sedimentation coefficient of the deletion mutants was reduced in comparison to that of <sup>mCh</sup>Spindly<sup>FL</sup>, indicative of larger Stokes' radii and frictional ratios and therefore of a more elongated conformation (Figure 4E). Thus, deletion of regions in CC1a and CC2 predicted to interact with each other in the folded conformation of Spindly cause an at least partial opening of the Spindly structure.

**Opening up Spindly with point mutations** The region of Spindly downstream of the Spindly box (residues 281-322) is very conserved among Spindly orthologues, but not among other members of the BICD adaptor family (Figure 1 – Supplement 1L). Within this region, we mutated the positively charged residue pair R295-K297 to glutamate. The resulting construct, indicated as Spindly<sup>CC2\*</sup> (where CC2\* indicates the R295E-K297E mutant in the CC2 coiled-coil) bound the PE complex in SEC (Figure 5A). The interaction with the PE complex was mediated by the Spindly motif, because combining the CC2\* mutation with a mutation in the Spindly motif (F258A, indicated as SM\*) to generate the <sup>mCh</sup>Spindly<sup>SM\*-CC2\*</sup> mutant, abolished the interaction (Figure 5B). In an orthogonal approach, we developed <sup>mCh</sup>Spindly<sup>ΔRV</sup>, a construct deleted of the Spindly-specific 2-residue insert (residues R154-V155) between the CC1a and CC1b coiled-coil segments (Figure 1 – Supplement 1F) with the goal of favoring a full extension of CC1 like in BICD2, which lacks the insertion. <sup>mCh</sup>Spindly<sup>ΔRV</sup> bound the PE complex (Figure 5C), albeit more weakly, suggesting that that the 2-residue insertion into the CC1 of Spindly favors autoinhibition.

In mass photometry measurements, <sup>mCh</sup>Spindly<sup>CC2\*</sup> and <sup>mCh</sup>Spindly<sup>ΔRV</sup> had masses expected of dimers (Figure 5D) and essentially indistinguishable from those of <sup>mCh</sup>Spindly<sup>FL</sup> (Figure 1C), suggesting that their ability to interact with the PE complex does not result from changes in stoichiometry. This was further confirmed by introducing a GST in Spindly<sup>FL</sup> and Spindly<sup>CC2\*</sup> to reinforce dimerization of both. GST-Spindly<sup>FL</sup> did not bind the PE complex, whereas GST-Spindly<sup>CC2\*</sup> did (Figure 5 – Supplement 1A-B). Analytical ultracentrifugation demonstrated a decreased sedimentation coefficient for <sup>mCh</sup>Spindly<sup>CC2\*</sup> and <sup>mCh</sup>Spindly<sup> $\Delta RV$ </sup> (Figure 5E), indicative of a more extended conformation, as already shown for the Spindly deletion mutants in Figure 4D-E. Further, we developed an mCherry-tagged BICD2-Spindly chimeric construct that combined the BICD2 N-terminal region until the Spindly motif (residues 1-292<sup>BICD2</sup>), which is believed to contain an uninterrupted CC1, with the Spindly motif and C-terminal RZZ-binding domain of Spindly (residues 251-605<sup>Spindly</sup>; Figure 5F). Indeed, AF2 modelled this construct (Spindly<sup>Chimera</sup>) with a continuous CC1 until the flexible region that precedes the Spindly box (Figure 5 – Supplement 1C). Spindly<sup>Chimera</sup> promoted Spindly-dependent oligomerization of the RZZ complex in filaments in vitro (Figure 5G), which requires MPS1 kinase and mimics kinetochore corona assembly (Raisch et al., 2021; Rodriguez-Rodriguez et al., 2018; Sacristan et al., 2018), thus indicating that the Spindly segment in Spindly<sup>Chimera</sup> is sufficient for polymerization. In fact, Spindly<sup>Chimera</sup> was more prone to promoting filaments than Spindly<sup>WT</sup>, and filaments were even observed at room temperature in absence of Mps1, when Spindly<sup>WT</sup> did not stimulate filament formation (Figure 5 - Supplement

1D). In SEC experiments, <sup>mCh</sup>Spindly<sup>Chimera</sup> bound the PE complex (Figure 5H), in agreement with our expectation that an uninterrupted CC1 allows the PE to access the Spindly motif.

Finally, we asked whether relief of autoinhibition would increase the affinity of Spindly for Dynein in addition to increasing the affinity for the PE complex of Dynactin. As the CC1 box, required for the interaction with the Dynein LIC, is directly involved in the autoinhibitory interaction, we asked whether we could see increasing binding of LIC upon straightening Spindly to render the CC1 box more accessible. As the affinity of the LIC for CC1 box containing adaptors is too low for accurate study by SEC, we used a pull-down assay. We produced recombinantly a GST-tagged construct of the LIC2 isoform, and we tested its ability to pull down Spindly wild-type and "open" mutants. As a negative control, we used the <sup>mCh</sup>Spindly<sup>AA/VV</sup> (A23V-A24V) mutant, which has been previously shown to inhibit the interaction with the LIC2 in a similar assay (Gama et al., 2017). As open mutants, we used <sup>mCh</sup>Spindly<sup>CC2\*</sup> and <sup>mCh</sup>Spindly<sup>ARV</sup>. There was a clear increase of affinity for the LIC of the open mutants, with <sup>mCh</sup>Spindly<sup>CC2\*</sup> showing an even slightly higher affinity than <sup>mCh</sup>Spindly<sup>ARV</sup>, possibly due to an only partial restoration of coiled-coil continuity in the latter (Figure 5I-J), and in line with the lower apparent affinity of <sup>mCh</sup>Spindly<sup>ARV</sup> for the PE. Collectively, these results indicate that open mutants of Spindly interfering with the stability of the CC1-CC2 interaction or with the bending of the CC1 coiled-coil are more easily accessible to DD.

*Spindly*<sup>CC2\*</sup> *binds Dynein and Dynactin with higher affinity than Spindly*<sup>WT</sup> After showing that Spindly<sup>WT</sup> does not interact with DD, we were eager to test whether the open Spindly mutants formed a super-complex with DD. For these assays, we were able to set up the production of recombinant human Dynactin, a significant technical challenge (Lau et al., 2021; Urnavicius et al., 2015). We produced Dynactin in the HEK293<sup>expi</sup> expression system, using a modified version of the pBiG2 plasmid from the biGBac system (Weissmann et al., 2016) (Figure 6 – Supplement 1A). The resulting pBiG2 was used to transfect insect cells to produce a baculovirus and then to infect Expi293F cells (see Methods). The purified human Dynactin is biochemically pure and similar to the protein complex purified from pig brains, except for the absence of the p135 isoform of p150<sup>glued</sup>, which is not expressed in our system (a band in the same position is likely caused by degradation of the p150<sup>glued</sup> subunits and is marked with an asterisk in Figure 6 – Supplement 1B). SEC combined with multiangle light scattering (SEC-MALS) measurements demonstrated that human recombinant purified Dynactin has the expected molecular mass (Figure 6 – Supplement 1C). Furthermore, recombinant Dynactin bound BICD2<sup>1.400</sup> like pig brain Dynactin (Figure 6 – Supplement 1D-E). Finally, recombinant human Dynactin appeared morphologically

indistinguishable from pig brain Dynactin, as judged by 2D classes from negative stain electron microscopy (Figure 6 – Supplement 1F; see also (Urnavicius et al., 2015)).

As we have shown that fragments from both Dynein (the LIC) or Dynactin (the PE) bind the CC2\* mutant independently, we performed binding assays using stoichiometric ratios of Spindly, Dynein tail, and recombinant human Dynactin to maximize complex formation (Figure 6A). <sup>mCh</sup>Spindly<sup>WT</sup> was unable to form the super-complex with DD, as expected, but <sup>mCh</sup>Spindly<sup>CC2\*</sup> and <sup>mCh</sup>Spindly<sup>Chimera</sup> were, as assessed by the shift of the adaptor into the expected super-complex peak (Figure 6A-B). We also found that <sup>mCh</sup>Spindly<sup>33-605</sup> interacted with Dynein-Dynactin, albeit apparently more weakly than <sup>mCh</sup>Spindly<sup>CC2\*</sup> or <sup>mCh</sup>Spindly<sup>Chimera</sup>. Both <sup>mCh</sup>Spindly<sup>CC2\*</sup> and <sup>mCh</sup>Spindly<sup>Chimera</sup> were able to interact with the RZZ complex, indicating that the mutations do not affect the cargo-binding region of RZZ (Figure 6C-D).

The localization of Spindly to human kinetochores Finally, we assessed the ability of the different Spindly mutants to reach kinetochores in human cells arrested in mitosis with a spindle poison. Endogenous Spindly was depleted by RNA interference (RNAi) and recombinant <sup>mCh</sup>Spindly variants were introduced in cells through electroporation as summarized in Figure 7 - Supplement 1A-B. As shown previously (Gassmann et al., 2010), depletion of Spindly prevented kinetochore recruitment of Dynactin (Figure 7A-D). Electroporation of <sup>mCh</sup>Spindly<sup>WT</sup> largely rescued these effects, but both the <sup>mCh</sup>Spindly<sup>CC2\*</sup> and the <sup>mCh</sup>Spindly<sup>Chimera</sup> constructs showed strongly reduced kinetochore levels and, accordingly, greatly reduced levels of Dynactin, as measured with an antibody against the p150<sup>glued</sup> subunit. The Spindly and Dynactin signals at kinetochores were well correlated for all four constructs (Figure 7E and Figure 7 – Supplement 1C-G). This argues that the reduction in kinetochore levels of Dynactin arises due to a reduction in the kinetochore levels of the <sup>mCh</sup>Spindly<sup>CC2\*</sup> and <sup>mCh</sup>Spindly<sup>Chimera</sup> constructs, rather than to an inability of Dynactin to interact with them. This conclusion is also in line with our biochemical data showing that these Spindly mutants interact with Dynein and Dynactin. Because Spindly<sup>Chimera</sup> promoted RZZ filamentation with enhanced efficiency in vitro (Figure 5 - Supplement 1D), we reasoned that its reduction at kinetochores may reflect the assembly of ectopic complexes with DD no longer limited to kinetochores, thus reducing the pool available for kinetochore binding. Indeed, <sup>mCh</sup>Spindly<sup>Chimera</sup> formed ectopic corona-like filaments with Dynactin in interphase cells in the absence of kinetochores (Figure 7F).

Kinetochore localization by Spindly<sup>CC2\*</sup> was evidently severely impaired but this construct did not form ectopic corona-like filaments (unpublished results). Filamentation assays with RZZ showed

it was almost unable to promote MPS1-dependent filament assembly (Figure 7G). Thus, it appears that the CC2\* mutations (R295E-K297E) affect at the same time Spindly's filament formation and kinetochore recruitment, without impairing DD binding. CC2\* shares these characteristics with Spindly<sup>Δ276-306</sup> (Figure 3E and (Raisch et al., 2021)). Collectively, these observations suggest that the region of CC2 containing these residues is at the same time implicated in kinetochore binding and filament formation, suggesting a possible mechanism of kinetochore regulation of corona expansion.

<sup>mCh</sup>Spindly<sup>33-605</sup> not only decorated kinetochores to levels that were indistinguishable from those of <sup>mCh</sup>Spindly<sup>WT</sup>, but also promoted recruitment of comparable levels of Dynactin (Figure 7A-D and Figure 7 – Supplement 1G). By suggesting that binding of the CC1 box to LIC1 is not required for robust DD recruitment at human kinetochores, this result may seem at odd with the observation that the Spindly<sup>A23V</sup> mutant (where the CC1 box is mutated rather than absent) strongly impairs kinetochore recruitment of Dynactin (Sacristan et al., 2018). Our <sup>mCh</sup>Spindly<sup>AA/VV</sup> mutant could not be used to further investigate the issue, as – for unclear reasons – it was unable to reach kinetochores (unpublished results), preventing us from comparing it to <sup>mCh</sup>Spindly<sup>33-605</sup> in the same assay. Nonetheless, the new results with <sup>mCh</sup>Spindly<sup>33-605</sup> suggest that the deletion of the CC1 box or its mutation result in fundamentally distinct behaviors, and support a role of the LIC subunits as triggers of adaptor opening more than as decisive contributors to the binding affinity of the interaction, a speculative conclusion that will require further investigations.

## Discussion

Spindly has been mainly studied as a mitotic adaptor of Dynein-Dynactin (DD), but recent observations suggest that its role might extend to interphase cells (Del Castillo et al., 2020). Here, we have dissected a mechanism of conformational control that addresses the functions of Spindly at kinetochores, and that might inform future studies of Spindly in other cellular locales and cell cycle phases. More generally, our studies have implications for the control of DD in time and space. Adaptors are known to activate the motility and processivity of Dynein by stabilizing a complex with Dynactin (Hoogenraad and Akhmanova, 2016; McKenney et al., 2014; Schlager et al., 2014b). Studies so far have focused on N-terminal segments of adaptors that overcome intramolecular regulation, exemplified by the BICD2<sup>1-400</sup> construct. Previous studies had demonstrated that the C-terminal cargo-binding region decreases the affinity of BICD2 for DD (Hoogenraad et al., 2003; Liu et al., 2013; Splinter et al., 2012; Terawaki et al., 2015). This effect was clearly evident also with Spindly, whose full-length form failed to bind DD in a variety of assays. However, we demonstrate

that Spindly undergoes an apparently more complex regulation, that is not limited to cargo binding, but also includes a folded-back conformation of the N-terminal coiled-coil.

The RZZ complex, considered the Spindly cargo at kinetochores, binds directly to farnesylated Spindly (Mosalaganti et al., 2017; Raisch et al., 2021). The observation that binding of RZZ to Spindly was insufficient to unleash a conformational change compatible with DD binding motivated our investigation of the detailed mechanisms that hold Spindly in a closed conformation. A Spindly fragment encompassing residues 354-605 is sufficient to bind RZZ (Henen et al., 2021; Raisch et al., 2021), but the autoinhibited conformation requires a fragment of Spindly comprised between residues 276 and 309, and therefore positioned upstream of the minimal RZZ-binding region (Figure 7H). AF2 modeling (Evans et al., 2021; Jumper et al., 2021), which became available in the late phases of our work and proved to be instrumental as a validation tool, suggests that Spindly<sup>1-275</sup> adopts an open conformation that is closely reminiscent of that of BICD2<sup>1-400</sup>, whereas Spindly<sup>1-309</sup> may adopt a closed conformation. While these predictions must be taken with caution in the absence of supporting experimental evidence, they reinforce conclusions from our detailed biochemical and biophysical analysis, which are entirely consistent with this idea. Specifically, this model is supported by crosslinking analyses that predict the Spindly CC1 box to be in close proximity with residues 295-305, and by evidence that the closed conformation correlates with a break in the CC1 coiled-coil that allows Spindly to bend back on itself around residue 155. This model also explains a pattern of intra-molecular contacts, revealed by crosslinking-mass spectrometry, that would otherwise not be expected for a parallel coiled-coil like the one in Spindly. Our mutational analysis, combined with a binding assay monitoring the interaction with the pointed-end complex of Dynactin, and thus measuring the accessibility of the Spindly motif, was entirely consistent with the hypothesis that these two structural determinants, namely the 2-residue insertion in CC1 and the 295-305 region, are crucial for the auto-inhibitory mechanism of Spindly. This model was later confirmed by demonstrating binding of Spindly mutants to the entire Dynein-Dynactin complex. For these experiments, we established the expression of a recombinant form of Dynactin, a highly desirable development that will enable the production of significantly more homogeneous samples and the design of mutants, as well as enabling the use of tags for the individual subunits.

If RZZ binding is insufficient for unleashing the potential of Spindly to bind DD, what else might be required? Previous work has demonstrated that the RZZ complex is necessary for the recruitment of Spindly to the kinetochore and that Spindly, in turn, recruits DD (Barisic et al., 2010; Chan et al., 2009; Cheerambathur et al., 2013; Gassmann et al., 2008; Gassmann et al., 2010;

Griffis et al., 2007; Raaijmakers et al., 2013; Starr et al., 1998; Yamamoto et al., 2008). Because the RZZ complex does not appear to be sufficient to promote binding of Spindly to DD, our analysis implies that a second, unknown interaction in the kinetochore elicits the opening of Spindly. The identity of the binding partner is unknown and will represent the focus of our future investigations. A possible working model is that the still unknown kinetochore binder interacts with the region comprised between residues 276-306, relieving it from its intra-molecular control of the Spindly closed conformation, and promoting the transition to the open conformation. Evidence supporting this idea is that mutations in the 276-306 region, including the deletion of this entire fragment or the introduction of charge-inverting point mutations at residues 295 and 297 respectively abolish or largely decrease the kinetochore recruitment of Spindly ((Raisch et al., 2021)) and this study), implying that they impinge directly not only on the closed-open transition but also on the Spindly recruitment mechanism. *In vitro*, the 276-306 region is also required for the assembly of RZZ-Spindly filaments (this study and (Raisch et al., 2021)). However, this is unlikely to explain the kinetochore localization defect, because corona expansion is not required for robust recruitment of Spindly to the kinetochore (Raisch et al., 2021; Rodriguez-Rodriguez et al., 2018).

Our recent report of the cryo-EM structure of the RZZ complex (Raisch et al., 2021) represents a milestone of mechanistic studies of corona assembly at the kinetochore. We also investigated the mechanism of corona assembly and the role of Spindly in this process, identifying important roles for protein phosphorylation, in agreement with previous studies (Barbosa et al., 2020; Rodriguez-Rodriguez et al., 2018). Nonetheless, how Spindly promotes assembly of the corona, and the detailed organization of the corona, remain poorly understood. The RZZ is evolutionarily and structurally related to precursors of the coats that surround membrane vesicles during intra-cellular transport (Civril et al., 2010; Mosalaganti et al., 2017), and is thus likely to polymerize through similar mechanisms. Plausibly, the solution to this conundrum will require biochemical reconstitutions addressing the spectrum of interactions that this protein establishes at the kinetochore.

In conclusion, we have studied in mechanistic detail the basis of Spindly activation at the kinetochore and shown it to reflect a structural transition from a closed to an open conformation. We identified several structural determinants likely involved in the transition, and proposed a 2-step activation mechanism comprising RZZ binding and binding to an unknown second kinetochore receptor ultimately required for binding of Spindly to Dynein-Dynactin. These studies have important general implications for the mechanism of activation of adaptors, as they imply that the definition of "cargo" must be nuanced to every particular situation in which an adaptor is

involved. In our particular analysis, the RZZ complex continues to be legitimately considered an element of the adaptor's cargo, but our data imply that a second, equally important component remains to be identified. We speculate that a similar 2-step or multistep mechanism may apply to additional cargo-adaptor systems.

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#### Materials and Methods

Mutagenesis and cloning cDNA segments encoding for wildtype Spindly or truncated constructs were subcloned into a pET28-mCherry plasmid, with an intervening PreScission cleavage site, or into a pLib plasmid for insect cell expression, with a 5' insert coding for a His6-tag. Mutations were introduced by site-directed mutagenesis by Gibson assembly (Gibson et al., 2009). All constructs were sequence-verified. The BicD2-Spindly chimeric construct (mChSpindlyChimera) was created by Gibson Assembly. We generated a sense primer containing the last 25bp of the coding sequence for the BicD2 (1-292) segment, and the first 25bp of the coding sequence for the Spindly (251-605) segment, as well as an antisense primer containing the same region. We used the former to expand the Spindly (251-605) sequence with an additional 3' overlap with the MCS of a pLibmCherry plasmid, and the latter to expand the BicD2 (1-292) coding sequence with an additional 5' overlap with the pLib-mCherry plasmid MCS. The two PCR products were then assembled by Gibson Assembly into a pLib-mCherry plasmid opened by restriction cloning (BamHI-HindIII). A plasmid for the expression of the Dynactin pointed end was generated using the biGBac system (Weissmann et al., 2016). Coding sequences for Arp11, p62, p25-6His and p27 were subcloned into individual pLib plasmids, which were then used to build a pBiG1 plasmid, which was then used for expression of the entire complex. For the GST-LIC2 construct, cDNA encoding for human LIC2 was subcloned into a pLib vector containing an N-terminal TEV-cleavable GST.

The following Dynactin coding genes were codon-optimized for protein expression in insect cells and synthesized (Epoch Life Science): DCTN1 (Protein name p150, Uniprot Isoform 1, NCBI Reference Sequence NM\_004082.4), DCTN2(p50, Isoform 1, NM\_006400.4), DCTN3 (p24, Isoform 1, NM\_007234.4), DCTN4 (p62, Isoform 1, NM\_016221.3), DCTN5 (p25, Isoform 1, NM\_032486.3), DCTN6 (p27, NM\_006571.3), CAPZA1 (CapZ $\alpha$ -1, NM\_006135.2), CAPZB (CapZ $\beta$ , Isoform 2, NM\_004930.4), ACTR1A (Arp1, NM\_005736.3), ACTR10 (Arp11, NM\_018477.2), ACTB ( $\beta$ -actin, NM\_001101.4). All genes were cloned into the pACEBac1 vector (Geneva Biotech) in between the polyhedron (polH) promoter and Simian virus 40 (SV40) poly A sequences by Gibson cloning. A sequence encoding the ZZ affinity tag plus a TEV-cleavage site was fused in-frame to the 5'-end of the DCTN1 gene. For cloning of mammalian cell expression vectors, the genes were cloned out of pACEBac1 into pcDNA<sup>TM</sup>4/TO (Thermo Fisher Scientific) in between the CMV promoter and the BGH poly A signal. To assemble the dynactin genes into a single expression plasmid, biGBac cloning was used and adapted for mammalian expression. The biGBac cloning plasmids (pBig1a, pBig1b, pBig1c, pBig2abc) were generated by Gibson cloning and using the pACEBac1 vector as the backbone. To amplify the gene expression cassettes (GECs)

by PCR and to assemble them into pBig1 plasmids, a set of CasMam oligonucleotides was designed and synthesized (Sigma-Aldrich/Merck, desalt purification). The oligonucleotides contained the optimized linker sequences  $(\alpha/\beta/\gamma/\delta/\epsilon/\omega)$  as described in Weissmann et al., 2016 and were modified complementary to the 5'-end CMV to be of promoter sequence (GTTGACATTGATTATTGACTAG - forward oligonucleotide) and reverse-complementary to the 3'-end of the BGH polyA sequence (CCATAGAGCCCACCGCATCC - reverse oligonucleotide). The GECs were then used to build three pBiG1 plasmids: pDCTN A, containing the ZZ-TEV-tagged p150<sup>glued</sup>, p50, and p24; pDCTN B, containing Arp1, Arp11, β-actin, CapZα, and CapZ<sub>β</sub>; and pDCTN C, containing p25, p27, and p62 subunits. The thee pBiG1 plasmids were then used to build a single pDCTN FL plasmid. Successful assembly of pDCTN FL was confirmed by complete plasmid sequencing (CCIB DNA Core Facility at Massachusetts General Hospital (Cambridge, MA)).

Expression and purification of RZZ, mCherry-Spindly and Spindly constructs The RZZ complex was expressed and purified using the biGBac system, with an mCherry N-terminally fused tag on the ROD subunit, as previously described (Sacristan et al., 2018). Expression of all mCherry-Spindly constructs and mutants except the "ChSpindlyChimera was carried out in E. coli. BL21 CodonPlus cells were transformed with the plasmid, and grown in TB at  $37^{\circ}$  C to an OD<sub>600</sub> of 0.5. Expression was induced with 0.4 mM IPTG. The culture was then transferred into an incubator pre-cooled to 18° C, and grown overnight before harvesting. The pellet was then snap-frozen and stored at -80° C until purification. <sup>mCh</sup>Spindly mutants containing the unnatural amino acid Bpa were expressed in E. coli BL21 strains containing the pEVOL-pBpF plasmid (Chin et al., 2002). Cells were cultured in selective (kanamycin, chloramphenicol) TB media, supplemented with 0.2% arabinose, to trigger expression of the tRNA synthetase/tRNA pair. Cells were grown at 37° C until an  $OD_{600}$  of 0.6 was reached. Expression was induced with 0.5 mM IPTG, and the Bpa was added to the bacterial culture at a concentration of 1 mM. The culture was then transferred into an incubator pre-cooled to 18° C, and grown overnight before harvesting. The pellet was then snap-frozen and stored at -80° C until purification. Spindly constructs without the mCherry tag and the <sup>mCh</sup>Spindly<sup>Chimera</sup> were expressed using the biGBac system as His<sub>6</sub> fusions. Baculovirus was generated in Sf9 culture and used to infect TnAO38 cells, which were grown for 72 hours at 27° C before harvesting. The pellet was then snap-frozen and stored at -80° C until purification. All <sup>mCh</sup>Spindly and Spindly constructs were purified using the same protocol. Pellets were resuspended in lysis buffer (50 mM HEPES pH 8.0, 250 mM NaCl, 50 mM imidazole, 2 mM TCEP) supplemented with protease inhibitor cocktail and lysed by sonication. The lysate was clarified by centrifugation followed by sterile filtration, and loaded onto a HisTrap HP column (Cytiva), which

was then washed with at least 10 column volumes (CV) lysis buffer. Elution was performed with lysis buffer with 300 mM imidazole. The eluate was diluted 1:5 in no salt buffer (50 mM HEPES pH 8.0, 2 mM TCEP), and applied to a 6 ml Resource Q anion exchange column (Cytiva). Elution was then performed over a 50-500 mM NaCl gradient. Fractions of the peak were analysed by SDS-PAGE and those containing the protein of interest were pooled and concentrated. The concentrated sample was loaded onto a Superdex 200 10/300 pre-equilibrated in Spindly buffer (50 mM HEPES pH 8.0, 250 mM NaCl, 2 mM TCEP). The eluate was concentrated to 10 mg/ml, flash-frozen and stored at -80° C until use. The Spindly<sup>250-275\_307-605</sup> construct was expressed in bacteria as an mCherry fusion, and the mCherry tag was removed after the SEC purification step by overnight incubation with PreScission protease purified in-house. The protease and the cleaved mCherry tag were then separated from the Spindly sample by a further run of SEC on a Superdex 200 10/300 column pre-equilibrated in Spindly buffer, and the eluate was concentrated, flash-frozen and stored at -80° C until use.

Expression and purification of Spindly<sup>1-100</sup> The N-terminal 1-100 residues of Spindly were cloned in the pET-NKI-His-3C-LIC vector (Luna-Vargas et al., 2011) for expression in E. coli BL21 (DE3) cells. Cells were grown in LB medium at 37°C to an OD<sub>600</sub> of 0.6. Overexpression was induced by adding isopropyl β- d-1-thiogalactopyranoside (IPTG) to a final concentration of 0.2 mM. Post induction, the cells were further grown for 18 hours at 18°C. For purification, cell pellet from a 2 l culture was resuspended in lysis buffer (40 mM HEPES/HCl pH 7.5, 500 mM NaCl, 10 mM Imidazole, 2 mM Tris(2-carboxyethyl)phosphine (TCEP, ThermoFisher)) supplemented with a protease inhibitor tablet (Roche) and 0.1 mM phenylmethylsulfonyl fluoride (PMSF). Cells were lysed by sonication (10 sec ON / 30 sec OFF; 70% Amplitude; 180 sec) and the lysate was centrifuged at 53000 g for 30 minutes. The supernatant was filtered through a 0.45  $\mu$ M filter (Millipore) and incubated with 1 ml of Ni-Sepharose beads (Qiagen) on a rotator for 1 hour at 4°C. The protein was eluted from the beads with elution buffer (40 mM HEPES pH 7.5, 100 mM NaCl, 500 mM Imidazole, 2 mM TCEP. Protein containing fractions were pooled together and incubated with 3 mg/ml of 3C protease (1:100 molar ratio) overnight at 4°C to cleave off the N-terminal 6x-His tag. The protein was further purified by ion-exchange chromatography using a 6ml Porous XQ column and eluted with a linear NaCl gradient (50-1000 mM). As a final purification step, the protein was loaded onto a S200 10/300 SEC column equilibrated with buffer containing 40 mM HEPES/HCl pH 7.5, 100 mM NaCl, 2 mM TCEP. The elution fractions were analyzed by SDS-PAGE and the protein containing fractions were pooled together, concentrated to 11 mg/ml and stored at -80°C.

**Expression and purification of GST-LIC2** Baculoviruses for GST-LIC2 expression were generated in Sf9 culture and used to infect TnAO38 cells, which were grown for 72 hours at 27° C before harvesting. The pellet was then snap-frozen and stored at -80° C until purification. The pellet was resuspended in Spindly buffer supplemented with protease inhibitor cocktail, and lysed by sonication. The lysate was clarified by centrifugation followed by sterile filtration, and loaded onto a GSTrap column. The column was washed with 10 CV Spindly buffer. Elution was performed in SEC buffer supplemented with 50 mM glutathione, pre-buffered to pH 8.0. The eluate was pooled and concentrated for gel filtration. Gel filtration was performed on a Superdex 200 10/300 column pre-equilibrated in Spindly buffer. The fractions of the peak were analysed by SDS-PAGE, and the ones containing the protein of interest were pooled, concentrated, snap-frozen and stored at -80° C until use.

**Expression and purification of the Dynactin pointed end** Baculoviruses for Pointed end complex expression were generated in Sf9 culture and used to infect TnAO38 cells, which were grown for 72 hours at 27° C before harvesting. The pellet was then snap-frozen and stored at -80° C until purification. The pellet was resuspended in lysis buffer supplemented with protease inhibitor cocktail and 1 mg/ml DNAse, and lysed by sonication. The lysate was clarified by centrifugation followed by sterile filtration, and loaded onto a HisTrap HP column, which was then washed with at least 10 CV lysis buffer. Elution was performed with lysis buffer with 250 mM imidazole. The eluate was diluted 1:5 in no salt buffer, and applied to a 6 ml Resource Q anion exchange column. Elution was performed over a 50-500 mM NaCl gradient. Fractions of the peak were analysed by SDS-PAGE and those containing the entire complex were pooled and concentrated. The concentrated sample was loaded onto a Superdex 200 16/60 column pre-equilibrated in Spindly buffer. The eluate was concentrated to 15 mg/ml, flash-frozen and stored at -80° C until use.

*Expression and purification of recombinant Dynein tail* Expression of the Dynein tail (residues 1-1455) construct was performed using a previously described plasmid and protocol (Schlager et al., 2014a; Urnavicius et al., 2015) . Pellets were resuspended on ice in Dynein buffer (50 mM HEPES-KOH pH 7.3, 150 mM KCl, 5 mM MgCl<sub>2</sub>, 2 mM TCEP, 0.2 mM ATP, 10% v/v glycerol) supplemented with protease inhibitor cocktail and 1 mg/ml DNAse, and lysed by sonication. The lysate was cleared by centrifugation and sterile filtration. The lysate was loaded on a 5 ml IgG column three times, and the column was washed with 20 CV Dynein buffer. Elution was performed by TEV cleavage of the ZZ tag overnight at 4° C. The cleaved protein was eluted in Dynein lysis buffer with fractionation. The eluate was pooled and concentrated for gel filtration. The concentrated sample was loaded on a Superose 6 10/300 column pre-equilibrated in Dynein buffer.

The eluate from gel filtration was pooled and concentrated to approximately 5 mg/ml, snap-frozen, and stored at  $-80^{\circ}$  C until use.

Purification of Dynactin from pig brain Pig brain Dynactin was purified essentially as previously described (Zhang et al., 2017). Pig brains were homogenised by blending at 4° C in a Waring blender, in PMEE buffer (35 mM PIPES-KOH pH 7.2, 5 mM MgSO<sub>4</sub>, 1 mM EGTA, 0.5 mM EDTA, 0.1 mM ATP, 2 mM TCEP), supplemented with 1 mg/ml DNAse and protease inhibitor cocktail. The lysate was cleared by two rounds of centrifugation, first at 30,000 g for 20 min at 2° C, then at 235,000 g for 45 min at 4° C, followed by two rounds of filtration, first through a glass fibre filter, then through a 0.45 µm filter. The cleared lysate was loaded on 300 ml SP Sepharose XL resin packed in an XK 50/30 column (Cytiva) and equilibrated in PMEE buffer. The column was washed with 4 CV PMEE. Elution was then performed with a 0 to 250 mM KCl gradient. Fractions containing Dynactin were initially identified by blotting for p150<sup>glued</sup>, pooled, and diluted 1:1 in PMEE. The diluted eluate was loaded on a MonoQ HR 16/10 column equilibrated in PMEE. The column was then washed with 10 CV PMEE. Elution was performed over a 150-350 mM KCl gradient. Dynactin eluted with a peak around 34 mS/cm conductivity. The fractions containing Dynactin were identified by SDS-PAGE, pooled, concentrated, snap-frozen, and stored at -80° C. As a final step of purification, the products of three MonoQ runs were pooled together and loaded on a Superose 6 10/300 column equilibrated in Dynactin buffer (25 mM HEPES-KOH pH 7.4, 150 mM KCl, 1 mM MgCl<sub>2</sub>, 2 mM TCEP, 0.1 mM ATP). The fractions of the peak were analysed by SDS-PAGE, and those containing Dynactin were pooled and concentrated to a final concentration of 3 mg/ml. The sample was snap-frozen and stored at -80° C until use.

*Expression and purification of recombinant human Dynactin* Recombinant human Dynactin was expressed using Expi293F cells (Invitrogen) and Expi293 expression medium (Invitrogen). Cells were freshly thawed for each expression, and passed at least three times before infection. All Expi293F cell cultures were performed at  $37^{\circ}$  C, 8% CO<sub>2</sub>, on an orbital shaker set to 125 rpm. Baculovirus for expression was made fresh for every expression. Sf9 cells were transfected with bacmid produced in EMBacY cells to produce baculovirus, which was then amplified through three 4-day rounds of amplification. The final V<sub>2</sub> virus to be used for expression was made by infecting 60 ml V<sub>2</sub> Sf9 culture supplemented with 5% FBS, and incubating it for 4 days at  $27^{\circ}$  C. The culture was then centrifuged at low speed to remove Sf9 cells, and the supernatant was decanted. 20 ml of sterile-filtered PEG solution (32% PEG6000, 400 mM NaCl, 40 mM HEPES pH 7.4) was added to the supernatant, mixed thoroughly, and the solution was allowed to precipitate overnight at 4° C in the dark. The precipitated virus was pelleted by centrifugation at

4000 rpm at 4° C for 30 minutes. The pellet was solubilized in 10 ml Expi293 expression medium prewarmed to 37° C and immediately used to infect 500 ml Expi293F culture, at a density of 3.5-5 x 10<sup>6</sup> cells per ml. After 8 h from infection, 5 ml of a 1 M stock solution of sterile sodium butyrate in PBS was added to 500 ml Expi293F culture. The expression was incubated at 37° C, 8% CO<sub>2</sub> for 48 h after infection. Cells were harvested by centrifugation at 350 g for 15 minutes, the pellet was washed with PBS, snap-frozen, and stored at -80° C until purification.

The pellet was resuspended in Dynein buffer, supplemented with 1 mg/ml DNAse and protease inhibitor cocktail, and lysed by sonication. The lysate was cleared by centrifugation, followed by sterile filtration, and loaded on a 5 ml IgG column. The flowthrough from the IgG column was further loaded three times on 4x 1 ml IgG beads in gravity flow columns. The column and the beads were washed with 10 CV each Dynein buffer. Elution was performed by TEV cleavage of the ZZ-tag overnight at 4° C. Fractions containing Dynactin were identified by SDS-PAGE, pooled and diluted 1:4 in MonoQ buffer (50 mM HEPES pH 7.3, 100 mM KCl, 5 mM MgCl<sub>2</sub>, 2 mM TCEP, 0.1 mM ATP, 10% glycerol). The diluted eluate was loaded on a MonoQ HR 16/10 column. After loading, the column was washed with 10 CV MonoQ buffer. Elution was performed over a 100-500 mM KCl gradient. Recombinant Dynactin eluted around 34 mS/cm conductivity, matching the results for pig brain Dynactin. The eluate was then concentrated in an Amicon 0.5 ml concentrator with a 100 kDa molecular mass cut-off to a final concentration of around 3-4 mg/ml. Typical yield from a 500 ml culture was in the range of 100-200 µg.

In vitro dephosphorylation and phosphorylation RZZ and Spindly were dephosphorylated using  $\lambda$ -phosphatase. Proteins were diluted to 10 µM in Spindly buffer, and  $\lambda$ -phosphatase was added to a final concentration of 500 nM, and incubated on ice for 15 minutes. Afterwards, MnCl<sub>2</sub> was added at 10 mM concentration. The reaction mixture was incubated overnight at 10° C, and loaded on a Superose 6 10/300 column equilibrated in SEC buffer. The eluate was pooled, concentrated, snap-frozen and stored at -80° C until use. Pre-dephosphorylated RZZ and Spindly were phosphorylated with the mitotic kinases CDK1:CyclinB, MPS1, and Aurora B, which were all purified in-house (Huis In 't Veld et al., 2021; Raisch et al., 2021; Sessa et al., 2005). Proteins were diluted to 10 µM in Spindly buffer, and kinases were added at 1 mM, together with 10 mM MgCl<sub>2</sub>. The reaction mixture was incubated overnight at 10° C, and loaded on a Superose 6 10/300 column

In vitro farnesylation Farnesyltransferase  $\alpha/\beta$  mutant (W102T/Y154T) was expressed and purified as previously described (Mosalaganti et al., 2017). Spindly was diluted to 100 µM in farnesylation buffer (50 mM HEPES pH 8.0, 250 mM NaCl, 10 mM MgCl<sub>2</sub>, 2 mM TCEP), and farnesyltransferase was added to a final concentration of 30 µM. Farnesyl pyrophosphate was added stepwise to a final concentration of 300 µM. The reaction mixture was incubated at RT for 6 hours, after which it was centrifuged at 16,000 g for 10 minutes to remove precipitate that formed during the reaction. The cleared reaction mixture was then loaded on a Superose 6 column equilibrated in SEC buffer to remove the farnesyltransferase. The fractions containing Spindly were identified by SDS-PAGE and pooled, concentrated, snap-frozen, and stored at -80 °C until use.

Analytical Size Exclusion Chromatography (SEC) Analytical SEC was performed under isocratic conditions at 4° C in Spindly buffer on an ÄKTAmicro system. Elution profiles were obtained by monitoring absorbance at 280 nm wavelength. 50 µl fractions were collected and analysed by SDS-PAGE. Complex formation assays were performed by mixing the samples at the indicated concentrations in 60 µl Spindly buffer and incubating them for at least 1 h on ice before the SEC assay was performed.

*Mass Photometry (MP)* Mass photometry experiments were performed essentially as described in Sonn-Segev et al., 2020. Standard microscope coverslips were cleaned with MilliQ water and isopropanol, and dried under an air stream. Silicon buffer gaskets were attached to the glass slides, and the slides were then mounted on a Refeyn TwoMP mass photometer (Refeyn Ltd). For measurement, Spindly construct samples were diluted to 100 nM in Spindly buffer immediately before measurement. The gasket was filled with Spindly buffer and the focal plane was automatically estimated. Proteins were diluted 1:10 into the buffer-filled gasket, to a final concentration of 10 nM. A 60 second movie was then recorded through AcquireMP (Refeyn Ltd). Data was processed using the DiscoverMP program. Contrast-to-mass calibration was performed with several known-mass proteins. Mass distributions were plotted with DiscoverMP and mean mass peaks determined by Gaussian fitting.

*Analytical Ultracentrifugation (AUC)* Sedimentation velocity analytical ultracentrifugation was performed at 42,000 rpm at 20° C in an An-60 Ti rotor (Beckman Coulter), using standard charcoal double-sector centrepieces. Protein samples were diluted in buffer containing 40 mM HEPES pH 8.0, 200 mM NaCl, and 2 mM TCEP. Approximately 300 radial absorbance scans at 587 nm were collected per each run, with a time interval of 1 minute. Buffer density and viscosity, as well as the protein partial specific volume, was estimated using the program SEDN/TERP. Sedimentation peaks were identified through an analysis using the SEDFIT suite (Brown and Schuck, 2006), in

terms of continuous distribution function of sedimentation coefficients (c(S)). Frictional ratios were manually set to match the sedimentation coefficient of the main peak to the theoretical mass of the sample, at the stoichiometry predicted by mass photometry data. A fitting run was then performed within SEDFIT, and the resulting estimations for frictional ratio and Stokes' radius were collected as output. Figures were produced with the program GUSSI.

*Size-exclusion chromatography-multi angle light scattering (SEC-MALS)* SEC-MALS was performed using a Heleos II 18 angle light scattering instrument (Wyatt), which was coupled to an Optilab rEX online refractive index detector (Wyatt). 100 μl of pig- or human dynactin (0.15 mg/ml) were loaded onto a TSKgel G4000SWXL column with a TSKgel SWXL guard column (TOSOH Bioscience) equilibrated in GF150 buffer (25 mM HEPES pH 7.4, 150 mM KCl, 1 mM MgCl2, 5 mM DTT, 0.1 mM ATP) at room temperature. The column outlet was directly connected to the light scattering instrument and the refractive index detector. Data collection and determination of molecular weight was performed using the ASTRA 5.3.4 software (Wyatt).

Negative stain electron microscopy (EM) For negative stain EM, 400-square-mesh copper grids (Electron Microscopy Sciences) were plasma cleaned, and 3 µl of protein ( $\approx$ 100 nM in GF150 buffer) were applied to the grid. 20 µl of 2 % (w/v) uranyl acetate were added to the sample, and excessive stain was removed with a filter paper. Micrographs were recorded on an FEI Tecnai G2 Spirit transmission (120 kV) transmission electron microscope equipped with a Gatan Ultrascan 1000 XP CCD detector. Image acquisition was performed at 26000x nominal magnification and 1.5 µm underfocus. For 2D classification of recombinant human dynactin, approximately 5000 particles were picked semi-automatically using EMAN2 (Tang et al., 2007) and 2D-classified using RELION 2.1 (Scheres, 2012).

DSBU and UV crosslinking For DSBU crosslinking, all proteins in the assay were diluted to 5  $\mu$ M in Spindly buffer. DSBU was added at a final concentration of 3 mM, and the reaction was incubated for 1 hour at RT. At the end of the reaction, TRIS-HCl pH 8.0 at a final concentration of 100 mM was added and the solution was thoroughly mixed to quench the reaction. Successful crosslinking was evaluated by SDS-PAGE analysis. For UV crosslinking, the BPA-containing Spindly mutants were diluted in Spindly buffer to a final concentration of 5  $\mu$ M. The samples were irradiated with LED UV light at 365 nm wavelength for 15 minutes to induce complete crosslinking. Successful crosslinking was evaluated by SDS-PAGE analysis.

**Processing for LC/MS** Crosslinked samples were precipitated by dilution into 4 volumes of acetone pre-cooled to -20° C, followed by overnight incubation at -20° C. Pellets were resuspended and denatured in 8 M urea, 1 mM DTT, and alkylated with 5.5 mM chloroacetamide. The

concentration of urea was then lowered to 4 M by dilution in 20 mM ammonium bicarbonate, pH 8.0, and the protein solution was then digested overnight at room temperature with Trypsin. Digestion was stopped by adding TFA to a final concentration of 0.2%. Samples were subjected to SEC on a Superdex 30 Increase 3.2/300 (Cytiva), followed by purification on a tC18 Sep-Pak column (50 mg, Waters) as previously reported (Pan et al., 2018). For the <sup>mCh</sup>Spindly<sup>Y26BPA</sup> and <sup>mCh</sup>Spindly<sup>Q29BPA</sup>, the SEC step was omitted.

*LC-MS/MS analysis* LC-MS/MS analysis was performed as described in Pan et al., 2018. Data analysis was performed for the DSBU-crosslinked samples in MeroX 2.0.0.8. The analysis of Bpacrosslinked samples was performed in MeroX 2.0.1.4. An artificial amino acid "Z" was added with the composition C16H13NO2 (mass 251.09463), and Bpa was defined as a crosslinker with parameters: 'Specificity site 1:Z', 'Specificity Site 2: [ABCDEFGHIKLMPQRSTVWY]', and 'Maximum C $\alpha$ -C $\alpha$  -distance: 30 Å'. For the Spindly<sup>F258BPA</sup> sample, the analysis was performed within MeroX 2.0.1.4 with a False Discovery Rate (FDR) of 50%. Crosslinks were then exported into XiView (Graham et al., 2019), and crosslinks with a score lower than 50 were filtered out. Visual representations of crosslinking data were produced using the xVis website (https://xvis.genzentrum.lmu.de (Grimm et al., 2015)), and then edited to produce the final figures.

Molecular modelling and bioinformatical methods AlphaFold 2 (AF2) was used for all molecular modelling (Jumper et al., 2021). For the Spindly<sup>1-275</sup>, Spindly<sup>1-309</sup>, and Spindly<sup>1-440</sup> constructs, the original version of AlphaFold Multimer was used (Evans et al., 2021). For all other constructs the ColabFold version of AlphaFold 2 was used (Mirdita et al., 2021). A general feature of AF2 Multimer is the higher sensitivity to intra- and intermolecular interactions compared to ColabFold and thus a tendency to predict extended protein structures as a more or less compact model, making it difficult to distinguish artificial contacts from structurally significant ones. This problem is worse for fragments containing long, disordered stretches. Therefore, the disordered C-terminal region of Spindly (441-605) was not included in the predicted sequences, unless otherwise specified. In absence of clear predicted intramolecular contacts from the PAE plots, the models of the adapters shown in Figure 2 - Supplement 1 (obtained with ColabFold) were manually "unfolded" with PyMol and Coot to extend them and make them comparable. To determine the putative interactions in the autoinhibited Spindly fragments, the pLDDT scores of the models and especially the "predicted alignment errors" (PAE) were scrutinized to see which parts of the models were predicted to have defined relative orientations to each other (Figure 1 – Supplement 2). Also, some features, like the "kink" at Spindly residues 155-156, were very consistently predicted in almost all models. Other features, like predicting extended conformations of e.g. the Spindly N-terminal

coiled coil, were more or less frequent, depending on the presence of the interacting, autoinhibitory regions, which was interpreted as being indicative of the strength of the co-evolutionary signal present in those regions.

Crystallographic structure of Spindly<sup>1-100</sup> Crystallization trials were set up in MRC 2-well sitting drop plates by mixing the protein and the reservoir solution in a 1:1 ratio. Initial crystallization hits were optimized and final crystals were obtained in a condition containing Bis-Tris Propane/HCl pH 6.5, 19% PEG3350 and 0.2M KSCN. For data collection, crystals were cryoprotected in mother liquor containing 20% glycerol prior to flash cooling in liquid  $N_2$ . Data for native crystals were collected at the MASSIF-1 beamline (Bowler et al., 2015) at the ESRF, Grenoble. For SelenoMethionine protein production, E. coli BL21 DE3 cells were grown in ready to use SelenoMethionine medium (Molecular Dimensions) and purified using the same protocol as for the native protein. SeMet crystals could be reproduced in similar crystallization conditions, and anomalous data were collected at I04 beamline of Diamond Light Source. Details for all steps are available in Table S1. Data processing was done with XDS (Kabsch, 2010) and the structure was solved by Se-SAD using the CRANK2 (Pannu et al., 2011) pipeline from the CCP4 (Winn et al., 2011) software suite. The initial model contained 206 residues built in 8 fragments and clearly showed the parallel dimeric coiled coil. Further model building was performed in Coot (Emsley et al., 2010) and subsequent refinement cycles were carried out in REFMAC (Murshudov et al., 2011) and PDB-REDO (Joosten et al., 2014) using non-crystallographic and jelly-body restraints. The structure was refined to 2.8 Å resolution to an R<sub>free</sub> of 29%. All residues are in the favorable regions of the Ramachandran plot and the structure is in the 97th percentile of Molprobity (Williams et al., 2018).

*Cell culture and drug treatment* HeLa cells were grown in Dulbecco's Modified Eagle's Medium (DMEM; PAN Biotech) supplemented with 10 % tetracycline-free FBS (PAN Biotech), and L-Glutamine (PAN Biotech). Cells were grown at 37°C in the presence of 5 % CO2.

*Cell transfection and electroporation* Depletion of endogenous Spindly was achieved through reverse transfection with 50 nM Spindly siRNA (5'-GAAAGGGUCUCAAACUGAA-3' obtained from Sigma-Aldrich, (Gassmann et al., 2010)) for 48 hours with RNAiMAX (Invitrogen, Carlsbad, California, United States). For rescue experiments, 24 hours after Spindly depletion, we electroporated recombinant Spindly constructs labeled with an N-terminal mCherry, at a concentration of 7  $\mu$ M in the electroporation slurry (as previously described in (Alex et al., 2019)) (Neon Transfection System, Thermo Fisher). Control cells were electroporated with mCherry. Following an 8 hour recovery, cells were treated with 9  $\mu$ M RO3306 (Calbiochem) for 15 hours.

Subsequently, cells were released into mitosis in presence of  $3.3 \ \mu\text{M}$  Nocodazole (Sigma) for 1 hour.

**Immunofluorescence** Cells were grown on coverslips pre-coated with Poly-L-lysine (Sigma-Aldrich). Cells were fixated with 4% PFA in PHEM for 10 minutes. Subsequently, the cells were permeabilized for 10 minuets with PHEM supplemented with 0.5% Triton-X100 (PHEM-T). After blocking with 5% boiled goat serum (BGS) in PHEM buffer, the cells were incubated for 2 hours at room temperature with the following primary antibodies: CENP-C (guinea pig, MBL, #PD030, 1:1000), Dynactin-p150 (mouse, BD Trans. Lab., #610473, 1:400), Spindly (rabbit, Bethyl, A301-354A, 1:1000) diluted in 2.5 % BGS-PHEM-T (PHEM supplemented with 0.1% Triton-X100). Subsequently, cells were incubated for 1 hour at room temperature with the following secondary antibodies: Goat anti-mouse Alexa Fluor 488 (Invitrogen, A11001), donkey anti-rabbit Rhodamine Red (Jackson Immuno Research 711-295-152), goat anti-guinea pig Alexa Fluor 647 (Invitrogen, A-21450). All washing steps were performed with PHEM-T buffer. DNA was stained with 0.5 µg/ml DAPI (Serva) and Mowiol (Calbiochem) was used as mounting media.

*Cell imaging* Cells were imaged at room temperature using a spinning disk confocal device on the 3i Marianas system equipped with an Axio Observer Z1 microscope (Zeiss), a CSU-X1 confocal scanner unit (Yokogawa Electric Corporation, Tokyo, Japan),  $100 \times /1.4$ NA Oil Objectives (Zeiss), and Orca Flash 4.0 sCMOS Camera (Hamamatsu). Images were acquired as z sections at 0.27 µm (using Slidebook Software 6 from Intelligent Imaging Innovations or using LCS 3D software from Leica). Images were converted into maximal intensity projections, exported, and converted into 16-bit TIFF files. Automatic quantification of single kinetochore signals was performed using the software Fiji with background subtraction. Measurements were exported in Excel (Microsoft) and graphed with GraphPad Prism 9.0 (GraphPad Software). Statistical analysis was performed with a nonparametric t-test comparing two unpaired groups (Mann-Whitney test). Symbols indicate: n.s. = p > 0.05, \* = p  $\leq 0.05$ , \*\* = p  $\leq 0.01$ , \*\*\* = p  $\leq 0.001$ , \*\*\*\* = p  $\leq 0.001$ 

*GST-LIC2 pulldown assay* GST and GST-LIC baits were added to 10  $\mu$ L GSH beads (Serva) at a final concentration of 4  $\mu$ M in Spindly buffer, in Pierce micro-spin columns (Thermo Scientific), and incubated at 4° C for 1 h. The unbound baits were removed by centrifugation, and the beads were washed twice. The Spindly preys were added at a final concentration of 8  $\mu$ M in Spindly buffer. The bait-bound beads were incubated with the preys at 4° C for 1 h, after which the supernatant was removed, and the beads washed twice. After the final wash, both prey and bait were eluted in Spindly buffer supplemented with 50 mM glutathione. The inputs and eluates were analysed by SDS-PAGE, and the unstained gels were imaged in the 546 nm channel. The signal

from the main band was quantified using ImageLab (Biorad), in comparison to the signal of the SpindlyFL band. The gels were then stained with Coomassie and imaged.

**RZZS filaments** RZZS filaments were formed and imaged essentially as described in (Raisch et al., 2021). 4  $\mu$ M RZZ was incubated with 8  $\mu$ M prefarnesylated Spindly in presence of 1  $\mu$ M MPS1, in M-buffer (50 mM HEPES pH 7.5, 100 mM NaCl, 1 mM MgCl2, 2 mM ATP, 1 mM TCEP) at room temperature overnight. Flow chambers were assembled by placing two parallel strips of double-sided tape onto a glass slide, with a standard coverslip on top, creating a chamber of 5 to 10  $\mu$ l volume. The filament sample was diluted to 0.5  $\mu$ m and loaded into the flow chamber. Imaging was performed on a 3i Marianas system at 100X magnification in the 561 nm channel. Sample images were acquired as 5-stacks of z-sections at 0.27  $\mu$ m, converted into maximal intensity projections, and processed in Fiji (Schindelin et al., 2012).

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## Figure 1. Spindly is a folded adaptor

(A) Schematic representation of the organisation of the coiled-coil regions of Spindly, and relevant coiled coil prediction (COILS, ExPaSy suite). (**B**) Organisation of the kinetochore and corona. (**C**) Results of mass photometry measurement of a sample of mCherry-tagged Spindly (<sup>mCh</sup>Spindly<sup>FL</sup>). The measurements are consistent with Spindly being a dimer in solution, even at very low concentration (10 nM). (D) Crystallographic structure of a Spindly<sup>1-100</sup> construct. Only residues 2-97 were visible in the electron density. (E) Structure of the Spindly CC1 box (orange) and surrounding sequence. (F) Structure of the Spindly CC1 SCM (also known as CC2 box) (red) and surrounding sequence. (G) Summary of XL-MS data reporting Spindly intramolecular crosslinks. For ease of viewing, only crosslinks detected  $\geq 3$  times and involving sites  $\geq 40$  residues apart are depicted. See also Table S2 for a detailed list of all crosslinks. (H) Summary of XL-MS data reporting Spindly intramolecular crosslinks found through amber codon suppression experiments. Red arrows indicate the sites where the BPA residues were introduced. Crosslinking results from three mutants are merged: Y26BPA (orange), Q29BPA (cyan), and F258BPA (magenta). A few crosslinks identified between the BPA residues and the mCherry tag were considered spurious and not displayed. See also Table S2 for a detailed list of crosslinks. (I) AlphaFold2 Multimer prediction of Spindly structure. The CC1 box is in orange, the CC1 SCM in red, the Spindly motif in cyan. The C-terminal unstructured tail of Spindly (aa 440-605) was omitted from the model due to the very low confidence index (pLDDT) of the prediction for this region (unpublished results).

## Figure 2. Spindly autoinhibition prevents its interaction with Dynein-Dynactin

(A) AlphaFold2 Multimer was used to predict a model of BicD2. The flexible region between 270 and 332 has very low reliability and has therefore been artificially linearized for visualization purposes (see Methods). The tail region has been omitted due to limited reliability of the predictions. The vertical like with an asterisk marks the accumulation of unknown contaminants in the upper part of the gel. (B) AlphaFold2 Multimer model of Spindly<sup>1-275</sup>. (C) AlphaFold2 Multimer model of Spindly<sup>1-309</sup>. The PAE plots and pLDDT scores are displayed in Figure 1 – Supplement 2. (D-G) Analytical size-exclusion chromatography on a G4000SWXL column (void volume  $\approx 5.5$  ml) to assess complex formation between Dynein tail (green), pig brain Dynactin (grey), and an adaptor - cargo/adaptor complex (blue), with the complex run shown in red. (D) BicD2<sup>1-400</sup>; (E) Spindly<sup>FL</sup>; (F) RZZ-Spindly<sup>FL</sup> treated with  $\lambda$ -phosphatase; (G) RZZ-Spindly<sup>FL</sup>

tail and Dynactin controls are both shared between panels C and D, and panels E and F. In all shown SEC experiments, Spindly<sup>FL</sup> was farnsylated.

#### Figure 3. Spindly autoinhibition is relieved by N- and C-terminal deletions

(A) Schematic representation of the pointed end complex in the context of Dynactin, and SDS-PAGE of its chromatographic peak in gel filtration. (**B-G**) Analytical SEC binding assays between the Dynactin pointed end (brown) and Spindly constructs. The complex run is always represented with a continuous line, the Spindly construct with a dashed line. (**B**) <sup>mCh</sup>Spindly (purple); (**C**) Spindly<sup>1-275</sup> (red); (**D**) Spindly<sup>1-250</sup> (red); (**E**) <sup>mCh</sup>Spindly<sup>Δ276-306</sup> (blue); (**F**) Spindly<sup>156-275</sup> (purple); (**G**) <sup>mCh</sup>Spindly<sup>22-605</sup> (blue); <sup>mCh</sup>Spindly<sup>33-605</sup> (red). The control gels with the pointed-end (PE) alone are shared between panels B and G, C and F, D and Figure 3 – Supplement 1A.

#### Figure 4. Spindly autoinhibition involves a direct interaction between N- and C-terminal regions

(A) Analytical SEC elution profile and SDS-PAGE analysis for interaction assays between the Spindly N-terminal and C-terminal domains. Spindly<sup>1-250</sup> (red, dashed) interacts with Spindly<sup>250-605</sup> (alone: purple, dashed; complex: red, continuous), but not with Spindly<sup>250-275\_307-605</sup> (alone: green, dashed; complex: green, continuous). Spindly<sup>51-250</sup> (blue, dashed) does not interact with Spindly<sup>250-605</sup> (complex: blue, continuous). This result shows that the Spindly regions 1-51 and 276-306 are required for Spindly folding. (**B**) schematic representation of Spindly constructs referred to in panels C,D and E. (**C**) Mass photometry results for the constructs in panel B. (**D**) Analytical ultracentrifugation results for the constructs in panel B. The smaller sedimentation coefficient indicates higher drag, which is caused by an increased Stokes' radius. (**E**) Data table from hydrodynamic and mass photometry results in panels C-D. The Stokes' radius and frictional ratio were estimated from the AUC-measured sedimentation coefficient and from the theoretical molecular weights.

#### Figure 5. Point mutations relieve Spindly autoinhibition

(A-C, H) Analytical SEC analyses on a Superdex 200 5/150 column to assess complex formation between the Dynactin pointed end (brown) and various Spindly constructs. The complex run is always represented with a continuous line, the Spindly construct with a dashed line. (A)  $^{mCh}$ Spindly<sup>CC2\*</sup> (red); (B)  $^{mCh}$ Spindly<sup>SM\*-CC2\*</sup> (blue); (C)  $^{mCh}$ Spindly<sup> $\Delta RV$ </sup> (purple). (H)  $^{mCh}$ Spindly<sup>Chimera</sup>

(green). (**D**) Mass photometry results for <sup>mCh</sup>Spindly<sup>CC2\*</sup> (red) and <sup>mCh</sup>Spindly<sup>ARV</sup> (purple). The main peaks' "shoulders" are consistent with minor sample degradation. (**E**) AUC profile of <sup>mCh</sup>Spindly<sup>CC2\*</sup> (red), <sup>mCh</sup>Spindly<sup>ARV</sup> (purple), and <sup>mCh</sup>Spindly<sup>WT</sup> (green). (**F**) schematic representation of the <sup>mCh</sup>Spindly<sup>Chimera</sup>. (**G**) Spinning-disk confocal fluorescence microscopy-based filamentation assay at 561 nm shows the indicated <sup>mCh</sup>RZZS species (4  $\mu$ M RZZ, 8  $\mu$ M Spindly) form filaments when incubated at 20°C with MPS1 kinase. Scale bar: 5  $\mu$ m. (**I**) SDS-PAGE analysis of pulldown assay with either GST or GST-tagged LIC2 as bait, and mCherry-tagged Spindly as prey. Coomassie staining and fluorescent signal in the red channel are displayed. Asterisks mark contaminants or degradation products. (**J**) quantification of the <sup>mCh</sup>Spindly fluorescent signal and SD calculated from three technical repeats. The PE alone controls in panels A and C are shared with the control in **Figure 3E**.

#### Figure 6. complex formation assay between Dynein-Dynactin and Spindly mutants

(A) Elution profiles and SDS-PAGE of complex formation assays between Dynein tail, recombinant Dynactin and Spindly constructs. Experiment run on a Superose 6 5/150 column, in stoichiometric conditions. Only selected gels are displayed. The gray dotted box indicates the fraction loaded in the SDS-PAGE shown in panel B. (B) Comparison of the fractions of the expected DDS complex peak shown in panel A. A minus sign indicates adaptor-only runs, a plus indicates full complex runs. The arrow points at the expected position of <sup>mCh</sup>Spindly. (C-D) Analytical SEC experiments on a Superose 6 5/150 column to assess complex formation (blue) between the RZZ complex (red) and the indicated Spindly constructs (green). (C) <sup>mCh</sup>Spindly<sup>Chimera</sup>. (D) <sup>mCh</sup>Spindly<sup>CC2\*</sup>.

# Figure 7. Kinetochore levels of Dynactin in presence of Spindly mutants

(A) Representative images showing the effects of a knockdown of the endogenous Spindly in HeLa cells on Dynactin recruitment monitored through the p150<sup>glued</sup> subunit. RNAi treatment was performed for 48 h with 50 nM siRNA (see Figure 7 – Supplement 1A). Before fixation, cells were synchronized in G2 phase with 9  $\mu$ M RO3306 for 16 h and then released into mitosis. Subsequently, cells were immediately treated with 3.3  $\mu$ M nocodazole for an additional hour. CENP-C was used to visualize kinetochores and DAPI to stain DNA. Scale bar here and in panel F: 5  $\mu$ m. (B) Quantification of residual Spindly levels at kinetochores. A representative image is shown in Figure 7 – Supplement 1B. (C) Quantification of kinetochore levels of the indicated

electroporated <sup>mCh</sup>Spindly proteins. (**D**) Kinetochore levels of Dynactin in cells depleted of endogenous Spindly and electroporated with the indicated Spindly proteins. (**E**) Least square linear fitting through the distribution of data points reporting for each kinetochore the CENP-Cnormalized <sup>mCh</sup>Spindly intensity on the X-axis and the CENP-C normalized p150<sup>glued</sup> intensity on the Y-axis. The individual distributions are shown in Figure 7 – Supplement 1C-G. (**F**) Electroporated <sup>mCh</sup>Spindly<sup>Chimera</sup> is observed forming polymers in Spindly-depleted cells in interphase, causing ectopic recruitment of p150<sup>glued</sup>. (**G**) Spinning-disk confocal fluorescence microscopy-based filamentation assay at 561 nm with the indicated <sup>mCh</sup>RZZS species (4  $\mu$ M RZZ, 8  $\mu$ M Spindly) at 20°C in presence of MPS1 kinase. Scale bar: 5  $\mu$ m. (**H**) Model for the activation of Spindly.

# Figure 1 – Supplement 1. Additional analyses of Spindly motifs and their influence on Spindly conformation

(A) Multiple sequence alignment of the first part of the CC1 region of the indicated adaptors containing the CC1 box and the CC1 SCM (or CC2 box). Hs, Homo sapiens; Mf, Macaca fascicularis; Mm, Mus musculus; Gg, Gallul gallus; Xt, Xenopus tropicalis; Ce, Caenorhabditis elegans. (B) Multiple sequence alignment of the Spindly motif. (C) Cartoon model of PDB ID 6PSE (Lee et al., 2020) showing the mode of binding of a LIC peptide (grey) to the CC1 box (yelloworange). (D) ColabFold prediction model of the Spindly CC1:LIC peptide complex. Coloring as in panel C. (E) Coiled-coil propensity was measured with the COILS program within ExPasY suite (Duvaud et al., 2021) and displayed for all indicated adaptors from the first residue of the CC1 box (see panel A). The coiled-coil propensity for Spindly has a deep that corresponds to a 2-residue insertion shown in panel F. (F) Multiple sequence alignment of the region of CC1 around the 2-residue insertion in Spindly that causes a deep in the coiled-coil prediction profile (see panel E). (G) SDS-PAGE documenting crosslinking of the full-length Spindly proteins with DSBU. (H) SDS-PAGE documenting crosslinking of the Spindly<sup>1-440</sup> proteins with DSBU. Panel G and H were obtained from the same original gel and the marker lane is the same in the two panels. (I) Summary of XL-MS data reporting Spindly intramolecular crosslinks. for ease of viewing, only crosslinks detected  $\geq 3$  times and involving sites  $\geq 40$  residues apart are depicted. See also Table S2. (J-K) Coomassie-stained SDS-PAGE gels documenting crosslinking of the indicated BPA mutants upon treatment with UV light. (L) Multiple sequence alignment of the indicated adaptors in the main region targeted by Q29<sup>BPA</sup> and Y26<sup>BPA</sup>. The CC2\* mutant discussed in the text

is the charge reversal mutant (E-E) at the two indicated positively charged residues (R295 and K297 in human Spindly).

## Figure 1 – Supplement 2. PAE plots and pLDDT scores

(A-C) Predicted Alignment Error (PAE) plots (left) and per-residue confidence score (pLDDT) of the three Spindly models discussed in the main text: (A) Spindly<sup>1-275</sup>. (B) Spindly<sup>1-309</sup>. (C) Spindly<sup>1-<sup>440</sup>. The pLDDT scores are displayed on the AF2 Multimer predictions of Spindly shown on the right (Blue: high confidence; Red: low confidence). The models are shown, with the same orientation, in Figure 1 and Figure 2. The PAE matrices refer to models of Spindly dimers, and correspondingly numbering of residues on the left and bottom of the plot is the number of residues in each chain multiplied by 2, and the second chain is plotted directly following the first. The parallel coiled-coils give rise to off-diagonal signals (blue) parallel to the main diagonal. Besides straight models like the one shown, a few predicted models of Spindly<sup>1-275</sup> also showed a foldedback conformation. However, there are no additional off-diagonal signals for the Spindly<sup>1-275</sup> construct in the PAE plots, suggesting that even if Spindly<sup>1-275</sup> explores folded-back conformations, these are not stable. Off-diagonal signals perpendicular to the main diagonal are instead clearly visible in the Spindly<sup>1-309</sup> and Spindly<sup>1-440</sup> constructs, consistent with a folded back conformation. A predicted folded conformation of Spindly<sup>1-440</sup> was also observed with orthologous sequences (unpublished results).</sup>

# Figure 2 – Supplement 1. Gallery of AF2 predictions of the structure of representative adaptors

(A) AF2 ColabFold models of the indicated adaptors. The scissor symbol indicates that the displayed cartoon models were truncated after the CC2 region. SM is the Spindly motif. The length of CC1 coiled-coil is indicated. As indicated in the Methods section, AF2 and variants can predict different quaternary structures for the adaptors, including trimeric or tetrameric coiled coil formation, if three or four chains respectively are used as input. However, with trimers or tetramers, the PAE values estimated between the same positions on different protomers are high to very high (i.e. insignificant), indicating that the predictions are less likely to be accurate (unpublished results). We only present predictions of dimers here, as there is experimental evidence for several of them that their active conformation is the dimer (Isabet et al., 2009; Kelkar et al., 2000; Lee et al., 2018; Urnavicius et al., 2015; Wu et al., 2005). For clarity, models were straightened as discussed in Methods. Even if Spindly is predicted to adopt a closed conformation

when CC2 is present (see main text), here for comparison we show the extended open conformation expected to bind DD. The model of the DD complex with BicD2 (PDB ID 6F3A) (Urnavicius et al., 2018) shows that the length of the experimentally modelled BicD2 coiled-coil is approximately identical to the length of CC1 predicted by AF2 for many of the displayed adaptors. (**B**) Coomassie-stained SDS-PAGE and ProQ diamond staining of phosphorylated and dephosphorylated RZZ and Spindly. Samples were initially dephosphorylated with  $\lambda$ -PPase (samples indicated as 'D') and later re-phosphorylated with the mitotic kinases CDK1/CyclinB, MPS1, and Aurora B (samples indicated as 'P').

## Figure 3 – Supplement 1. Additional analyses of the Spindly N-terminal autoinhibitory region

(**A-D**) Additional analytical SEC interaction assays between the Dynactin pointed end (brown) and the indicated Spindly constructs. The complex run is always represented with a continuous line, the Spindly construct with a dashed line. (**A**) Spindly<sup>1-440</sup> (blue); Spindly<sup>1-354</sup> (green); (**B**) <sup>mCh</sup>Spindly<sup>76-605</sup> (blue); (**C**) <sup>mCh</sup>Spindly<sup> $\Delta$ 26-28</sup> (green); (**D**) <sup>mCh</sup>Spindly<sup> $\Delta$ 26-32</sup> (blue). The PE alone control in panel A is shared with Figure 3D, between panels C and D, and between panel B and panels Figure 5 – Supplement 1A-B.

# Figure 5 – Supplement 1. Additional characterization of Spindly binding to PE

(**A**) Size-exclusion chromatography (SEC) separation of <sup>GST</sup>Spindly (dotted green line), the pointed end (PE) complex (brown line), and their mixture (continuous green line). (**B**) SEC separation of <sup>GST</sup>Spindly<sup>CC2\*</sup> (dotted green line), the pointed end (PE) complex (brown line), and their mixture (continuous green line). The PE control is shared between the two shown experiments and with Figure 3 – Supplement 1B. (**C**) The AF2 ColabFold model of the BicD2-Spindly chimera shows CC1 is continuous. (**D**) Spinning-disk confocal fluorescence microscopy-based filamentation assay at 561 nm with the indicated <sup>mCh</sup>RZZS species (4 μM RZZ, 8 μM Spindly) at 20°C in absence of MPS1 kinase. Scale bar: 5 μm.

#### Figure 6 – Supplement 1. Recombinant human Dynactin

(A) Map of the Dynactin expression plasmid. Individual subunits are labelled according to list below. CMV promoters and enhancers are labelled in yellow. PolyA signals are labelled in light blue. (B) Comparison of SDS-PAGE of pig brain Dynactin and recombinant human Dynactin, in

Coomassie staining. (**C**) SEC-MALS analysis of human (purple) and pig brain (red). Pig brain Dynactin contains a mix of the p150 isoforms p150<sup>glued</sup> and p135, yielding two different expected masses. (**D**-**E**) SEC assays on a Superose 6 5/150 column to assess binding of Dynein tail, Dynactin, and BicD2<sup>1-400</sup> with pig brain (**D**) and human (**E**) Dynactin. The Dynein tail and BicD2<sup>1-400</sup> controls are shared between the two experiments. (**F**) 2D class averages from negative stain imaging of recombinant human Dynactin.

#### Figure 7 - Supplement 1. Additional data on kinetochore levels of Dynactin with Spindly mutants

(A) Schematic of the RNAi and complementation by electroporation with recombinant proteins. (B) Representative images of RNAi control cells and cells depleted of Spindly by RNAi. C-G) Least square fitting through the distribution of data points reporting for each kinetochore the CENP-C-normalized <sup>mCh</sup>Spindly intensity on the X-axis and the CENP-C normalized p150<sup>glued</sup> intensity on the Y-axis. (C) Fit curves with data points. (D) Individual best fit for <sup>mCh</sup>Spindly<sup>FL</sup>. (E) Individual best fit for <sup>mCh</sup>Spindly<sup>CC2\*</sup>. (F) Individual best fit for <sup>mCh</sup>Spindly<sup>CC2\*</sup>. (G) Individual best fit for <sup>mCh</sup>Spindly<sup>33-605</sup>.



Figure 1







Figure 3



Figure 4



Figure 5



Figure 6



Figure 7

# Table S1 Crystallographic data

Dataset	Spindly <sup>1-100</sup>	SeMet-Spindly <sup>1-100</sup>	
Spacegroup	P 3 <sub>2</sub> 2 1	P 1 2 <sub>1</sub> 1	
a=b, c (Å)	112.51, 49.64	112.33, 48.65	
Wavelength (Å)	0.9999	0.91587	
Resolution (Å)	48.8 - 2.8 (2.95 - 2.80)	97.3 - 3.4 (3.58 - 3.4)	
R <sub>merge</sub>	0.079 (1.558)	0.139 (0.769)	
R <sub>pim</sub>	0.050 (0.970)	0.026 (0.136)	
$I/\sigma(I)$	10.8 (1.5)	15.0 (4.9)	
$CC_{1/2}$	0.99 (0.72)	0.99 (0.99)	
Completeness(%)	99.3 (99.6)	100 (100)	
Redundancy	5.4 (5.6)	33.2 (33.0)	
Refinement		Structure determination (SAD) - CR	ANK2
Resolution (Å)	2.8	Phasing	
Reflections (work, test)	9068, 342	CFOM substructure	69.4
R <sub>work</sub> / R <sub>free</sub>	0.247 / 0.290	Se sites used	18
		FOM Phasing	0 2031
<u>No. atoms</u>			0.2031
Protein (chain A, B)	1570, 1589	FOM Density Modification	0.3764
B factors (Å <sup>2</sup> )		Automated Model Building	
Protein (chain A B)	751 734	B from	0 3287
rotein (enan 11, D)	7511,7511	1 circe	0.9207
R.m.s deviations (RMSZ score)		Built Residues	206
Bond lengths	0.0149 (0.564)	Built Fragments	8
Bond angles	1.831 (0.908)	Docked Residues (%)	72.8
Favorable/Allowed/Outliers	189/0		
Molprobity score (%ile)	2.93 (97)		



Figure 1 – Supplement 1



Figure 1 – Supplement 2



Figure 2 – Supplement 1





Figure 3 – Supplement 1







Figure 6 – Supplement 1



Peptide 1	AbsPos1		Peptide 2	AbsPos2	Score
AAQxGLQLVI		26	VHMEGSVN(	-207	72
AAQxGLQLVI		26	HPADIPDYLK	-163	77
AAQxGLQLVI		26	AKKPVQLPG/	-51	81
AAQxGLQLVI		26	TMGWEASSI	-99	93
AAQxGLQLVI		26	EEVNELQYR(	198	95
AAQxGLQLVI		26	VMNFEDGG	-133	99
AAQxGLQLVI		26	NLYDSmESKI	361	109
AAQxGLQLVI		26	HPADIPDYLK	-160	114
AAQxGLQLVI		26	LQIATLLQMK	300	118
AAQxGLQLVI		26	LQIATLLQMK	303	123
AAQxGLQLVI		26	VHMEGSVN	-204	124
AAQxGLQLVI		26	NLYDSMESKI	360	124
LKAAQxGLQL		26	MLESLSBEBE	74	129
AAQxGLQLVI		26	SRMLESLSBE	73	139
AAQxGLQLVI		26	LQIATLLQMK	306	149
AAQxGLQLVI		26	AAQxGLQLVI	30	163
AAQxGLQLVI		26	AAQxGLQLVI	31	163
LKAAQxGLQL		26	AAQxGLQLVI	29	170
AAQxGLQLVI		26	AAQxGLQLVI	28	172

Peptide 1	AbsPos1		Peptide 2	AbsPos2	Score
[AAQYGLxLV		29	[NLYDSMESK	353	82
[AAQYGLxLV		29	[AKKPVQLPG	-51	86
[LKAAQYGLx		29	[MLESLSBEB	73	95
[AAQYGLxLV		29	[VMNFEDGG	-143	97
[AAQYGLxLV		29	[EEVNELQYR	194	107
[AAQYGLxLV		29	[HPADIPDYLk	-160	131
[AAQYGLxLV		29	[VMNFEDGG	-131	135
[AAQYGLxLV		29	[TMGWEASS	-99	135
[AAQYGLxLV		29	[MKLQIATLLC	297	135
[AAQYGLxLV		29	[MKLQIATLLC	298	144
[AAQYGLxLV		29	[LQIATLLQM	299	146
[AAQYGLxLV		29	[AAQYGLxLV	30	162
[AAQYGLxLV		29	[AAQYGLxLV	32	164

Peptide 1	AbsPos1	Peptide 2	AbsPos2	Score
SRMLESLSCc	74	GNSLZAEVE	258	17
SRMLESLSCc	74	GNSLZAEVE	258	21
GNSLZAEVE	258	GNSLZAEVE	258	22
GNSLZAEVE	258	LQIATLLQMo	302	28
GNSLZAEVE	258	GNSLZAEVE	258	28
GNSLZAEVE	258	EEVNELQYR	186	29
GNSLZAEVE	258	GNSLZAEVE	258	30
GNSLZAEVE	258	GNSLZAEVE	258	31
GNSLZAEVE	258	GNSLZAEVE	258	35
GNSLZAEVE	258	GNSLZAEVE	258	36
GNSLZAEVE	258	GNSLZAEVE	258	36
GNSLZAEVE	258	GNSLZAEVE	258	36
GNSLZAEVE	258	GNSLZAEVE	258	37
SRMLESLSCc	74	GNSLZAEVE	258	39
GNSLZAEVE	258	GNSLZAEVE	258	39
GNSLZAEVE	258	GNSLZAEVE	258	40
GNSLZAEVE	258	LIGVPADAEA	539	41
GNSLZAEVE	258	GNSLZAEVE	258	41
MLESLSCcm	76	GNSLZAEVE	258	42
GNSLZAEVE	258	GNSLZAEVE	258	43
GNSLZAEVE	258	GNSLZAEVE	258	45
GNSLZAEVE	258	GNSLZAEVE	258	49
GNSLZAEVE	258	GNSLZAEVE	258	50
GNSLZAEVE	258	GNSLZAEVE	258	51
GNSLZAEVE	258	GNSLZAEVE	258	52
GNSLZAEVE	258	GNSLZAEVE	258	52
GNSLZAEVE	258	GNSLZAEVE	258	53
GNSLZAEVE	258	GNSLZAEVE	258	54
GNSLZAEVE	258	GNSLZAEVE	258	54
GNSLZAEVE	258	GNSLZAEVE	258	55
GNSLZAEVE	258	GNSLZAEVE	258	55
GNSLZAEVE	258	CcmLQLSESE	420	55
GNSLZAEVE	258	GNSLZAEVE	258	55
GNSLZAEVE	258	GNSLZAEVE	258	55
GNSLZAEVE	258	GNSLZAEVE	258	56
GNSLZAEVE	258	GNSLZAEVE	258	56
GNSLZAEVE	258	GNSLZAEVE	258	58
GNSLZAEVE	258	GNSLZAEVE	258	59
SRMLESLSCc	74	GNSLZAEVE	258	59
GNSLZAEVE	258	GNSLZAEVE	258	59
GNSLZAEVE	258	GNSLZAEVE	258	59
GNSLZAEVE	258	GNSLZAEVE	258	60
GNSLZAEVE	258	GNSLZAEVE	258	60
LIGVPADAEA	539	GNSLZAEVE	258	61
SRMoxLESLS	74	GNSLZAEVE	258	62
GNSLZAEVE	258	GNSLZAEVE	258	62
GNSLZAEVE	258	EVLPVDITTAI	458	63

GNSLZAEVE	258	GNSLZAEVE	258	63
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	63
SRMLESLSCc	74	GNSLZAEVE	258	64
GNSLZAEVEI	258	GNSLZAEVE	258	64
GNSLZAEVEI	258	CcmLQLSESE	420	64
LIGVPADAEA	539	GNSLZAEVE	258	64
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	65
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	65
GNSLZAEVEI	258	GNSLZAEVE	258	65
SRMoxLESLS	74	GNSLZAEVE	258	66
GNSLZAEVEI	258	GNSLZAEVE	258	66
GNSLZAEVEI	258	GNSLZAEVE	258	66
GNSLZAEVEI	258	GNSLZAEVE	258	66
GNSLZAEVEI	258	GNSLZAEVE	258	66
GNSLZAEVEI	258	GNSLZAEVE	258	67
GNSLZAEVEI	258	GNSLZAEVE	258	67
GNSLZAEVEI	258	GNSLZAEVE	258	67
GNSLZAEVEI	258	GSQTEFEQQ	312	68
GNSLZAEVEI	258	GNSLZAEVE	258	69
GNSLZAEVEI	258	GNSLZAEVE	258	70
GNSLZAEVEI	258	GNSLZAEVE	258	70
GNSLZAEVEI	258	GNSLZAEVE	258	70
GNSLZAEVEI	258	GNSLZAEVE	258	71
GNSLZAEVEI	258	GNSLZAEVE	258	71
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	71
GNSLZAEVEI	258	GNSLZAEVE	258	72
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	72
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	72
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	72
GNSLZAEVEI	258	GNSLZAEVE	258	73
GNSLZAEVEI	258	GNSLZAEVE	258	73
GNSLZAEVEI	258	GNSLZAEVE	258	73
SRMLESLSCc	74	GNSLZAEVE	258	73
GNSLZAEVEI	258	GNSLZAEVE	258	73
GNSLZAEVEI	258	EVLPVDITTAI	458	73
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	74
SRMLESLSCc	74	GNSLZAEVE	258	74
SRMLESLSCc	74	GNSLZAEVE	258	74
SRMLESLSCc	74	GNSLZAEVE	258	75
LIGVPADAEA	539	<b>GNSLZAEVE</b> [	258	76
GNSLZAEVEI	258	GNSLZAEVE	258	77
GNSLZAEVEI	258	GNSLZAEVE	258	78
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	78
GNSLZAEVEI	258	GNSLZAEVE	258	78
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	79
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	79
GNSLZAEVEI	258	CcmLQLSESE	420	79
GNSLZAEVEI	258	GNSLZAEVE[	258	80

GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	80
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	81
LIGVPADAEA	539	GNSLZAEVE	258	81
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	81
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	81
LIGVPADAEA	539	GNSLZAEVE	258	81
LIGVPADAEA	539	<b>GNSLZAEVE</b> [	258	81
SRMLESLSCc	74	GNSLZAEVE[	258	82
LIGVPADAEA	539	<b>GNSLZAEVE</b> [	258	82
LIGVPADAEA	539	GNSLZAEVE	258	82
GNSI7AFVFI	258	GNSIZAEVE[	258	82
GNSI7AFVFI	258	GNSIZAEVE[	258	82
GNSIZAEVEI	258	GNSIZAEVE	258	82
GNSIZAEVEI	258	GNSLZAEVEL	258	82
GNSLZAEVEL GNSLZAEVEL	258	GNSLZAEVEL	258	83
	250		258	83
SRMIESISCO	230		250	8/
SRMLESLSCC	74		250	84
	74 250		258	84
	230		230	04
	70 2E0		200	04 0E
GNSLZAEVEL	200	GINSLZAEVEL	200	65 0F
GNSLZAEVEL	258	GNSLZAEVEL	258	85
GNSLZAEVEL	258	GNSLZAEVEL	258	86
GNSLZAEVEL	258	GNSLZAEVEL	258	86
GNSLZAEVEL	258	GNSLZAEVEL	258	86
GNSLZAEVEL	258	GNSLZAEVEL	258	86
SRMLESLSCC	74	GNSLZAEVEL	258	87
GNSLZAEVE	258	GNSLZAEVE	258	87
GNSLZAEVE	258	GNSLZAEVE	258	87
GNSLZAEVE	258	GNSLZAEVE	258	87
GNSLZAEVE	258	CcmLQLSESE	420	87
GNSLZAEVEI	258	GNSLZAEVE	258	87
GNSLZAEVEI	258	GNSLZAEVE	258	88
GNSLZAEVEI	258	GNSLZAEVE	258	88
SRMLESLSCc	74	GNSLZAEVE	258	89
GNSLZAEVEI	258	GNSLZAEVE	258	89
GNSLZAEVE	258	GNSLZAEVE	258	89
GNSLZAEVE	258	CcmLQLSESE	420	90
GNSLZAEVE	258	GNSLZAEVE	258	90
GNSLZAEVEI	258	GNSLZAEVE	258	91
SRMLESLSCc	74	GNSLZAEVE	258	91
GNSLZAEVE	258	GNSLZAEVE[	258	91
GNSLZAEVEI	258	GNSLZAEVE[	258	91
GNSLZAEVEI	258	GNSLZAEVE[	258	91
SRMLESLSCc	74	GNSLZAEVE	258	92
GNSLZAEVE	258	GNSLZAEVE	258	92
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	92
GNSLZAEVE	258	GNSLZAEVE	258	92

GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	92
GNSLZAEVE	258	<b>GNSLZAEVE</b> [	258	92
GNSLZAEVE	258	<b>GNSLZAEVE</b> [	258	92
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	93
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	93
GNSLZAEVEI	258	GNSLZAEVE	258	93
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	93
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	93
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	94
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	94
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	94
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	94
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	94
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	94
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	94
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	95
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	96
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	97
GNSLZAEVEI	258	GNSLZAEVE	258	97
SRMLESLSCc	74	GNSLZAEVE	258	97
GNSLZAEVEI	258	GNSLZAEVE	258	97
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	97
GNSLZAEVEI	258	GNSLZAEVE	258	98
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	98
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	98
GNSLZAEVEI	258	GNSLZAEVE	258	99
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	99
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	99
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	99
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	100
GNSLZAEVEI	258	GNSLZAEVE	258	100
GNSLZAEVEI	258	GNSLZAEVE	258	100
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	101
GNSLZAEVEI	258	GNSLZAEVE	258	101
GNSLZAEVEI	258	GNSLZAEVE	258	102
GNSLZAEVEI	258	GSQTEFEQQ	312	102
GNSLZAEVEI	258	GNSLZAEVE	258	103
GNSLZAEVEI	258	GNSLZAEVE	258	103
GNSLZAEVEI	258	GNSLZAEVE	258	103
GNSLZAEVEI	258	GNSLZAEVE	258	103
GNSLZAEVEI	258	GNSLZAEVE	258	103
GNSLZAEVEI	258	GNSLZAEVE	258	103
GNSLZAEVEI	258	GNSLZAEVE	258	104
SRMLESLSCc	74	GNSLZAEVE	258	104
GNSLZAEVEI	258	GNSLZAEVE	258	104
GNSLZAEVEI	258	<b>GNSLZAEVE</b> [	258	104
SRMLESLSCc	74	<b>GNSLZAEVE</b> [	258	104
GNSLZAEVE	258	GNSLZAEVE	258	104

GNSLZAEVE	258	GNSLZAEVE	2	58	105
GNSLZAEVE	258	GNSLZAEVE	2	58	105
LIGVPADAEA	539	GNSLZAEVE	2	58	105
GNSLZAEVEI	258	GNSLZAEVE	2	58	106
GNSLZAEVEI	258	GNSLZAEVE	2	58	106
GNSLZAEVEI	258	GNSLZAEVE	2	58	106
GNSLZAEVE	258	GNSLZAEVE	2	58	107
GNSLZAEVE	258	CcmLQLSESE	42	20	108
GNSLZAEVE	258	GNSLZAEVE	2	58	108
MLESLSCcmE	76	GNSLZAEVE[	2	58	108
GNSLZAEVE	258	GNSLZAEVE[	2	58	108
GNSLZAEVE	258	GNSLZAEVE	2	58	108
GNSLZAEVEI	258	GNSLZAEVE	2	58	108
GNSIZAEVEI	258	GNSI7AEVEI	2	58	109
GNSIZAEVEI	258	GNSI7AFVF	2	58	109
	74	GNSI7AFVF	2	58	110
GNSI 7 AFVFI	258	GNSI7AFVE	2	58	110
	258		2	58	110
	258		2	58	110
	258		2	58	110
	250		2.	58	110
	250		2.	58	111
	258		2.	50	111
	250	GNSLZAEVEL	2.	50	111
	250	GNSLZAEVEL	2.	50	117
	250		2.	50	112
	230		2.	50	112
	44Z 2EQ		2.	50	112
GNSLZAEVEL	200		2.	20	112
	230		2.	50 50	112
	200		2:	20 E0	114
GNSLZAEVEL	200	GNSLZAEVEL	2:	58 50	114
GNSLZAEVEL	258	GNSLZAEVEL	2:	58 50	114
GNSLZAEVEL	258	GNSLZAEVEL	2:	58 50	114
GNSLZAEVEL	258	GNSLZAEVEL	2:	58 50	115
SRIVILESLSCC	/4	GNSLZAEVEL	2:	58	110
GNSLZAEVEL	258	GNSLZAEVEL	2:	58	116
GNSLZAEVEL	258	GNSLZAEVEL	2:	58	116
GNSLZAEVEL	258	GNSLZAEVEL	2:	58	116
GNSLZAEVEL	258	GNSLZAEVEL	2	58	116
GNSLZAEVEL	258	GNSLZAEVEL	2	58	11/
GNSLZAEVEL	258	CCMLQLSESE	4.	20	11/
GNSLZAEVEL	258	GNSLZAEVE	2	58	117
SRMLESLSCc	74	GNSLZAEVE[	2.	58	118
GNSLZAEVE[	258	GNSLZAEVE[	2.	58	118
GNSLZAEVE[	258	GNSLZAEVE[	2.	58	118
GNSLZAEVE	258	GNSLZAEVE[	2	58	118
GNSLZAEVE	258	GNSLZAEVE	2	58	119
GNSLZAEVEI	258	GNSLZAEVE	2	58	120

GNSLZAEVEI	258	GNSLZAEVE	258	121
GNSLZAEVEI	258	GNSLZAEVE	258	121
GNSLZAEVEI	258	GNSLZAEVE	258	122
GNSLZAEVEI	258	GNSLZAEVE	258	123
GNSLZAEVEI	258	GNSLZAEVE	258	124
GNSLZAEVEI	258	GNSLZAEVE	258	124
GNSLZAEVEI	258	GNSLZAEVEI	258	124
GNSLZAEVEI	258	GNSLZAEVEI	258	124
GNSLZAEVEI	258	GNSLZAEVE	258	124
MLESLSCcmE	76	GNSLZAEVEI	258	124
GNSLZAEVEI	258	GNSLZAEVE	258	127
GNSLZAEVEI	258	GNSLZAEVE	258	128
GNSLZAEVEI	258	GNSLZAEVEI	258	128
GNSLZAEVEI	258	GNSLZAEVE	258	129
GNSLZAEVEI	258	GNSLZAEVE	258	129
GNSLZAEVEI	258	GNSLZAEVE	258	129
GNSLZAEVEI	258	GNSLZAEVE	258	129
GNSLZAEVEI	258	GNSLZAEVE	258	130
GNSLZAEVEI	258	GNSLZAEVEI	258	137
GNSLZAEVEI	258	GNSLZAEVEI	258	138
GNSLZAEVEI	258	GNSLZAEVEI	258	140
GNSLZAEVEI	258	GNSLZAEVEI	258	140
GNSLZAEVEI	258	GNSLZAEVEI	258	141
GNSLZAEVEI	258	GNSLZAEVEI	258	141
GNSLZAEVEI	258	GNSLZAEVEI	258	148
GNSLZAEVEI	258	GNSLZAEVEI	258	149
GNSLZAEVEI	258	GNSLZAEVEI	258	151
GNSLZAEVEI	258	GNSLZAEVEI	258	155
GNSLZAEVEI	258	GNSLZAEVEI	258	157
GNSLZAEVEI	258	GNSLZAEVEI	258	158

Peptide1	AbsPos1		Peptide2	AbsPos2	Score
LKEAEEER		25	LKEAEEER	25	210
LKEAEEERLK		25	EVELKSR	81	101
LKEAEEER		25	ΤΚΙΕΚ	123	47
LKEAEEER		25	IEKLK	126	32
LKEAEEERLK		25	LSEKQLK	139	62
LKEAEEER		25	LKEEK	220	78
LKEAEEER		25	YQSLKK	294	100
LKEAEEER		25	NLEKFK	353	102
LKEAEEERLK		25	mKALFESQR	406	119
LKEAEEERLK		25	KLFANER	420	175
LKEAEEER		25	AKLDELK	441	139
LKEAEEERLK		25	LDELKLK	446	96
LKEAEEERLK		25	NLPVDMQLK	538	58
LKEAEEER		25	SGNTPNSPR	560	87
LKEAEEER		25	SGNTPNSPR	566	42
LKEAEEER		25	LAAESK	573	100
LKEAEEERLK		25	LQTEVKEGK	580	103
LKEAEEER		25	EGKETSSK	587	122
LKEAEEER		25	ETSSKLEK	588	108
LKEAEEER		25	LEKETBK	593	26
LKAAQYGLQI		33	MKLQIATLLQ	313	77
LKAAQYGLQI		33	EIESTKGELSI	396	41
LKAAQYGLQI		33	MKALFESQR	411	136
LKAAQYGLQI		33	BLQLSESENN	437	144
LKAAQYGLQI		33	YEPEETVEVP	461	120
LKAAQYGLQI		33	LAAESKLQTE	577	67
LKAAQYGLQI		37	YEPEETVEVP	461	98
AAQYGLQLV		54	KQNVFNR	295	130
AAQYGLQLV		54	MKLQIATLLQ	308	134
AAQYGLQLV		54	MKLQIATLLQ	313	58
AAQYGLQLV		54	MKALFESQR	411	48
AAQYGLQLV		54	LKYEPEETVE	448	108
BRNEMMTN		61	LKEAEEER	25	90
NEMMTMTE		70	KQNVFNR	295	33
EVELKSR		80	EVELKSR	80	213
EVELKSR		80	EVELKSR	81	244
EVELKSR		80	ТКІЕК	123	102
EVELKSR		80	IEKLK	126	119
EVELKSR		80	LSEKQLK	139	169
EVELKSR		80	YQSLKK	294	87
EVELKSR		80	NLEKFK	353	176
EVELKSR		80	AKLDELK	441	177
EVELKSR		80	LDELKLK	446	33
EVELKSR		80	ЕТВКК	595	105
EVELKSR		81	TKIEK	123	52
EVELKSR		81	LSEKQLK	139	127
EVELKSR		81	AKLDELK	441	145

4 EVELKSR	81	149
4 QQQKMHLEK	99	108
4 MHLEKLEEQL	110	143
4 TKIEK	123	82
4 LSEKQLK	139	85
4 EAVSYYNALE	239	92
4 LAAESKLQTE	577	101
4 ETSSKLEK	588	49
0 LKEAEEERLK	25	132
0 QQQKMHLEK	99	68
0 EEKEEREK	223	27
0 EAVSYYNALE	239	160
0 NLEKFK	353	34
0 LDELKLK	446	54
0 LQTEVKEGK	580	36
1 EVELKSR	81	124
1 SHGQEVNEL	112	83
1 TKIEK	123	58
1 LKVELDEAR	128	150
2 LKEAEEER	25	41
2 EVELKSR	80	141
2 TKIEK	123	126
2 LKVELDEAR	128	199
2 LSEKQLK	139	35
2 LKEEK	220	167
2 EEKEER	223	46
2 KLFANER	420	85
3 TKIEK	123	66
6 TKIEK	123	81
6 LKEEK	220	81
8 EVELKSR	81	128
8 TKIEKLK	122	52
8 IEKLK	126	160
8 EEKEER	223	52
8 AKLDELK	441	25
9 TKIEK	123	52
9 IEKLK	126	81
9 LSEKQLK	139	219
9 SEELR	156	35
9 LKEEK	220	66
9 NLEKFK	353	98
9 AKLDELK	441	125
2 EVELKSR	81	48
9 LKEAEEER	25	37
9 EVELKSR	81	105
9 IEKLK	126	30
9 ELLSBKSEELF	155	40
9 ELLSBKSEELF	156	127
	24   EVELKSR     24   QQQKMHLEK     24   MHLEKLEEQL     24   TKIEK     24   LSEKQLK     24   EAVSYYNALE     24   EAVSYYNALE     24   EAVSYYNALE     24   EAVSYYNALE     24   ETSSKLEK     26   EKEEREK     27   EKEEREK     28   EKEEREK     29   EKEKER     20   LOTEVKEGK     21   TKIEK     22   LKEAEER     23   TKIEK     24   LKVELDEAR     25   EKER     26   IKVELDEAR     27   IKIEK     28   IKVELDEAR     29   IKEK     29   IKEK	04 EVELKSR 81   04 QQQKMHLEK 99   04 MHLEKLEEQL 110   04 TKIEK 123   04 LSEKQLK 139   04 LSEKQLK 139   04 LAAESKLQTE 577   04 EAVSYYNALE 239   04 LAAESKLQTE 577   04 ETSSKLEK 588   05 LKEAEEERLK 25   06 QQQKMHLEK 99   10 EEKEEREK 223   10 DQUKKEK 353   10 LDELKLK 446   10 LQTEVKEGK 580   21 EVELKSR 81   22 IKKAEEER 25   23 IKIEK 123   24 LKAEEER 25   25 EVELKSR 80   22 IKVELDEAR 128   23 IKIEK 123   24 LKEEK 220   25 EKEER 223   26 IKEKK

HQVDHQKEL	149 LKEEK	220	39	
HQVDHQKEL	149 NLEKFK	353	84	
HQVDHQKEL	149 MKALFESQR	406	35	
HQVDHQKEL	149 KLFANER	420	25	
ELLSBKSEELF	153 LQTEVKEGK	580	32	
QLKHQVDHQ	153 ETSSKLEK	588	26	
ELLSBKSEELF	155 EVELKSR	80	112	
ELLSBKSEELF	155 LSEKQLK	139	99	
ELLSBKSEELF	155 LKEEK	220	41	
ELLSBKSEELF	155 KLFANER	420	99	
ELLSBKSEELF	156 LKEAEEER	25	91	
ELLSBKSEELF	156 LQTEVKEGK	580	113	
TTLKEEVNEL	189 VKYQSLK	289	73	
TTLKEEVNEL	189 MKALFESQR	406	41	
TTLKEEVNEL	190 IEKLK	126	47	
TTLKEEVNEL	192 LKEAEEER	25	65	
TTLKEEVNEL	192 EVELKSR	81	143	
TTLKEEVNEL	192 LSEKQLK	139	60	
TTLKEEVNEL	192 ELLSBK	153	63	
TTLKEEVNEL	192 SEELR	156	97	
TTLKEEVNEL	192 KQNVFNR	295	100	
TTLKEEVNEL	192 EGKETSSK	586	55	
TTLKEEVNEL	200 LKEAEEER	25	43	
TTLKEEVNEL	200 TKIEK	123	26	
TTLKEEVNEL	200 LSEKQLK	139	32	
TTLKEEVNEL	200 ELLSBKSEELF	155	39	
TTLKEEVNEL	200 NLEKFK	353	72	
TTLKEEVNEL	200 LAAESKLQTE	574	66	
LKEEKEER	220 LKEAEEER	25	40	
LKEEK	220 TKIEK	123	81	
LKEEK	220 IEKLK	126	118	
LKEEK	220 LKEEK	220	135	
LKEEKEEREK	223 TKIEK	123	99	
EEKEEREK	223 IEKLK	126	106	
EEKEER	223 LKEEK	220	30	
EKEAVSYYN/	228 EVELKSR	81	114	
EKEAVSYYN/	228 SHGQEVNEL	122	81	
EKEAVSYYN/	228 TKIEK	123	37	
EKEAVSYYN <i>F</i>	228 LSEKQLK	137	49	
EKEAVSYYN/	228 LKEEK	220	62	
EKEAVSYYN <i>F</i>	228 EEKEEREK	223	54	
EKEAVSYYN <i>F</i>	228 EAVSYYNALE	239	147	
EKEAVSYYN <i>F</i>	228 NLEKFK	353	58	
EKEAVSYYN <i>F</i>	228 KLFANER	420	102	
EKEAVSYYN <i>F</i>	228 AKLDELK	441	69	
EKEAVSYYN <i>F</i>	228 EGKETSSK	587	72	
EAVSYYNALE	232 EEKEER	223	30	
EAVSYYNALE	234 EEKEER	223	52	
EAVSYYNALE	239	LKEAEEER	25	139
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EAVSYYNALE	239	EVELKSR	81	144
EAVSYYNALE	239	QQQKMHLEK	99	151
EAVSYYNALE	239	MHLEKLEEQL	104	170
EAVSYYNALE	239	MHLEKLEEQL	110	171
EAVSYYNALE	239	SHGQEVNEL	122	138
EAVSYYNALE	239	TKIEK	123	43
EAVSYYNALE	239	LKVELDEAR	128	143
EAVSYYNALE	239	LSEKOLK	139	87
EAVSYYNALE	239	ELLSBKSEELF	156	111
EAVSYYNALE	239	EAVSYYNALE	234	
EAVSYYNALE	239	EAVSYYNALE	239	141
FAVSYYNALF	239		289	74
FAVSYYNALF	239	YOSIKK	294	97
FAVSYYNALF	239	KONVENR	295	160
FAVSYYNALF	239		253	142
ΕΛΛΟΣΥΥΝΔΙΕ	233		391	110
ΕΛΛΟΣΥΥΝΔΙΕ	233	MKALEESOR	406	162
	233	KIEANER	400	160
FAVSVVNALE	233		420	201
FAVSVVNALE	233		538	38
FAVSVVNALE	233		574	102
	233		580	166
	233	EGRETSSK	583	100 Q1
	233	EUKEISSK	588	1/2
	233	IEVETDV	502	140 70
	233		120	70 //1
	207		280	41 1/2
	207		209	142
	207		292	155
	207		295	24
	207	EISSNLEN	500	122
	20/	EGREISSK	201	125
	289		294	130
VKYQSLK	290		80	25
VKYQSLK	290		220	35
VKYQSLK	292	LSEKQLK	139	55 170
VKYQSLKK	294	YQSLKK	294	1/8
KQNVFNR	295	EVELKSR	80	121
KQNVFNR	295	EVELKSR	81	132
KQNVFNR	295	LKEEK	220	32
KQNVFNR	295	YQSLKK	294	4/
KQNVFNR	295	NLEKFK	353	81
KQNVFNR	295	KLFANER	420	51
KQNVFNR	295	AKLDELK	441	128
KQNVFNR	295	LDELKLK	446	150
KQNVFNR	295	EGKETSSK	587	76
mKLQIATLLQ	308	KQNVFNR	295	178
mKLQIATLLQ	308	NGEIKHLLGE	342	76

MKLQIATLLQ	308	NLEKFK	353	67
MKLQIATLLQ	308	LDNLNKEIES <sup>-</sup>	396	85
mKLQIATLLQ	308	KLFANER	420	55
mKLQIATLLQ	308	AKLDELK	441	198
MKLQIATLLQ	308	LDELKLK	446	174
MKLQIATLLQ	308	LAAESKLQTE	574	93
MKLQIATLLQ	308	LQTEVKEGK	580	38
MKLQIATLLQ	308	ETSSKLEK	588	27
GSQTEFEQQ	320	LKEAEEER	25	49
LLAmLEQKN	337	NGEIKHLLGE	342	149
LLAMLEQKN	337	NLEKFK	353	64
LLAmLEQKN	337	MKALFESQR	411	56
NGEIKHLLGE	342	LKEAEEERLK	25	119
NGEIKHLLGE	342	EVELKSR	81	102
NGEIKHLLGE	342	ТКІЕК	123	80
NGEIKHLLGE	342	LSEKQLK	139	55
NGEIKHLLGE	342	mKALFESQR	406	155
NGEIKHLLGE	342	KLFANER	420	103
NGEIKHLLGE	342	AKLDELK	441	82
NGEIKHLLGE	342	SGNTPNSPR	563	25
NGEIKHLLGE	342	LAAESK	573	127
NGEIKHLLGE	342	LAAESKLQTE	574	54
NGEIKHLLGE	342	LAAESKLQTE	577	71
NGEIKHLLGE	342	LQTEVKEGK	580	143
NGEIKHLLGE	342	EGKETSSK	586	140
NGEIKHLLGE	342	EGKETSSK	587	128
NGEIKHLLGE	342	ETSSKLEK	588	86
NGEIKHLLGE	342	LEKETBK	591	72
NGEIKHLLGE	342	ЕТВКК	595	103
NLEKFK	353	NLEKFK	353	191
LDNLNKEIES <sup>-</sup>	391	LKEAEEER	25	41
LDNLNKEIES <sup>-</sup>	391	NGEIKHLLGE	342	111
LDNLNKEIES <sup>-</sup>	391	LDNLNKEIES <sup>-</sup>	395	34
LDNLNKEIES <sup>-</sup>	391	LDNLNKEIES <sup>-</sup>	396	35
LDNLNKEIES <sup>-</sup>	391	LAAESKLQTE	577	81
LDNLNKEIES <sup>-</sup>	391	EGKETSSK	587	58
LDNLNKEIES <sup>-</sup>	395	LKEAEEERLK	25	34
LDNLNKEIES <sup>-</sup>	395	ТКІЕК	123	30
LDNLNKEIES <sup>-</sup>	395	VKYQSLK	289	45
LDNLNKEIES <sup>-</sup>	395	YQSLKK	294	79
LDNLNKEIES <sup>-</sup>	395	NLEKFK	353	155
LDNLNKEIES <sup>-</sup>	395	MKALFESQR	406	106
LDNLNKEIES <sup>-</sup>	395	KLFANER	420	145
LDNLNKEIES <sup>-</sup>	396	LKEAEEER	25	170
LDNLNKEIES <sup>-</sup>	396	EVELKSR	80	48
LDNLNKEIES <sup>-</sup>	396	NGEIKHLLGE	342	91
LDNLNKEIES <sup>-</sup>	396	NLEKFK	353	175
LDNLNKEIES <sup>-</sup>	396	MKALFESQR	406	105

LDNLNKEIES	396	KLFANER	420	154
LDNLNKEIES <sup>-</sup>	396	AKLDELK	441	149
LDNLNKEIES	396	LQTEVKEGK	580	164
EIESTKGELSI	397	EVELKSR	80	79
EIESTKGELSI	397	EVELKSR	81	65
EIESTKGELSI	397	TKIEK	123	74
EIESTKGELSI	397	EEKEEREK	223	28
EIESTKGELSI	397	KQNVFNR	295	80
EIESTKGELSI	397	NGEIKHLLGE	342	119
EIESTKGELSI	397	NLEKFK	353	110
EIESTKGELSI	397	LDNLNKEIES <sup>-</sup>	391	152
EIESTKGELSI	397	EIESTK	395	144
FIESTKGELSI	397		396	149
FIESTKGELSI	397	KIFANFR	420	157
FIESTKGELSI	397	LAAFSKIOTE	574	47
FIESTKGELSI	397	FGKFTSSK	587	34
FIESTKGELSI	397	FTSSKIFK	588	98
FIESTKGELSI	397	LEKETBK	500	56
mKALEESOR	406		25	125
MKALEESOR	406	EVELKSR	23 81	123
MKALFESOR	406		126	36
mKALESQR	406		139	34
MKALEESOR	400	KONVENR	295	96
mKALEESQR	400		253	154
mKALESQR	400	FIESTK	306	69
	400	mKALEESOR	406	125
	400		400	12J 81
	400		411	196
	400		420	100
	400		441	110
	400		440 560	02
	400		500	97
	406		580	115
	406	EGREISSK	507	128
	406	EISSKLEK		22
	406		595	88
	420		80	158
KLFANER	420	EVELKSK	130	160
KLFANER	420	IEKLK	126	98
KLFANER	420	LSEKQLK	139	42
KLFANER	420	LKEEK	220	89
KLFANER	420	VKYQSLK	289	95
KLFANER	420	NLEKFK	353	102
KLFANER	420	ALFESQR	411	46
KLFANER	420	KLFANER	420	217
KLFANER	420	AKLDELK	441	148
KLFANER	420	LDELKLK	446	61
KLFANER	420	LAAESK	573	74
KLFANER	420	EGKETSSK	586	119

KLFANER	420	EGKETSSK	587	113
BLQLSESENN	431	NLEKFK	353	42
BLQLSESENN	431	ETSSKLEK	588	36
BLQLSESENN	433	mKALFESQR	406	84
BLQLSESENn	433	EGKETSSK	583	36
BLQLSESENN	437	LKEAEEER	25	28
BLQLSESENN	437	MKLQIATLLQ	308	60
BLQLSESENN	437	LDELKLK	446	108
BLQLSESENn	437	LAAESKLQTE	574	25
LRAKLDELK	441	LKEAEEER	25	26
AKLDELK	441	TKIEK	123	43
AKLDELK	441	IEKLK	126	44
AKLDELK	441	NLEKFK	353	93
AKLDELK	441	AKLDELK	441	134
LRAKLDELK	441	LDELKLK	446	50
AKLDELK	441	LQTEVK	577	31
LDELKLK	446	TKIEK	122	41
LDELKLK	446	LSEKQLK	139	57
LDELKLK	446	YQSLKK	294	180
LDELKLK	446	AKLDELK	441	198
LKYEPEETVE	448	LKEAEEER	25	109
LKYEPEETVE	448	VKYQSLK	292	44
LKYEPEETVE	448	KQNVFNR	295	132
LKYEPEETVE	448	MKALFESQR	411	93
LKYEPEETVE	448	KLFANER	420	39
LKYEPEETVE	448	BLQLSESENN	437	136
LKYEPEETVE	448	LDELKLK	446	123
LKYEPEETVE	448	YEPEETVEVP	461	80
LKYEPEETVE	448	LAAESKLQTE	574	124
YEPEETVEVP	449	AKLDELK	441	85
LKYEPEETVE	461	LKEAEEER	25	74
YEPEETVEVP	461	EVELKSR	80	71
YEPEETVEVP	461	EVELKSR	81	106
YEPEETVEVP	461	EAVSYYNALE	239	46
YEPEETVEVP	461	QLISmKVK	287	86
YEPEETVEVP	461	YQSLKK	294	61
YEPEETVEVP	461	KQNVFNR	295	121
YEPEETVEVP	461	mKLQIATLLQ	308	150
LKYEPEETVE	461	KLFANER	420	149
YEPEETVEVP	461	BLQLSESENn	433	106
YEPEETVEVP	461	BLQLSESENn	437	140
YEPEETVEVP	461	AKLDELK	441	164
YEPEETVEVP	461	YEPEETVEVP	461	152
LKYEPEETVE	461	LAAESKLQTE	574	33
LKYEPEETVE	461	LAAESKLQTE	577	56
YEPEETVEVP	461	EGKETSSK	587	47
YEPEETVEVP	461	ETSSKLEK	588	42
REVLPVDITT/	472	LKEAEEER	25	31

<b>REVLPVDITT</b>	472 mKALFESQR	406	96
EVLPVDITTAI	472 BVKLIGVPAD	545	104
EVLPVDITTAI	473 LKAAQYGLQI	33	122
EVLPVDITTAI	473 EAVSYYNALE	239	132
EVLPVDITTAI	473 mKLQIATLLQ	308	112
<b>REVLPVDITT</b>	473 mKALFESQR	406	86
EVLPVDITTAI	473 KLFANER	420	26
EVLPVDITTAI	473 BLQLSESENN	437	78
EVLPVDITTAI	473 LAAESKLQTE	574	81
EVLPVDITTAI	475 MKLQIATLLQ	308	121
EVLPVDITTAI	475 MKALFESQR	411	138
RREVLPVDIT	475 KLFANER	420	71
EVLPVDITTAI	475 BLQLSESENn	433	113
RREVLPVDIT <sup>®</sup>	475 AKLDELK	441	57
EVLPVDITTAI	475 EVLPVDITTAI	489	33
EVLPVDITTAI	475 EETQSBPNSL	504	63
EVLPVDITTAI	475 LAAESKLQTE	577	42
EVLPVDITTAI	482 MKLQIATLLQ	308	36
EVLPVDITTAI	482 BLQLSESENN	437	143
EVLPVDITTAI	489 MKLQIATLLQ	308	47
DABVNNSAL	489 MKALFESQR	406	31
EVLPVDITTAI	489 LKYEPEETVE	448	89
EVLPVDITTAI	489 YEPEETVEVP	461	129
LPPQKEETQS	495 LKAAQYGLQI	37	25
LPPQKEETQS	495 EVELKSR	80	29
LPPQKEETQS	495 KQNVFNR	295	28
LPPQKEETQS	495 MKLQIATLLQ	308	50
LPPQKEETQS	495 mKALFESQR	406	95
LPPQKEETQS	495 BLQLSESENn	433	25
LPPQKEETQS	495 YEPEETVEVP	461	139
LPPQKEETQS	495 LAAESKLQTE	577	25
EETQSBPNSL	498 EVELKSR	80	25
LPPQKEETQS	498 LKYEPEETVE	448	41
LPPQKEETQS	498 YEPEETVEVP	461	98
LPPQKEETQS	498 LAAESKLQTE	574	34
LPPQKEETQS	500 EIESTKGELSI	397	65
LPPQKEETQS	500 BLQLSESENN	437	31
EETQSBPNSL	500 LAAESKLQTE	574	77
EETQSBPNSL	504 KQNVFNR	295	105
LPPQKEETQS	504 MKALFESQR	406	45
EETQSBPNSL	504 KLFANER	420	39
EETQSBPNSL	504 AKLDELK	441	63
EETQSBPNSL	504 LAAESKLQTE	574	73
EETQSBPNSL	504 LQTEVKEGK	580	65
EETQSBPNSL	504 EGKETSSK	583	64
EETQSBPNSL	504 ETSSKLEK	588	49
NLPVDMQLK	538 LKEAEEER	25	41
NLPVDMQLK	538 EVELKSR	80	38

NLPVDMQLK	538	EVELKSR	81	125
NLPVDMQLK	538	LSEKQLK	139	33
NLPVDMQLK	538	NLEKFK	353	98
NLPVDMQLK	538	MKALFESQR	406	85
NLPVDMQLK	538	MKALFESQR	411	50
NLPVDmQLK	538	KLFANER	420	147
NLPVDMQLK	538	AKLDELK	441	43
NLPVDMQLK	538	LQTEVKEGK	580	99
NLPVDMQLK	538	EGKETSSK	587	53
BVKLIGVPAD	545	EVELKSR	81	105
BVKLIGVPAD	545	LAAESKLOTE	574	96
BVKLIGVPAD	545	EGKETSSK	587	35
LIGVPADAEA	557	NGEIKHLLGE	342	84
SGNTPNSPR	560	EVELKSR	80	174
SGNTPNSPR	560	NIFKFK	353	81
SGNTPNSPR	560	AKIDELK	441	133
SGNTPNSPR	560	FGKFTSSK	583	151
SGNTPNSPR	566	KIFANFR	420	75
SGNTPNSPR	566	FGKFTSSK	583	44
	574		25	110
	574	YTI OR	72	42
	574	FVFLKSR	81	42
	574		287	32
	574	mKALEESOR	406	165
	574	KIFANFR	400	75
	57/		420	11/
	57/		538	16
	57/		573	63
	574		588	144
	577		25	144
	577		20	100
	577		122	138
	577		125	20
	5//		120	28
	5//		159	99 100
	5//		207	20
	5//		294	38
	5//		290	150
	5//		303	188
	5//		406	1/5
	5//		420	202
	5//	AKLDELK	441	135
	5//		446	89
LAAESKLQTE	5//	NLPVDMQLK	538	88
	5//		580	95
LAAESKLQTE	577	EISSKLEK	588	147
LAAESKLQTE	577	LEKETBK	591	119
LQTEVKEGK	580	LKEAEEER	25	42
LQTEVKEGK	580	EVELKSR	81	113

LQTEVKEGK	580	ТКІЕК	123	59
LQTEVKEGK	580	LKEEK	220	35
LQTEVKEGK	580	KQNVFNR	295	60
LQTEVKEGK	580	NLEKFK	353	100
LQTEVKEGK	580	KLFANER	420	54
LQTEVKEGK	580	LDELKLK	446	77
LQTEVKEGK	580	SGNTPNSPR	560	76
LQTEVKEGK	580	SGNTPNSPR	566	28
LQTEVKEGK	580	LQTEVKEGK	580	140
LQTEVKEGK	580	EGKETSSK	587	91
LQTEVKEGK	580	LEKETBK	593	148
LQTEVKEGK	580	ЕТВКК	595	70
EGKETSSK	583	EVELKSR	81	116
EGKETSSK	583	NLEKFK	353	163
EGKETSSK	583	EGKETSSK	587	146
EGKETSSK	586	EVELKSR	80	121
EGKETSSK	586	TKIEK	123	39
ETSSKLEK	586	NLEKFK	353	87
EGKETSSK	586		446	65
EGKETSSK	586	ETBKK	595	57
EGKETSSK	587	EVELKSR	80	158
EGKETSSK	587	IEKLK	126	56
EGKETSSK	587	LSEKOLK	139	83
EGKETSSK	587	LKEEK	220	70
ETSSKLEK	587	YQSLKK	294	49
EGKETSSK	587	NLEKFK	353	108
EGKETSSK	587	AKLDELK	441	96
EGKETSSK	587	ЕТВКК	595	57
ETSSKLEK	588	EVELKSR	80	148
ETSSKLEK	588	EVELKSR	81	142
ETSSKLEK	588	TKIEK	123	46
ETSSKLEK	588	IEKLK	126	41
ETSSKLEK	588	LSEKQLK	139	72
ETSSKLEK	588	LKEEK	220	60
ETSSKLEK	588	VKYQSLK	289	88
ETSSKLEK	588	VKYQSLK	292	60
ETSSKLEK	588	YQSLKK	294	135
ETSSKLEK	588	KLFANER	420	80
ETSSKLEK	588	AKLDELK	441	130
ETSSKLEK	588	LDELKLK	446	61
ETSSKLEK	588	LAAESK	573	84
ETSSKLEK	588	EGKETSSK	587	152
ETSSKLEK	588	LEKETBK	593	62
LEKETBK	591	EVELKSR	81	116
LEKETBK	591	LSEKQLK	139	42
LEKETBK	591	LKEEK	220	27
LEKETBK	591	NLEKFK	353	96
LEKETBK	593	EVELKSR	80	142

LEKETBK	593 LKEEK	220	58
ЕТВКК	595 TKIEK	123	25
EGKETSSKLE	595 ELLSBKSEELF	156	29

Score	Peptide1	AbsPos1	Peptide2	AbsPos2
265	LSEKQLK	139	LSEKQLK	139
248	EVELKSR	80	EVELKSR	81
245	NLEKFK	353	NLEKFK	353
225	KLFANER	420	KLFANER	420
223	LKEAEEER	25	LKEAEEER	25
222	EVELKSR	80	NLEKFK	353
217	' EAVSYYNALE	239	EAVSYYNALE	239
212	EVELKSR	80	LSEKQLK	139
211	LDELKLK	446	AKLDELK	441
209	EVELKSR	81	NLEKFK	353
207	' EVELKSR	80	AKLDELK	441
203	EVELKSR	81	AKLDELK	441
202	mKLQIATLLQ	308	AKLDELK	441
191	EAVSYYNALE	239	MKALFESQR	406
186	SHGQEVNEL	122	LKVELDEAR	128
185	mKALFESQR	406	mKALFESQR	406
184	EAVSYYNALE	239	MHLEKLEEQI	110
183	LKEAEEER	25	KLFANER	420
182	LDNLNKEIES	396	MKALFESQR	406
182	ELLSBKSEELF	156	KLFANER	420
181	mKLQIATLLQ	308	KQNVFNR	295
179	EVELKSR	81	LSEKQLK	139
178	EKEAVSYYNA	228	EAVSYYNALE	239
178		396	NLEKFK	353
178	LKEAEEER	25	LKEEK	220
174	YQSLKK	294	YQSLKK	294
173	MKALFESQR	406	EVELKSR	81
172	LDNLNKEIES	396	LKEAEEER	25
172	mKLQIATLLQ	308	KLFANER	420
172	EVELKSR	80	EVELKSR	80
172	mKALFESQR	406	KLFANER	420
171	KLFANER	420	EVELKSR	81
170	EAVSYYNALE	239	LKEAEEER	25
169	MKLQIATLLQ	308	LDELKLK	446
166	AKLDELK	441	AKLDELK	441
165	LKEEK	220	LKEEK	220
164	KLFANER	420	AKLDELK	441
164	TTLKEEVNEL	192	KQNVFNR	295
163	mHLEKLEEQL	110	EAVSYYNALE	239
161	mKALFESOR	406	ALFESQR	411
161	MHLEKLEEQL	104	EVELKSR	81
159		446	YOSLKK	294
158	mKLQIATLLO	308	mKALFESOR	406
157	EAVSYYNAI F	239	EVELKSR	81
157	SHGOEVNEL	122	KLFANER	420
156		54	MKLOIATIIO	308
156	KQNVFNR	295	EVELKSR	81

156 EAVSYYNALE	239	KLFANER	420
153 EKEAVSYYNA	228	<b>EKEAVSYYN</b>	228
152 LLAmLEQKN(	337	NGEIKHLLGE	342
151 mKLQIATLLQ	308	NLEKFK	353
151 LKVELDEAR	128	EVELKSR	81
151 BLQLSESENn	437	LDELKLK	446
151 EIESTKGELSI	397	KLFANER	420
151 LDNLNKEIES	396	KLFANER	420
150 EIESTKGELSI	397	LDNLNKEIES	396
150 MHLEKLEEQL	104	MHLEKLEEQL	110
149 MKALFESQR	406	KQNVFNR	295
147 LKAAQYGLQI	37	mKLQIATLLQ	308
147 LDNLNKEIES	396	AKLDELK	441
146 EIESTKGELSI	397	EIESTK	395
145 LKAAQYGLQI	33	MKLQIATLLQ	308
144 QLISMKVK	287	KLFANER	420
142 EAVSYYNALE	239	ELLSBKSEELF	156
141 EAVSYYNALE	239	MKLQIATLLQ	308
141 LSEKQLK	139	AKLDELK	441
140 HQVDHQKEL	149	EVELKSR	81
140 mKALFESQR	406	NLEKFK	353
140 SHGQEVNEL	122	EVELKSR	80
138 mHLEKLEEQL	104	EAVSYYNALE	239
138 NGEIKHLLGE	342	EVELKSR	81
138 LKAAQYGLQI	33	EIESTKGELSI	396
137 MKLQIATLLQ	308	LDNLNKEIES	396
137 MHLEKLEEQL	104	AKLDELK	441
137 LDNLNKEIES	395	KLFANER	420
137 EAVSYYNALE	239	KQNVFNR	295
136 LKAAQYGLQI	33	mKLQIATLLQ	313
136 LSEKQLK	139	NLEKFK	353
136 LKEAEEER	25	NLEKFK	353
135 EAVSYYNALE	239	LKVELDEAR	128
135 BLQLSESENn	437	mKLQIATLLQ	308
132 VKYQSLK	289	EVELKSR	81
132 EAVSYYNALE	239	ELLSBKSEELF	155
132 LKVELDEAR	128	KLFANER	420
132 LKEAEEER	25	AKLDELK	441
131 EVELKSR	80	LDELKLK	446
130 EAVSYYNALE	239	NLEKFK	353
130 KQNVFNR	295	NLEKFK	353
130 QLISMKVK	287	AKLDELK	441
130 ELLSBKSEELF	156	EVELKSR	80
129 mHLEKLEEQL	104	NLEKFK	353
128 NLEKFK	353	LKEEK	220
127 MKLQIATLLQ	308	EVELKSR	81
127 LKAAQYGLQI	37	BLQLSESENN	437
127 NGEIKHLLGE	342	AKLDELK	441

127 TKIEKLK	126	LKEEK	220
126 NGEIKHLLGE	342	mKALFESQR	406
126 LKVELDEAR	128	IEKLK	126
126 LDELKLK	446	NLEKFK	353
126 HQVDHQKEL	149	ELLSBKSEELF	156
126 LKEEK	220	ТКІЕК	123
125 EKEAVSYYNA	228	KLFANER	420
125 LKAAQYGLQI	33	BLQLSESENN	431
125 LKEAEEERLK	25	EVELKSR	81
125 LKEAEEER	25	YQSLKK	294
125 KQNVFNR	295	KQNVFNR	295
124 MKALFESQR	406	LKEAEEER	25
124 BLQLSESENn	437	KQNVFNR	295
124 mKLQIATLLQ	308	mKLQIATLLQ	308
124 LKAAQYGLQI	33	LQIATLLQmK	318
123 EIESTKGELSI	397	AKLDELK	441
123 SHGQEVNEL	122	LKEEK	220
122 LDELKLK	446	LKYEP}	448
122 KLFANER	420	LKYEP}	448
122 VKYQSLK	289	VKYQSLK	292
121 LKEAEEERLK	25	EVELKSR	80
121 GSQTEFEQQ	320	LKEAEEER	25
121 LKAAQYGLQI	33	LQIATLLQMK	322
121 QLISMKVK	287	VKYQSLK	289
120 BLQLSESENn	437	LKEAEEER	25
120 MKLQIATLLQ	308	LKEAEEER	25
120 MKALFESQR	406	LDELKLK	446
120 LLAmLEQKN(	337	LKEAEEER	25
119 MKLQIATLLQ	308	LKEEK	220
119 KLFANER	420	LSEKQLK	139
119 AAQYGLQLV	54	LKYEP}	448
119 EAVSYYNALE	239	SHGQEVNEL	122
118 EAVSYYNALE	239	LDNLNKEIES	391
117 NGEIKHLLGE	342	EIESTK	396
116 LDNLNKEIES	396	KQNVFNR	295
116 ELLSBKSEELF	155	MKALFESQR	406
116 NGEIKHLLGE	342	KLFANER	420
115 QLISmKVK	287	VKYQSLK	292
115 EIESTKGELSI	397	EIESTKGELSI	397
114 MKLQIATLLQ	308	LKYEP}	448
113 TTLKEEVNEL	192	EVELKSR	81
113 EKEAVSYYNA	234	EAVSYYNALE	239
113 SHGQEVNEL	121	LKVELDEAR	128
113 LKEAEEERLK	25	mKALFESQR	406
112 LRAKLDELK	441	LDELKLK	446
112 ELLSBKSEELF	155	LKYEP}	448
112 BLQLSESENn	433	LKYEP}	448
112 LDNLNKEIES	396	LSEKQLK	139

111	QLISMKVK	287	KQNVFNR	295
111	EKEAVSYYN <i>A</i>	228	KQNVFNR	295
110	BLQLSESENN	437	LKYEP}	448
110	LKEEK	220	IEKLK	126
110	LLAMLEQKN(	337	MKALFESQR	406
109	LDELKLK	446	LSEKQLK	139
108	BLQLSESENn	437	LDNLNKEIES <sup>-</sup>	396
106	HQVDHQKEL	149	KLFANER	420
106	MHLEKLEEQL	104	LSEKQLK	139
106	mKALFESQR	406	AKLDELK	441
106	NGEIKHLLGE	342	LDELKLK	446
105	EAVSYYNALE	239	LDELKLK	446
104	KQNVFNR	295	KLFANER	420
104	TTLKEEVNEL	192	ELLSBKSEELF	156
104	KQNVFNR	295	LDELKLK	446
104	SHGQEVNEL	122	TKIEK	123
103	mKALFESQR	406	IEKLK	126
103	LKAAQYGLQI	33	MKALFESQR	411
103	EAVSYYNALE	239	MHLEKLEEQI	104
103	TTLKEEVNEL	192	ELLSBK	153
102	KLFANER	420	YTLQR	71
102	LKEAEEERLK	25	LDELKLK	446
102	EIESTKGELSI	397	NGEIKHLLGE	342
102	AAQYGLQLV	54	LKAAQYGLQI	33
102	NGEIKHLLGE	342	IEKLK	126
102	EAVSYYNALE	239	EAVSYYNALE	234
102	LSEKQLK	139	LKEEK	220
101	EAVSYYNALE	239	QQQKMHLEK	99
101	LDNLNKEIES	391	LDNLNKEIES	395
101	TKIEK	123	TKIEK	123
100	NGEIKHLLGE	342	LKEAEEER	25
99	BLQLSESENn	437	NLEKFK	353
98	NGEIKHLLGE	342	LKEEK	220
97	EIESTKGELSI	397	NLEKFK	353
97	KLFANER	420	IEKLK	126
97	EVELKSR	81	IEKLK	126
95	VKYQSLK	289	YQSLKK	294
94	mHLEKLEEQL	110	EVELKSR	80
93	EKEAVSYYNA	228	AKLDELK	441
93	LDELKLK	446	LKEEK	220
93	MHLEKLEEQL	110	mKLQIATLLQ	308
93	TTLKEEVNEL	200	<b>EKEAVSYYN</b>	228
92	NLYDSmESKI	364	MKALFESQR	411
92	SHGQEVNEL	122	AKLDELK	441
91	NGEIKHLLGE	342	KQNVFNR	295
91	SHGQEVNEL	121	SHGQEVNEL	122
91	TTLKEEVNEL	192	LSEKQLK	139
90	AAQYGLQLV	54	KQNVFNR	295

90 BLQLSESENN	433	mKALFESQR	406
90 LKEAEEER	25	IEKLK	126
90 VKYQSLK	292	AKLDELK	441
90 KLFANER	420	VKYQSLK	289
89 AAQYGLQLV	54	MKALFESQR	411
89 EAVSYYNALE	239	LKEEK	220
89 LDNLNKEIES	396	NGEIKHLLGE	342
89 EAVSYYNALE	239	LKYEP}	448
88 MHLEKLEEQL	110	KQNVFNR	295
88 KQNVFNR	295	LKYEP}	448
88 EEKEEREK	223	IEKLK	126
86 EAVSYYNALE	239	NGEIKHLLGE	342
86 BLQLSESENn	437	EVELKSR	81
86 LKEAEEER	25	KQNVFNR	295
86 EKEAVSYYNA	232	EVELKSR	80
86 lkaaqyglqi	33	BLQLSESENn	437
86 EKEAVSYYNA	228	EVELKSR	81
86 QLISmKVK	287	EVELKSR	81
84 TTLKEEVNEL	190	EVELKSR	80
83 HQVDHQKEL	149	HQVDHQKEL	149
83 mHLEKLEEQL	110	KLFANER	420
83 MHLEKLEEQL	104	QQQKMHLEK	99
83 EEREKEAVSY	228	LKVELDEAR	128
81 ELLSBKSEELF	155	LSEKQLK	139
81 EVELKSR	80	SEELR	156
80 LQIATLLQmK	320	MKALFESQR	406
80 BLQLSESENN	437	MKALFESQR	406
79 KLFANER	420	NLEKFK	353
79 LDNLNKEIES	396	EVELKSR	80
79 NLEKFK	353	LKYEP}	448
78 AAQYGLQLV	54	EAVSYYNALE	239
78 MKALFESQR	406	YTLQR	71
78 EEKEEREK	223	ΤΚΙΕΚ	123
77 EVELKSR	80	YQSLKK	294
76 LDNLNKEIES	396	LDELKLK	446
75 LDNLNKEIES	391	LKEAEEER	25
74 EVELKSR	80	LKYEP}	448
74 NGEIKHLLGE	342	ТКІЕК	123
73 AKLDELK	441	LKEEK	220
72 GELSIQR	401	NLEKFK	353
71 mKALFESQR	406	EIESTK	396
71 LKEAEEERLK	25	QLISmKVK	287
69 EKEAVSYYNA	234	mKALFESQR	406
69 mKALFESQR	406	LKYEP}	448
69 VKYQSLK	292	LDELKLK	446
68 LSEKQLK	139	IEKLK	126
68 ELLSBKSEELF	155	EVELKSR	81
68 KQNVFNR	295	AKLDELK	441

68 EVELKSR	81	EEKEER	223
66 mHLEKLEEQL	104	LKEAEEER	25
66 KLFANER	420	LDELKLK	446
65 AAQYGLQLV	54	mKLQIATLLQ	313
64 EEREKEAVSY	228	EEKEER	223
64 LLAmLEQKN(	337	KLFANER	420
64 LLAmLEQKN(	337	MKALFESQR	411
63 mKALFESQR	411	NLEKFK	353
63 EEREKEAVSY	228	IEKLK	126
62 LDNLNKEIES	391	LKYEP}	448
62 ELLSBKSEELF	155	SEELR	156
61 LLAMLEQKN(	337	NLEKFK	353
61 EIESTKGELSI	397	SHGQEVNEL	122
60 EKEAVSYYNA	232	MKALFESQR	406
59 NLYDSmESKI	376	mKLQIATLLQ	308
59 LQIATLLQmK	318	KLFANER	420
59 TTLKEEVNEL	200	MKALFESQR	406
59 EEKEER	223	EEKEER	223
58 EIESTKGELSI	397	ТКІЕК	123
58 EAVSYYNALE	239	YQSLKK	294
58 EVELKSR	80	EEKEER	223
57 KLFANER	420	EEKEER	223
57 ELLSBKSEELF	155	LDELKLK	446
57 LKAAQYGLQI	33	KLFANER	420
56 LQIATLLQMK	318	LKEAEEER	25
56 NGEIKHLLGE	342	LKYEP}	448
56 YTLQREVELK	81	EVELKSR	80
56 EKEAVSYYNA	228	MHLEKLEEQI	110
55 MHLEKLEEQL	104	TKIEK	123
55 MHLEKLEEQL	110	ТКІЕК	123
55 TTLKEEVNEL	192	TTLKEEVNEL	200
54 KQNVFNR	295	EVELKSR	80
54 EAVSYYNALE	239	EIESTKGELSI	397
53 EAVSYYNALE	234	EEKEER	223
53 TTLKEEVNEL	192	VKYQSLK	289
53 AKLDELK	441	IEKLK	126
53 MHLEKLEEQL	110	NLEKFK	353
52 TTLKEEVNEL	189	LSEKQLK	139
52 QLISmKVK	287	LKYEP}	448
52 IEKLK	126	TKIEK	123
52 SHGQEVNEL	122	KQNVFNR	295
51 NEmmTMTE	70	QLISmKVK	287
50 AAQYGLQLV	54	MKALFESQR	406
50 LSEKQLK	139	ELLSBK	153
50 LSEKQLK	139	LKYEP}	448
50 QLISMKVK	287	NLEKFK	353
50 ELLSBKSEELF	156	IEKLK	126
50 LDNLNKEIES	395	NLEKFK	353

50 LKVELDEAR	128	EEKEER	223
50 LDNLNKEIES	395	LDELKLK	446
49 mHLEKLEEQL	110	MKALFESQR	406
49 NLEKFK	353	YQSLKK	294
49 EKEAVSYYNA	228	SHGQEVNEL	122
49 LKAAQYGLQI	33	LDNLNKEIES	391
47 NLYDSmESKI	364	EIESTKGELSI	397
47 EKEAVSYYNA	228	NLEKFK	353
47 LSEKQLK	139	ТКІЕК	123
47 VKYQSLK	292	LKEEK	220
45 NEmmTMTE	70	KQNVFNR	295
43 AAQYGLQLV	54	EIESTKGELSI	397
43 VKYQSLKKQN	295	mKALFESQR	406
43 LDNLNKEIES	391	LDNLNKEIES <sup>-</sup>	396
42 GSQTEFEQQ	320	KLFANER	420
42 BLQLSESENn	433	NLEKFK	353
42 mKLQIATLLQ	308	YQSLKK	294
41 LQIATLLQMK	318	MKALFESQR	411
41 TTLKEEVNEL	200	LSEKQLK	139
40 EKEAVSYYNA	228	LKEEK	220
40 mKLQIATLLQ	308	LDNLNKEIES <sup>-</sup>	395
39 NGEIKHLLGE	342	QLISmKVK	287
39 BLQLSESENn	433	LKEAEEER	25
39 mKALFESQR	406	LKEEK	220
39 mHLEKLEEQL	104	EEKEER	223
38 EKEAVSYYNA	228	LSEKQLK	139
37 HQVDHQKEL	149	LKEAEEER	25
37 EKEAVSYYNA	228	LSEKQLK	137
37 MHLEKLEEQL	110	LKEAEEER	25
35 LRAKLDELK	441	LKEAEEER	25
35 SHGQEVNEL	122	EEKEER	223
35 mKALFESQR	406	LSEKQLK	139
35 LDNLNKEIES	391	LDELKLK	446
34 EIESTKGELSI	395	EVELKSR	80
34 TTLKEEVNEL	189	MKALFESQR	406
34 BLQLSESENn	437	LSEKQLK	139
33 EIESTKGELSI	397	LDELKLK	446
32 EAVSYYNALE	239	EEKEER	223
32 KQNVFNR	295	IEKLK	126
31 ELLSBKSEELF	155	EEKEER	223
31 LDNLNKEIES	395	IEKLK	126
31 LDNLNKEIES	396	YQSLKK	294
31 EAVSYYNALE	239	QLISMKVK	287
30 SHGQEVNEL	121	SHGQEVNEL	112
30 mHLEKLEEQL	104	mKALFESQR	411
29 LDNLNKEIES	395	YQSLKK	294
28 LQIATLLQMK	320	AKLDELK	441
28 GSQTEFEQQ	322	KLFANER	420

27	LQIATLLQMK	320	BLQLSESENn	437
27	BLQLSESENn	437	EIESTKGELSI	397
27	MHLEKLEEQL	110	EEKEEREK	223
27	EVELKSR	80	IEKLK	126
27	LKVELDEAR	128	LDELKLK	446
26	NGEIKHLLGE	342	mKALFESQR	411
24	EAVSYYNALE	239	ТКІЕК	123
24	mHLEKLEEQL	104	mKALFESQR	406
23	NLEKFK	353	ТКІЕК	123
23	LKEAEEER	25	GELSIQR	401