DYNAMIC KINETOCHORE SIZE REGULATION PROMOTES MICROTUBULE CAPTURE

AND CHROMOSOME BIORIENTATION IN MITOSIS

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ABSTRACT

Faithful chromosome segregation depends on the ability of sister kinetochores to attach to spindle microtubules. An outer layer of the kinetochore known as the fibrous corona transiently expands in early mitosis and disassembles upon microtubule capture. Neither the functional importance nor the mechanistic basis for this are known. Here we show that the dynein adaptor Spindly and the RZZ kinetochore complex drive fibrous corona formation in a dynein-independent manner. C-terminal farnesylation and MPS1 kinase activity cause conformational changes of Spindly that promote oligomerization of RZZ:Spindly complexes into a corona-like meshwork in cells and in vitro. Concurrent with corona expansion, Spindly potentiates corona shedding by recruiting dynein via three conserved short linear motifs. Expanded, nonsheddable fibrous coronas engage in extensive, long-lived lateral microtubule interactions that persist to metaphase and result in fused sister kinetochores, formation of merotelic attachments and chromosome segregation errors in anaphase. Thus, dynamic kinetochore size regulation in mitosis is coordinated by a single, Spindly-based mechanism that promotes initial microtubule capture and subsequent correct maturation of attachments.

INTRODUCTION

The kinetochore is a complex protein structure that physically anchors the chromosomes to spindle microtubules to ensure chromosome segregation during cell division ^{1,2}. Defects in kinetochore function cause chromosome segregation errors that lead to aneuploid cellular progeny. Aneuploidy is a major cause of embryonic maldevelopment and a prominent genomic abnormality in human cancers ^{3–9}.

To guarantee the fidelity of chromosome segregation in mitosis, chromosomes must capture microtubules and achieve stable bioriented attachments before anaphase, meaning they are bound to microtubules from opposite spindle poles. Microtubule capture and biorientation are complex processes that require an intricate kinetochore machinery, involving dozens of proteins that dynamically adapt to the microtubule attachment status of the kinetochore ^{1,2,10–12}. The core of the kinetochore in animals and fungi is made of the CCAN and the KMN network. The CCAN associates with centromeric chromatin and connects it to the microtubule-binding site that resides within the HEC1/NDC80 and NUF2 members of the KMN network ². HEC1/NDC80 is a key target in the biorientation pathway, as its phosphorylation by Aurora B kinase signals non-bioriented attachments that are then corrected accordingly ¹³.

The core kinetochore module is supplemented with proteins whose quantity at kinetochores changes during mitotic progression. These include proteins of the spindle assembly checkpoint (SAC), which bind predominantly to the scaffold KNL1 when HEC1/NDC80 is not bound to microtubules ¹⁴. Other transient kinetochore proteins, more distal to the centromeric chromatin, are dynein recruiters (ROD-ZW10-Zwilch (RZZ) and Spindly), dynein regulators (CENP-F/Nude1/Nde1/CLIP-170), the

kinesins CENP-E and Kif2b, and modifiers of microtubule dynamics such as CLASP1 and CLASP2 ^{15,16}. These proteins are involved in initial microtubule capture, chromosome transport, and microtubule-based force generation that enables, amongst others, anaphase.

The mammalian kinetochore exhibits great morphological plasticity during mitosis. While the core kinetochore module expands little if any (with the exception of CENP-C in *Xenopus* extracts), the more centromere-distal modules expand to large, crescent shapes in early prometaphase and collapse into compact spherical structures in metaphase ^{17–27}. Electron micrographs of kinetochores in their most expanded form show a halo of low-density material referred to as the fibrous corona, which is absent from kinetochores that are bound by microtubules ^{28–33}. Neither the molecular nature of the fibrous corona nor its functional relevance are known. Recent computational modeling suggested that the fibrous corona promotes spindle assembly, and that correct rotation of the sister kinetochores followed by their compaction then reduces the risk of erroneous attachments ^{17,34}. Experimental approaches to manipulate the fibrous corona in order to test this and other hypotheses are lacking ³⁴.

The dynamic behaviour of the fibrous corona correlates with that of the proteins that display early mitotic expansion and subsequent compaction. It is therefore likely that one or more of them can form the corona meshwork, and that regulation of its expansion/compaction will impinge on these components. A candidate is the RZZ complex. RZZ shows classic expansion/compaction behavior, is required for recruitment of several outer-kinetochore components, and has a molecular architecture resembling that of membrane coating proteins that can form polymeric states ^{25,35–44}. We here set out to identify the nature of the fibrous corona,

the molecular mechanisms of its ability to expand and compact, and its functional relevance for chromosome segregation. We show that the Spindly protein is at the heart of fibrous corona biology: It forms the fibrous corona by triggering polymerization of RZZ-Spindly complexes, and subsequently sets the stage for later corona compaction by recruiting the dynein motor complex. In this way, Spindly couples the mechanisms of physical expansion of kinetochores with subsequent compaction and we show this enables efficient capture and stabilization of correct microtubule attachments.

RESULTS

Spindly recruits dynein to compact kinetochores upon microtubule attachment

In agreement with previous observations ^{17,18,22}, kinetochores expand in nocodazole-treated cells shortly after NEB, as evidenced by the formation of ZW10-positive crescents (Figure 1a, examples 1 to 3). The crescents grew during mitosis and merged with those of the sister kinetochore to form large, ring-shaped structures (Figure 1a, example 4). Rings were absent from cells in which a spindle was allowed to form, and the attached kinetochore on a mono-oriented chromosome was more compact than its unattached sister kinetochore (Figure 1a, example 5). To examine the function of this expansion and subsequent compaction by microtubules, we wished to understand its molecular underpinnings. In *Xenopus* egg extracts, kinetochore size is regulated by mitotic phosphorylation ²⁴. Phosphatases were, however, unable to reverse kinetochore expansion in human cells, as small molecule inhibitors of Aurora B, PLK1 or MPS1, when added after expansion, did not cause compaction of kinetochores (Figure S1).

Dynein drives poleward transport of kinetochore proteins ^{25,45–50}. To investigate if dynein is required for compaction of expanded kinetochores, we examined Spindly, a predicted coiled-coil protein that is the dynein/dynactin adaptor at kinetochores ^{26,35,43,51–53}. Spindly localization to kinetochores requires a conserved C-terminal CAAX box (Figure 1b), whose farnesylation promotes binding to the RZZ complex ^{35,43,51,54,55}. Spindly in turn recruits dynein/dynactin through two motifs: the Spindly box (SB), and the recently identified CC1 box (Figure 1b and S2a) ^{35,43,51,52}. Our ConFeaX pipeline, which identifies conserved motifs in orthologous protein families ^{56,57}, additionally identified a Qxx[HY] motif close to the CC1 box, which we here

named the CC2 box (Figure 1b). CC2-like boxes can be found in other dynein/dynactin adaptors (BICD1/2, BICDR1/2, TRAK1/2, and HAP1, Figure S2a), suggesting they are involved in dynein/dynactin binding. Indeed, recruitment of the dynactin subunit p150^{glued} to kinetochores was compromised in cells expressing GFP-Spindly mutated in either of the three motifs [A24V (Spindly ACC1), Y60A (Spindly CC2), F258A (Spindly ASB), and, importantly, was nearly abolished when all three motifs were mutated (Spindly ACCS) (Figure 1c,d). Dynein recruitment by Spindly results in stripping ^{26,43,51–53,58,59} In of Spindly from kinetochores upon microtubule attachment accordance with this, the low kinetochore dynactin levels in cells expressing Spindly motif mutants caused those mutants to be retained on metaphase kinetochores (Figure 1e,f). Strikingly, Spindly ΔCCS, which was the most compromised in recruiting dynein (Figure 1e, f), frequently appeared as a large structure bridging the two sister kinetochores (Figure 1e). This resembled fully expanded kinetochores of nocodazoletreated cells (compare example 4 in Figure 1a with inset of Spindly $^{\Delta CCS}$ in Figure 1e). A similar phenotype was observed by deleting the N-terminal 65 amino acids of Spindly (Spindly $^{\Delta N}$) that contain the CC1 and CC2 boxes (Figure 1g). These findings suggest that Spindly, via the recruitment of dynein, is essential to compact previously expanded kinetochores (Figure 1h).

Kinetochores expand by forming a structurally stable fibrous corona

Having established the mechanism of kinetochore compaction, we next examined the molecular basis for the kinetochore expansion. Quantitative immunofluorescence imaging of kinetochores in nocodazole-treated cells showed that the RZZ complex, Spindly and MAD1 occupied the largest volumes (often showing partial or complete

encirclement of the sister kinetochores), followed by the KMN network and the CCAN (Figure 2a,b). This is in agreement with studies reporting that different kinetochore modules expand to different extents and inversely to their proximity to the chromosome's axis ^{17,18,24}. RZZ, Spindly and MAD1 are thought to be part of the fibrous corona, which in electron micrographs appears as a thick electron-dense structure that assembles on the outer kinetochore in early mitosis or upon addition of microtubule-deplymerizing drugs ^{17,19,21,22,25,28,30,33,36,42,55}. We thus deemed it likely that kinetochore expansion observed in immunofluorescence images represents corona assembly. RZZ, Spindly and MAD1 may therefore constitute the core fibrous corona meshwork and/or regulate its formation. This was supported by the observation that brief (20') CDK1 inhibition caused the detachment from the inner parts of the kinetochore of intact, crescent-shaped rods or extended filaments close to the kinetochores that contained Spindly, ZW10, MAD1 and CENP-E (Figure 2c). These data support the hypothesis that the expanded kinetochore module observed in the absence of microtubules is the fibrous corona and that it is a stable structure with a specific architecture, composition and regulation, distinct from other outer kinetochore modules such as the KMN network (Figure S3a).

Spindly and RZZ are essential for corona formation

Immunofluorescence imaging and electron microscopy showed absence of kinetochore expansion and absence of a fibrous corona, respectively, in ZW10 RNAi cells (Figure 3a,c,e,g and Figure S3b-d), further establishing that kinetochore expansion equates to formation of a fibrous corona. Interestingly, ZW10 RNAi also reduced the volumes occupied by the KMN network member HEC1 and the CCAN

member CENP-C (Figure 3a, c). RZZ is thus essential for fibrous corona formation, and controls the (limited) expansion of inner- and outer- kinetochore regions.

We next turned our attention to Spindly, and reasoned that if dynein recruitment were its predominant contribution to kinetochore size dynamics, Spindly depletion should phenocopy severe Spindly motif mutants, i.e. persistently expanded kinetochores due to an inability of microtubules to cause shedding of the corona (Figure 1g). In contrast, however, volume measurements and electron microscopy revealed that cells depleted of Spindly were unable to expand kinetochores and had severely compromised fibrous coronas (Figure 3b,d,f,g and Figure S3b-d). Spindly is therefore essential not only for corona disassembly through dynein binding, but also for corona assembly.

Spindly stimulates RZZ-Spindly polymerization in vitro and in vivo

RZZ is necessary for the recruitment of several fibrous corona components ^{22,26,37,38,40,53,58,60}, and its structure resembles that of coat scaffolds, which can self-assemble into polymeric states ^{35,44}. We therefore asked if RZZ could assemble ectopically into polymeric structures, which may be surmised to promote the expansion of the corona. Purified, recombinant RZZ, which assembles with 2:2:2 (ROD:Zwilch:ZW10) stoichiometry³⁵, did not oligomerize, as assessed by direct visualization of mCherry-ROD by fluorescence microscopy (Figure 4a). Addition of purified farnesylated Spindly (Spindly^{FAR}), however, caused spontaneous oligomerization into filamentous structures at 30°C (Figure 4c, Movie S1). *In vitro* filament formation of RZZ-Spindly (RZZS) complexes could be prevented by addition

of detergent, suggesting that hydrophobic interactions underlie the assembly reaction (Figure 4a). Notably, RZZS oligomerization in the presence of GFP-Spindly-coated agarose beads resulted in stable association of filaments with the beads (Figure 4b), in a configuration reminiscent of the fibrous corona at kinetochores.

Expression of Spindly^{FL} (full length) in interphase cells had no discernable effect on RZZ complexes, but expression of Spindly^{ΔN} caused spontaneous formation of cytoplasmic filaments containing ZW10, Zwilch and ROD (Figure 4c, S4a and Movie S2). These filaments strongly resembled the ones formed *in vitro* by purified components and were strikingly similar to the detached coronas apparent after brief CDK1 inhibition (see Figure 2c). Cytoplasmic filament formation was abolished upon ZW10 RNAi or mutation of the Spindly CAAX box (C602A), showing it relied on RZZ and RZZ's interaction with farnesylated Spindly (Figure 4c). Spindly^{ΔCCS}, which like Spindly^{ΔN} is unable to bind dynein/dynactin, did not induce filaments (Figure 4c), showing that the ability of Spindly^{ΔN} to cause RZZS oligomerization is unrelated to dynein/dynactin binding.

Spindly adopts a 3D configuration that impairs its ability to promote RZZS oligomerization

Collectively, these results indicate that Spindly can promote the formation of a stable RZZS meshwork, and that this process is greatly enhanced by deletion of the N-terminal 65 amino acids of Spindly. To examine how removal of the N-terminal 65 amino acids of Spindly promotes cytoplasmic RZZ filament formation, we next wished to obtain a better understanding of Spindly's structure and regulation. Residues 1-440 of Spindly are predicted to form a coiled-coil structure while the C-terminal region is

likely disordered (Figure 1b) ^{35,43,54}. Small Angle X-ray Scattering (SAXS) analysis of a recombinant Spindly fragment encompassing 1-440 showed intramolecular dimensions (D_{max}, length: 260Å, R_c, width: ~22Å) of which in particular the R_c did not fit with that of di-, tetra-, or hexameric coiled-coil models (R_c: 6.2, 9.6 and 12.4 Å, respectively) (Figure 5a). Spindly^{FL} showed an overall shape and dimensions similar to Spindly¹⁻⁴⁴⁰ (Figure 5a). Negative stain electron microscopy (EM) on single Spindly¹⁻⁴⁴⁰ and Spindly^{FL} particles confirmed the SAXS analyses: Spindly adopts elongated shapes with some characteristic "bends" along its length (Figure 5b). The overall dimensions from EM confirmed SAXS measurements, and suggested that purified Spindly may not form a typical coiled-coil. Finally, SAXS and SEC-MALLS (Size exclusion chromatography-multi angle laser light scattering) experiments indicated that Spindly¹⁻⁴⁴⁰ and Spindly^{FL} are dimeric and trimeric, respectively, in solution (Figure 5a and SSb,c).

Cross-link mass spectrometry of Spindly^{FL} supported the possibility of a more complicated Spindly fold, and revealed the existence of various long range intramolecular interactions (Figure S5c). Several of these interactions occurred around a cluster of cross-linked lysines at positions 276, 278, 283 and 284 (Figure S5c). Size exclusion chromatography showed interaction between an N-terminal fragment of Spindly (Spindly¹⁻²⁵⁰) and a Spindly fragment encompassing C-terminal sequences (Spindly²⁵⁰⁻⁶⁰⁵) (Figure 5c), which was verified by Surface Plasmon resonance (SPR) (Figure 5d). This further verified intramolecular Spindly interactions and implicated the N-terminal region in these. No interaction was observed between Spindly¹⁻²⁵⁰ and Spindly³⁰⁶⁻⁶⁰⁵ (Figure 5c), suggesting amino acids 259-305 have a crucial role in longrange interactions with the N-terminal Spindly helices. This was consistent with the

position of the lysine cluster in the cross-link mass spectrometry analysis (Figure S5c). Notably, while a truncated Spindly lacking the N-terminal 274 amino acids (Spindly $^{\Delta274}$) retained ability to form cytoplasmic filaments in cells, this was abolished by additional removal of 13 amino acids (Spindly $^{\Delta287}$) (Figure 5e). Thus, this region that is crucial for Spindly intramolecular interactions, is also important for *in vivo* filament formation.

Together, these data support the hypothesis that the Spindly N-terminal region imposes an auto-inhibited configuration that precludes RZZ-Spindly oligomerization.

Release of Spindly autoinhibition promotes its interaction with RZZ

We next performed SPR analyses with immobilized, purified RZZ to examine interactions of recombinant Spindly versions with the RZZ scaffold. In the absence of C-terminal farnesylation, ~2 molecules of Spindly^{FL} weakly bound one (dimeric) molecule of RZZ with a K_D of ~1 μ M (Figure 6a). Farnesylation had little impact on overal interaction affinity *in vitro* but increased the number of Spindly molecules accumulating on RZZ. Similar observations were made with an alternative source of Spindly protein (Figure S6a). The farnesyl group thus appears to target Spindly to multiple sites on RZZ or to other Spindly molecules already on RZZ, under these conditions. A version of Spindly lacking the N-terminal helices (Spindly⁵⁴⁻⁶⁰⁵) associated with RZZ with higher affinity (~0.7 μ M) and at higher stoichiometries: At least four molecules of Spindly could associate with RZZ. Notably, farnesylation no longer impacted interactions between RZZ and Spindly⁵⁴⁻⁶⁰⁵ (Figure 6a). Removal of

the N-terminal helices of Spindly therefore can achieve high occupancy and higher affinity RZZ-Spindly interactions independently of farnesylation. Based on our observation that Spindly^{AN} farnesylation was required for cytoplasmic filament formation (see Figure 4c) but was dispensible for Spindly^{AN}-RZZ interactions when RZZ was on the SPR chip, we reasoned that farnesylation might facilitate initial Spindly ^{ΔN}-RZZ interactions that promote additional, farnesyl-independent ones. To test this, we examined if RZZ complexes concentrated on mitotic kinetochores can recruit unfarnesylated Spindly molecules. As expected, mutation of the farnesylated cysteine (C602A) or treatment with the farnesyl transferase inhibitor Lonafarnib (SCH66336) 61 prevented Spindly^{FL} localization and fibrous corona formation (Figure 6b-e and S6c,e). Removal of the N-terminal helices (Spindly^{AN}) not only rescued the Spindly localization defect caused by impairing farnesylation, but also rescued fibrous corona formation (Figure 6b-d). Rescue of kinetochore localization was dependent on the 275-287 region and was independent of dynein (Figure S6c-f). We thus conclude that releasing the auto-inhibited state of Spindly, aided by C-terminal farnesylation, is necessary for formation of the fibrous corona.

Activity of the kinetochore kinase MPS1 promotes fibrous corona expansion

Our data thus far suggest a mechanism for fibrous corona formation in which release of Spindly auto-inhibition enables direct interactions with RZZ and other Spindly molecules to form the corona meshwork. We deemed it likely that such a release would occur at or near kinetochores in mitosis, and reasoned that kinetochore-localized kinases are likely candidates to cause the triggering event. Whereas addition of small molecule inhibitors of Aurora B or PLK1 prior to mitotic entry caused no

evident problem with fibrous corona formation in nocodazole/MG132-treated cells (Figure 7a,b), Spindly localization and the corona were substantially affected by MPS1 inhibition, to an extent similar to Spindly depletion (Figure 7b,c-f). Importantly, Spindly kinetochore levels and fibrous corona formation were rescued in MPS1-inhibited cells by deletion of the N-terminal helices of Spindly (Figure 7c-f). Thus, MPS1 is required for the release of the auto-inhibited state of Spindly. To examine if MPS1 is sufficient to trigger Spindly-dependent RZZ oligomerization, we targeted MPS1 to cytoplasmic Spindly by co-expressing GFP-Spindly and DARPin GFP-MPS1 (A200). This MPS1 variant is unable to localize to kinetochores (B2) but can bind Spindly's GFP tag by virtue of a GFP-binding DARPin moiety (B3). Strikingly, when bound by active but not inactive MPS1, wild-type GFP-Spindly was able to induce interphasic filament formation (Figure 7g), just like Spindly (AN) could in the absence of MPS1 activity (see Figure 4a).

We conclude that MPS1 kinase activity triggers the ability of Spindly to promote RZZ-Spindly oligomerization by releasing an inhibitory intra-molecular Spindly interaction.

The fibrous corona engages in extensive lateral microtubule interactions

The persistent presence of an expanded fibrous corona in cells expressing Spindly $^{\Delta CCS}$ and Spindly $^{\Delta N}$ provided a means to examine the function of the corona and the importance of its shedding in late mitosis. Imaging of congressed chromosomes in Spindly $^{\Delta N}$ -expressing cells showed that the corona of sister chromatids engaged with the sides of microtubules (Figure 8a) and had lower Astrin levels, indicative of fewer mature end-on kinetochore-microtubule interactions (Figure S7a,b) 64 . The laterally interacting microtubules were positive for PRC1 (Figure S7c) and thus likely were anti-

parallel bridging fibers ('b' in Figure 8a) ^{65–67}. Cold treatment depolymerized PRC1 positive bundles (Figure S7c) and revealed the additional presence of stable, end-on microtubule interactions (Figure 8b and S7c). Super-resolution imaging by Expansion Microscopy (ExM) ⁶⁸ confirmed the co-occurrence of lateral and end-on attachments on the same kinetochore ('l' and 'e' in Figure 8c) and showed an extensive fibrous corona surface capable of lateral microtubule interactions (Example 1 in Figure 8c). Live imaging of mCherry-Tubulin and GFP-Spindly^{ΔN} further showed that the corona maintained interactions with the lattices of microtubules that go through rounds of catastrophe and rescue ('Cat' and 'Res' in Figure 8d and Movies S3, S4), and with the lattices of microtubules that move away from the sister kinetochores (Figure 8d and Movies S3 and S4). The expanded fibrous corona thus efficiently captures and maintains interactions with the lattices of dynamic microtubules (Figure 8e).

Proper bi-orientation and chromosome segregation rely on dynamic kinetochore size regulation

We next examined the importance of fibrous corona shedding (i.e. kinetochore compaction) once microtubules are captured. We noticed that the expanded coronas of sister chromatids in Spindly $^{\Delta N}$ cells often appeared fused and frequently formed merotelic attachments ('m' in Figure 8b and Example 2 of Figure 8c). Live imaging captured these events: During successive cycles of capture-release, we observed short-lived events in which fibrous coronas simultaneously bound to dynamic microtubule plus ends and to stable k-fibers, briefly causing merotelic attachments (see second 15 in Example 1 of Figure 8d and Movie S3). Merotely is an important cause for chromosome segregation errors in healthy and cancerous cells 34,69,70 . To

examine if inability to shed the corona increases the frequency of chromosome

segregation errors, we live imaged Spindly mutant cells undergoing anaphase, induced

by treatment with an MPS1 inhibitor to bypass the SAC silencing defect resulting from

absence of kinetochore dynein ^{51,52}. We analyzed only cells that achieved full

chromosome alignment so as not to bias for alignment problems associated with

persistent coronas (Figure 8f). In contrast to cells expressing Spindly variants that

allowed corona shedding (Spindly^{FL} and Spindly^{ΔSB}), cells expressing Spindly^{ΔCCS}

showed a high rate of lagging chromosomes in anaphase, indicative of persistent

merotelic attachments.

Together these experiments show that the fibrous corona promotes lateral

microtubule capture, and that subsequent corona shedding is important to establish

proper, amphitelic attachments of sister chromatids.

DISCUSSION

While the understanding of the assembly mechanisms of the core kinetochore has

seen great progress in recent years ^{2,71}, the nature of the fibrous corona has remained

enigmatic. Here we have shown that the fibrous corona is composed of an RZZ-Spindly

filamentous network. The properties of this network resemble those of other cellular

polymers that assemble through a 'collaborative' mechanism ⁷². Collaborative

filaments form on a supporting matrix, such us DNA or membranes, and frequently

use hydrophobic interactions as a driving force for their assembly. By analogy, RZZ

assembles on the core kinetochore in a manner dependent on BUB1 and KNL1 39,73,

and subsequently forms a stable filamentous structure that is maintained even when

the core kinetochore is lost after inhibiting CDK1. Possibly, recruitment of BUB1 to the

KMN network creates a specific physical-chemical environment that is prone to shielding by the hydrophobic fibrous corona, causing the formation of rings embracing the centromere region, much like coating proteins covering vesicles ³⁵.

The RZZ-Spindly module provides the unique property of adapting kinetochore size to microtubule attachment status ¹⁷: In the absence of microtubules, RZZ-Spindly polymerize into a structurally stable filamentous meshwork, while microtubule binding promotes shedding of the meshwork by Spindly-dependent dynein/dynactin activity. This Spindly-centered mechanism promotes error-free chromosome birorientation by enabling efficient microtubule capture and preventing merotely by ensuring timely kinetochore compaction. Based on our observations we propose the following mechanistic model (Figure 8g): Before localizing to kinetochores, Spindly exists in an auto-inhibited state, where the N-terminal helices mask a surface required for polymerization of RZZ. Farnesylation-dependent targeting of Spindly to kinetochores by virtue of an initial interaction with ROD then enables kinetochorelocalized MPS1 activity to release Spindly auto-inhibition and stimulate RZZ-Spindly polymerization ^{35,43,51,54,55}. This regulation can be entirely bypassed if the N-terminal helices of Spindly are removed, suggesting that this region acts as a focal point of intramolecular regulation. Interestingly, intramolecular interactions similar to those in Spindly have been observed for the related dynein adaptor, BICD2. In BICD2, cargo binding releases auto-inhibition, thereby allowing its N-terminal region to bind dynein-dynactin ⁷⁴. Albeit BICD2 and Spindly share conserved short sequence motifs that affect dynein binding, our SAXS and EM data suggest an indirect role of the sequence motifs in dynein binding, consistent with a recent structural study 75. This raises interesting questions about how the structural organization of Spindly facilitates regulated RZZ interactions. It remains to be tested if the conformational change of Spindly is necessary to bind or activate dynein at the kinetochore. Interestingly, Spindly binds dynein efficiently but is quite heterogeneous in its ability to stimulate dynein mobility ⁵⁹, perhaps due interference from long-range intramolecular interactions.

Several proposed events in our model of Figure 8g require further investigations. For example, the details of how MPS1 promotes release of Spindly auto-inhibition are unknown, but given that MPS1 can stimulate Spindly-RZZ filamentation in interphase it likely involves phosphorylation of one or more components of the RZZ-Spindly complex and/or one of its binding partners. Likewise, the mechanisms by which the corona binds the microtubule lattice, and the type of microtubule attachments that trigger shedding of the fibrous corona are unknown. Several microtubule-binding proteins including CENP-E, CENP-F, CLASP1, CLASP2 and dynein show localization consistent with presence at the fibrous corona ^{17,19,20,27,30,76}. While dynein is important for chromosome congression ^{16,77}, our data indicate that it is not essential for the interaction of kinetochores with microtubule walls. It will be important to examine which of the other microtubule-binding proteins is the primary lattice-binding activity of the fibrous corona. CENP-E is a prime candidate ^{78,79}. Dynein is, on the other hand, crucial for shedding of the corona ^{26,45,51,52,80}. This is likely mediated by end-on microtubule attachments, because lateral interactions are not sufficient to remove MAD1 from kinetochores ⁷⁹. How such end-on interactions can form in the presence of the fibrous corona is unclear, but they may not need to be of the stable kind that is mediated by the NDC80 complex. We can envision [and have observed (CS and GJPLK, unpublished observations)] transient interactions between microtubule plus-ends and the fibrous corona that are sufficient to cause some corona shedding. This would not only expose NDC80 complexes, but would also remove the reported RZZ-dependent inhibition of their microtubule binding activity ^{81,82}. NDC80 complexes can then engage in more stable end-on microtubule interactions that further stimulate dynein activity. Fibrous corona removal could thus promote progressive maturation of end-on attachments in a positive feedback. Such a scenario combines efficient lateral capture of microtubules with efficient transition to end-on attachments while minimizing the risk of merotelic attachments and subsequent chromosome segregation errors.

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Author Contributions

C.S. and G.J.P.L.K. conceived the idea for this project. C.S., G.J.P.L.K., M.A. A.P., J.K. and A.M designed the experiments and interpreted the data. C.S. performed the *in vivo* experiments. M.A. and J.K. performed the *in vitro* experiments with the help of A.F.. J.F. and J.K. performed and analysed the electron microscopy experiments in cells with the help of C.S.. V.G. performed the cross-linking experiments. E.T. performed the comparative sequence analysis. R.M. and J.M.C performed the electron microscopy of Spindly. C.S. and G.J.P.L.K. wrote the manuscript with the help of A.P. and A.M. and the input of the rest of authors.

FIGURE LEGENDS

Figure 1. Spindly recruits dynein to compact kinetochores upon microtubule attachment. (a) Immunofluorescence images of ZW10 and HEC1 in HeLa cells treated with nocodazole or STLC. A, Attached; U, Unattached. (b) Overview of the secondary structure of human Spindly with predicted coiled-coils (grey bars) and disordered regions, and with sequence logos of four conserved motifs (supplementary sequences 1). (c,d) Representative images (c) and quantification (d) of nocodazole-treated HeLa cells transfected with siRNAs to Spindly and expressing the indicated GFP-Spindly variants, and immuno-stained for the indicated antigens. The graph in D shows mean kinetochore intensity (±s.d.) normalized to the values of Spindly^{FL} from at least four

independent experiments. Each dot represents one cell and at least 20 cells were quantified per experiment and condition. Asterisks indicate significance (analysis of variance with Tukey's multiple comparison test). ****P < 0.001. (e,f) Representative images (e) and quantification (f) of metaphase HeLa cells transfected with siRNA to Spindly and expressing the indicated GFP-Spindly variants, and immuno-stained for the indicated antigens. The graph shows the mean fold change in kinetochore intensity (±s.d.) normalized to the values of Spindly^{FL} from five independent experiments. Each dot represents one cell and at least 20 cells were quantified per experiment and condition. Asterisks indicate significance (analysis of variance with Tukey's multiple comparison test). ****P < 0.001. (g) Immunofluorescence images of HeLa cells transfected with siRNA to Spindly and treated or not with doxycycline (Dox) to induce the expression of GFP-Spindly variants, and immuno-stained for the indicated antigens. (h) Representative images of the morphology of Spindly and CENP-C in metaphase kinetochores expressing the indicated GFP-Spindly variants. Stripping activity is based on ability of Spindly variants to recruit dynein/dynactin.

Figure 2. Kinetochores expand by forming a structurally stable fibrous corona. (a,b) Representative images (a) and volume quantification (b) of the indicated GFP-tagged proteins in expanded kinetochores of HeLa cells entering mitosis in the presence of nocodazole. Maximum expanded kinetochores were selected based on ZW10 staining. The graph shows the mean kinetochore volume (\pm s.d.) of at least three independent experiments. Each dot represents one kinetochore and at least 10 kinetochores per experiment were quantified. (c) Images of kinetochores of HeLa cells entering mitosis

in the presence of nocodazole and subsequently treated with RO-3306 for 20 minutes,

and immunostained with the indicated antibodies. Color codes in B and C correspond

to behavior of protein after brief CDK1 inhibition.

Figure 3. Spindly and RZZ are essential for corona formation. (a-d) Representative

images (a, b) and volume quantification (c, d) based on the indicated antigens of

expanded kinetochores of nocodazole-treated HeLa cells transfected with siRNA to

ZW10 (a,c) or Spindly (b,d). The graph shows the mean kinetochore volume (±s.d.) of

at least three independent experiments. Each dot represents one kinetochore and at

least 10 kinetochores per experiment were quantified. Asterisks indicate significance

(analysis of variance with Tukey's multiple comparison test). ****P < 0.001; n.s., not

significant. (e-g) Electron micrographs (e,f) and surface quantification (g) of the fibrous

corona of nocodazole-treated cells depleted of ZW10 (e) or Spindly (f). The fibrous

corona is highlighted in green and the outer plate in blue. The graph shows the mean

surface of the fibrous corona (±s.d.) normalized to the value of the control from two

(control and siSpindly) or one (siZW10) experiment. At least 50 kinetochores per

condition and experiment were imaged. Quantification of the protein depletion

efficiency are shown in Figure S3c,d. Each dot represents a single kinetochore.

Asterisks indicate significance (Student's t-test, unpaired). ****P < 0.001. Control is

siGAPDH.

Figure 4. Spindly stimulates RZZ-Spindly polymerization in vitro and in vivo. (a)

Fluorescence microscopy of purified mCherry-RZZ alone or in the presence of

recombinant farnesylated Spindly (Spindly FAR) and incubated as indicated. See also

Movie S1. **(b)** GFP-Spindly immobilized on beads and incubated in the presence of mCherry-RZZ and farnesylated Spindly (RZZS). Control is empty beads. The intensity level of the green channel in the control were enhanced to allow the comparison with GFP-Spindly. **(c)** Immunofluorescence images of interphase HeLa cells expressing the

indicated versions of GFP-Spindly and treated and stained as indicated. See also

Movie S2.

Figure 5. Spindly adopts a 3D configuration that impairs its ability to promote RZZS

oligomerization. (a) Normalized Kratky plot (left graph) and paired distance

distribution (right graph) of purified Spindly^{FL} and Spindly¹⁻⁴⁴⁰. The intersection of the

red dotted lines represents the peak for a globular protein irrespective of its molecular

weight. Comparison with dimeric, tetrameric and hexameric coiled-coil models of

similar size are shown in shades of grey. Rg, radius of gyration; Rc, cross-sectional

radius of gyration; D_{max}, maximum inter-particle distance. **(b)** Representative class

averages from the negative stain EM of indicated Spindly variant proteins. (c) Elution

profiles and SDS-PAGE of SEC experiments on Spindly¹⁻²⁵⁰ and Spindly²⁵⁰⁻⁶⁰⁵ or their

stoichiometric combination. AU, arbitrary units. (d) Surface Plasmon Resonance (SPR)

analyses of the interaction between immobilized Spindly^{259-C} (blue) or Spindly^{306-C}

(red) and soluble Spindly¹⁻²⁵⁰. The response (y-axis) was normalized to the molecular

weight of the analyte to yield stoichiometry of binding. (e) Immunofluorescence

images of Spindly in interphase HeLa cells expressing the indicated truncations of GFP-

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Spindly.

Figure 6. Release of Spindly autoinhibition promotes its interaction with RZZ. (a) SPR analyses of the indicated Spindly variant proteins. The response (y-axis) was normalized to the molecular weight of the analyte to yield stoichiometry of binding. (b-e) Immunofluorescence (b,c) and quantification of kinetochore levels (d) or volumes (e) of Spindly in HeLa expressing GFP-Spindly^{FL} or GFP-Spindly^{ΔN} and treated with nocodazole and the farnesyl transferase inhibitor Lonafarnib (Lon). The graph shows the mean fold change in kinetochore intensity (\pm s.d.) normalized to the values of Spindly^{FL} from three independent experiments. Each dot in D represents one cell and at least 30 cells per experiment and condition were quantified. Each dot in E represents one kinetochore and at least 10 kinetochores per experiment were

quantified. Asterisks indicate significance (analysis of variance with Tukey's multiple

comparison test). ****P < 0.001.

Figure 7. Activity of the kinetochore kinase MPS1 promotes fibrous corona expansion. (a) Timeline of the treatments with kinase inhibitors and nocodazole (Noco) of the experiments shown in b-f. (b) Representative images of ZW10 immunostainings of cells treated as indicated in A. The intensity levels of the zoom ins (on right) were equalized to facilitate the direct comparison of the size of the kinetochores. (c,d) Representative images (c) and quantification (d) of kinetochore localization of the indicated GFP-Spindly variants in HeLa cells treated with nocodazole and Cpd-5 as indicated in a. The graph shows the mean fold change in kinetochore intensity (±s.d.) normalized to the values of Spindly^{FL} from three independent experiments. Each dot represents one cell and at least 30 cells per experiment and condition were quantified. Asterisks indicate significance (analysis of variance with

Tukey's multiple comparison test). ****P < 0.001. **(e,f)** Representative images (e) and volume quantification (f) of immunostained kinetochores of HeLa cells expressing the indicated versions of GFP-Spindly and treated as indicated in a. To measure the volume, imaging acquisition was set to obtain similar mean intensity levels for the different conditions. The graph shows the mean kinetochore volume (\pm s.d.) of at least three independent experiments. Each dot represents one kinetochore and at least 10 kinetochores per experiment were quantified. Asterisks indicate significance (analysis of variance with Tukey's multiple comparison test). ****P < 0.001. **(g)** Images of HeLa cells overexpressing GFP-Spindly^{FL} and an active (WT) or kinase dead (KD) version of mCherry-MPS1^{Δ 200} targeted to GFP-Spindly by DARPin^{GFP}.

Figure 8. Expanded kinetochores engage in extensive lateral microtubule interactions and prevent bi-orientation. (a,b) Representative single z-plane immunofluorescence images of HeLa cells in metaphase, transfected with siRNA to Spindly and expressing the indicated GFP-Spindly variants and mCherry-tubulin. Cells were fixed at 37°C (a) or after cold-shock (b). 'b', bridging fibers; 'k', k-fibers; 'm' merotelic attachment. (c) Two examples of ExM (expansion microscopy) images of HeLa cells transfected with siRNA to Spindly and expressing GFP-Spindly^{AN} and mCherry-tubulin. Single z-plane images show pairs of enlarged kinetochores with simultaneous end-on (e) and lateral (I) attachments (example 1) or end-on (e), lateral (I), and merotelic (m) attachments (example 2). (d) Live-cell imaging of HeLa cells transfected with siRNA to Spindly and expressing GFP-Spindly^{AN} and mCherry-tubulin. 'Cat', catastrophe; 'Res', rescue. Maximum projections of several z-planes are shown. See also Movies S3 (Example 1) and S4 (Example2). (e) Cartoon summarizing the

interaction of compacted or enlarged kinetochores with microtubule fibers. **(f)** Quantification of chromosome segregation errors by live-cell imaging of meptaphase HeLa cells transfected with siRNA to Spindly, expressing the indicated GFP-Spindly variants, and treated with Cpd-5. The bar graph shows the mean percentage (\pm s.d.) of cells showing lagging chromosomes of two independent experiments. Asterisk indicate significance (analysis of variance with Tukey's multiple comparison test). *P < 0.01; n.s., not significant. (FL, n=222 cells; Δ SB=179 cells; Δ CCS, n=167 cells). **(g)** Model of the mechanism of kinetochore expansion and compaction performed by the axis RZZ-Spindly-dynein and MPS1. See discussion for details.

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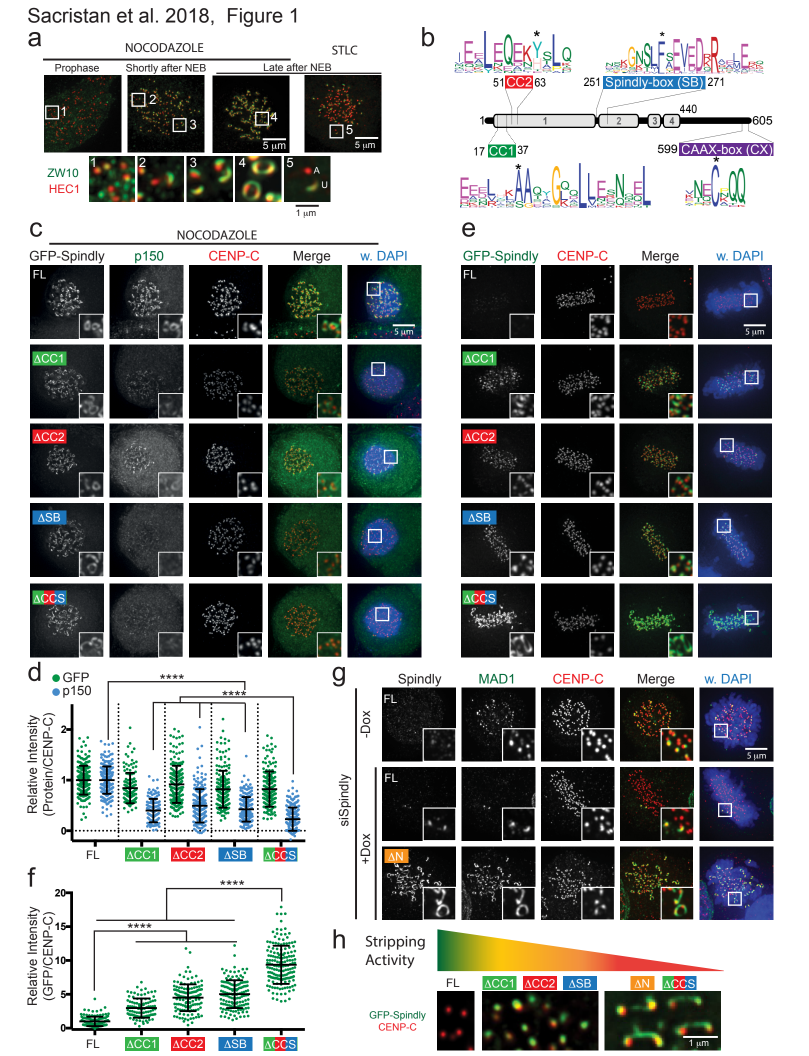
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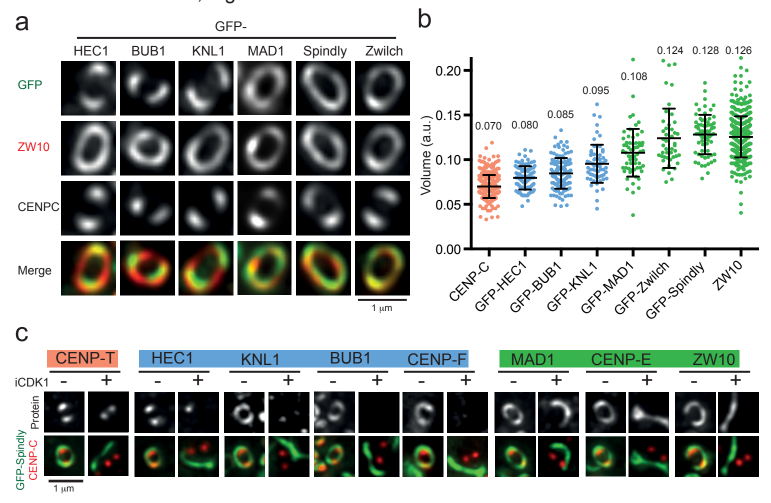
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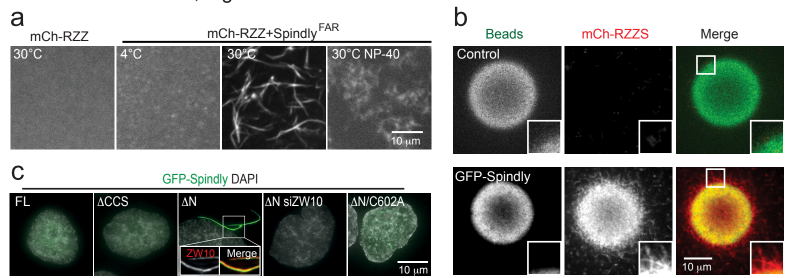


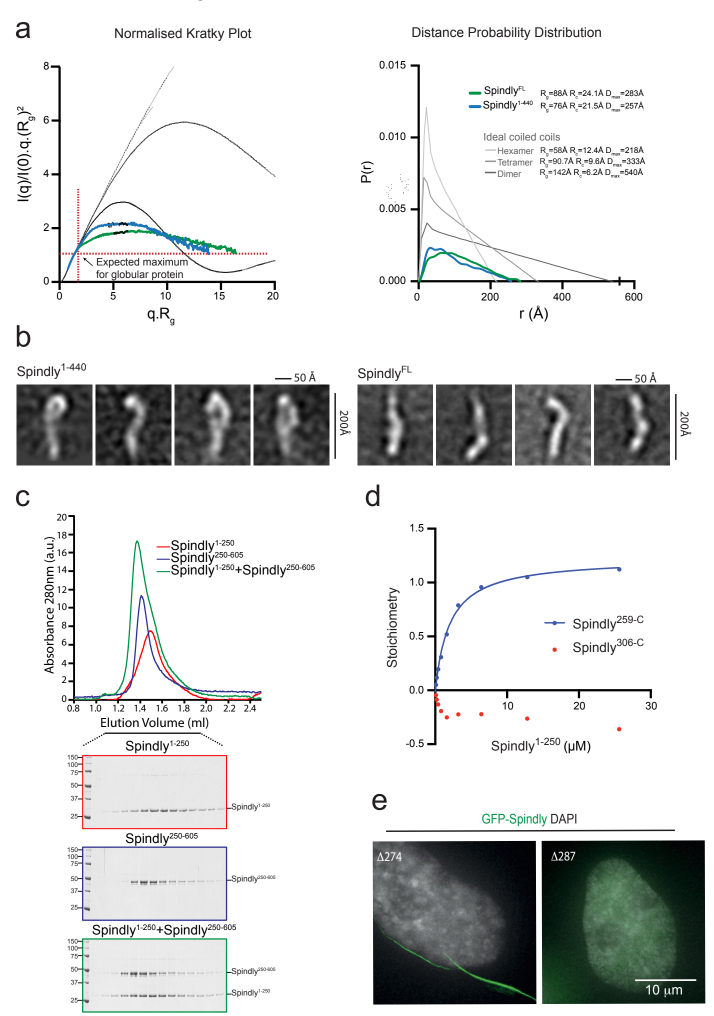
Sacristan et al. 2018, Figure 2



Sacristan et al. 2018, Figure 3 a ZW10 GFP-Zwilch HEC1 Merge ZW10 MAD1 CENP-C CENP-E CENP-C Merge Merge Control siZW10 b Spindly GFP-Zwilch HEC1 Merge Spindly MAD1 CENP-C Merge Control siSpindly d g C ControlsiZW10 ControlsiSpindly 0.25 0.25 Normalized Surface (a.u.) 0.092 0.119 0.20 0.20 0.085 Volume (a.u.) Volume (a.u.) 0.150.078 0.074 0.071 0.070 0.10 0.05 0.05 0.00 0.00 Control е siZW10 Control Control Fibrous corona Outer plate Fibrous corona Outer plate

Sacristan et al. 2018, Figure 4



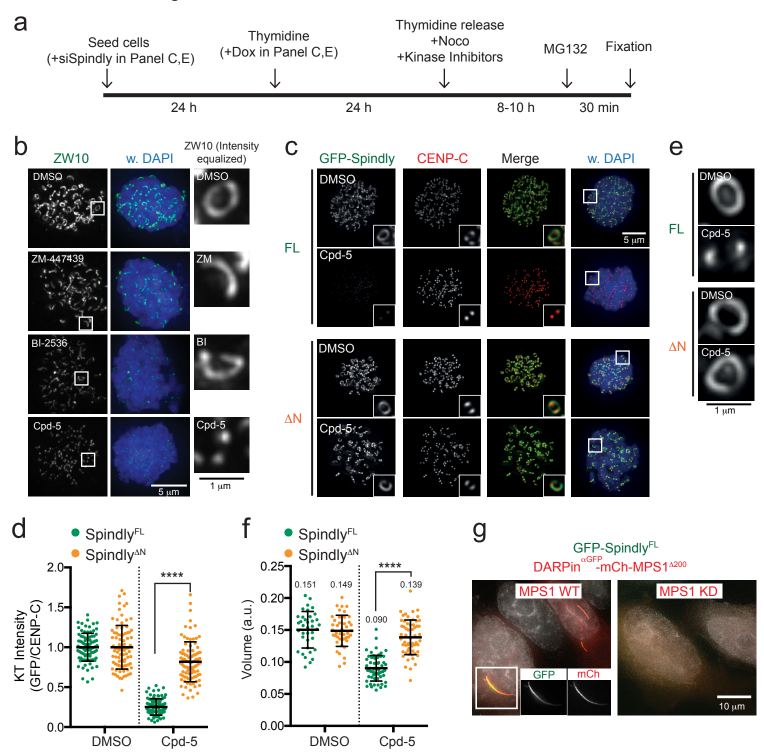


Sacristan et al. Figure 6 b a SpindlyFL w. DAPI **GFP-Spindly** CENP-C Merge Spindly^{FL} FAR **DMSO** Stoichiometry Spindly⁵⁴⁻⁶⁰⁵ Spindly⁵⁴⁻⁶⁰⁵ FAR 5 μm FL Lon 0.001 100 0.01 10 Spindly (µM) C GFP-Spindly ZW10 CENP-C DMSO DMSO FL Lon ΔN DMSO ΔΝ Lon 1 μm d е Spindly^{FL} Spindly^{FL} Spindly^{∆N} Spindly^{∆N} 0.25 0.129 2.0-0.136 0.20 Kinetochore Intensity (GFP/CENP-C) Volume (a.u.) 0.096 0.111 0.106 0.118 0.15 0.071 0.10 0.05 0.00 0.0 DMSO DMSO DMSO Lon Lon Lon

Spindly

ZW10

Sacristan et al. Figure 7



Sacristan et al. 2018, Figure 8 a b GFP-Spindly mCh-Tubulin (37°C) GFP-Spindly mCh-Tubulin (Cold-shock) HEC1 mCH Merge HEC1 mC Merge C d GFP-Spindly^{△N} mCh-Tubulin **ExM** Merge mCh-Tubulin GFP-Spindly^{△N} Example 1 Example 2 Example 1 0 sec mCh-Tubulin 0 sec mCh-Tubulin 15 sec 30 sec Example 2 50 sec 90 sec е Metaphase with compact kinetochores K-Fibers (k) Core kinetochore (HEC1) 1 μm Cpd5 0 min Cells with lagging chromosomes (%) Bridging Fibers (b) Metaphase with enlarged kinetochores Cpd-5 28 min Enlarged kinetochore Merotelic (m) End-on (e) FL ΔCCS ΔSB Lateral (I) g Lateral attachment Spindly MAD1 End-on attachment 1. The initial kinetochore 2. A conformational change of 3. The formation of Spindly, enabled by MPS1 recruitment of RZZ end-on attachments is determines the kinase activity, promotes coupled to kinetochore compaction through the RZZ-Spindly oligomerization maturation of the core and dynein recruitment. dynein-mediated kinetochore. shedding of the fibrous Increase in size enables high FC= fibrous corona efficiency capture of the wall of O= outer kinetochore corona. microtubules. I= inner kinetochore