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Dephosphorylation of the Ndc80 Tail Stabilizes Kinetochore-Microtubule Attachments via the Ska Complex

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SUMMARY

During cell division, genome inheritance is orchestrated by microtubule attachments formed at kinetochores of mitotic chromosomes. The primary microtubule coupler at the kinetochore, the Ndc80 complex, is regulated by Aurora kinase phosphorylation of its N-terminal tail. Dephosphorylation is proposed to stabilize kinetochore-microtubule attachments by strengthening electrostatic interactions of the tail with the microtubule lattice. Here, we show that removal of the Ndc80 tail, which compromises *in vitro* microtubule binding, has no effect on kinetochore-microtubule attachments in the *C. elegans* embryo. Despite this, preventing Aurora phosphorylation of the tail results in prematurely stable attachments that restrain spindle elongation. This premature stabilization requires the conserved microtubule-binding Ska complex, which enriches at attachment sites prior to anaphase onset to dampen chromosome motion. We propose that Ndc80 tail dephosphorylation promotes stabilization of kinetochore-microtubule attachments via the Ska complex and that this mechanism ensures accurate segregation by constraining chromosome motion following biorientation on the spindle.

Abstract

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D.K.C. and A.D. initiated the project; D.K.C conducted and analyzed the majority of experiments; B.P. performed SKA imaging and high-time-resolution chromosome dynamics assays; N.H. generated and analyzed the microtubule-binding domain deletion; L.L. performed the Aurora B mutant analysis; K.D.C. generated structural models and guided mutational and biochemical analysis; K.O. provided conceptual and experimental advice and helped guide the effort; the manuscript was prepared by D.K.C, K.O., and A.D., with input from all authors.



Phosphorylation by Aurora kinases regulates kinetochore-microtubule attachments during mitosis. Cheerambathur et al. show that in *C. elegans* one-cell embryos, Aurora phosphorylation of the Ndc80 complex, the major microtubule-binding factor at the kinetochore, controls kinetochore-microtubule attachments indirectly by regulating the recruitment of an effector, the microtubule-binding Ska complex.

Keywords

Centromere; Kinetochore; Chromosome Segregation; Mitosis; Cell Division; Ndc80 Complex; Ska Complex; Microtubule; Cell Polarity

INTRODUCTION

During cell division, the replicated genome is partitioned into daughter cells by interactions between the kinetochore, a multiprotein complex assembled on the centromere of mitotic chromosomes, and the spindle microtubules (Cheeseman, 2014). Accurate segregation requires bioriented kinetochore-microtubule attachments, in which sister chromatids are connected to microtubules from opposite spindle poles. Following initial capture by kinetochore-localized motors, kinetochores form end-on attachments to microtubules that are mediated by the conserved four-subunit Ndc80 complex (Cheeseman et al., 2006; Ciferri et al., 2008; Deluca et al., 2006; Wei et al., 2007). End-on kinetochore-microtubule attachments are initially dynamic to facilitate correct attachment geometry and chromosome alignment. These attachments are eventually stabilized, as best shown by reduced turnover of kinetochore microtubules in vertebrate cells (Kabeche and Compton, 2013). The suppression of kinetochore-microtubule dynamics after bi-orientation is associated with posttranslational modifications as well as alterations in kinetochore composition (Cheerambathur and Desai, 2014; Foley and Kapoor, 2012). Notably, misregulation of this step is thought to be the major source of chromosome segregation errors in cancer cells(Godek et al., 2014).

The microtubule binding activity of the Ndc80 complex resides in two distinct elements in the N-terminus of its Ndc80 subunit: a basic unstructured N-terminal tail with dispersed Aurora B kinase phosphorylation sites, and the adjacent Calponin Homology (CH) domain (Ciferri et al., 2008). High-resolution cryo-EM of Ndc80 complex-decorated microtubules indicates that the Ndc80 CH domain contacts the microtubule lattice (Alushin et al., 2012; 2010; Wilson-Kubalek et al., 2008). Mutations that disrupt this interface abrogate microtubule binding in vitro and impair attachments in vivo (Ciferri et al., 2008; Lampert et al., 2012; Sundin et al., 2011; Tooley et al., 2011). The N-terminal tail, whose deletion compromises *in vitro* microtubule binding, to a similar extent as CH domain mutations (Ciferri et al., 2008; Wei et al., 2007), is proposed to electrostatically interact with the negatively charged tubulin surface. However, the contribution of the Ndc80 tail to microtubule attachments in vivo differs across species. In mammalian cells, deletion of the N-tail destabilizes kinetochore-microtubule attachments, whereas, N-tail deletion does not obviously perturb chromosome segregation in one-cell C. elegans embryos and is not lethal in budding yeast (Cheerambathur et al., 2013; Guimaraes et al., 2008; Kemmler et al., 2009; Miller et al., 2008).

Phosphorylation of the Ndc80 tail, by Aurora family kinases, is implicated in regulating the dynamics of kinetochore-microtubule attachments. In mammalian cells, phosphorylation of Aurora target residues is high during early mitosis and low on bioriented chromosomes (DeLuca et al., 2011). Phosphorylation by Aurora kinase, or the introduction of phosphomimetic mutations in the Ndc80 tail, reduces microtubule binding affinity of the Ndc80 complex *in vitro* (Alushin et al., 2012; Cheeseman et al., 2006; Umbreit et al., 2012). In addition, phosphomimetic Ndc80 variants compromise kinetochore fiber stability and chromosome alignment in mammalian cells (Alushin et al., 2012; Guimaraes et al., 2008; Zaytsev et al., 2014). These observations have led to the proposal that N-tail phosphorylation by Aurora kinase regulates kinetochore-microtubule attachments by modulating the electrostatic affinity between the basic N-tail and the negatively-charged microtubule lattice in a graded manner (Zaytsev et al., 2015).

In addition to phosphorylation-based modulation of microtubule affinity, multiple additional factors, such as the Dam1 complex in fungi, the Ska complex in metazoans, and XMAP215 family proteins are known to enhance Ndc80's microtubule coupling *in vitro* and this cooperation is proposed to be integral to proper kinetochore-microtubule attachment dynamics *in vivo* (Miller et al., 2016; Schmidt et al., 2012; Tien et al., 2010). Although it is known that these microtubule-associated proteins depend on Ndc80 for their kinetochore localization, how this additional machinery collaborates with the Ndc80 complex to control the properties of microtubule attachments *in vivo* and whether phosphoregulation of the Ndc80 complex is involved is not well understood. Notably, neither the Dam1 complex nor the Ska complex interact with the Ndc80 complex requires the context of the microtubule lattice (Lampert et al., 2010; Schmidt et al., 2012; Tien et al., 2010).

Here, we employ sensitive assays in the one-cell *C. elegans* embryo to show that dephosphorylation of the N-tail alters kinetochore-microtubule attachment dynamics via the Ska complex. We find that, while the N-tail deletion has no effect on kinetochore-

microtubule attachments *in vivo*, abrogating N-tail phosphorylation suppresses attachment dynamics and prevents spindle elongation until the activation of a spindle-external force prior to anaphase. The suppression of attachment dynamics by the non-phosphorylatable tail is dependent on the Ska complex, indicating that N-tail phosphorylation maintains a dynamic kinetochore-microtubule interface by restraining the action of the Ska complex. Removal of the Ska complex prevents the dampening of chromosome motion that occurs prior to anaphase onset in wild-type embryos and compromises viability. These findings reveal a mechanism that acts after chromosome biorientation to suppress chromosome motion and ensure accurate segregation.

RESULTS

Subtle Kinetochore-Microtubule Attachment Defects Can be Detected *in vivo* by Analyzing Spindle Pole Separation Rates Prior to Anaphase

To investigate the role of NDC-80 N-tail phosphorylation, we established an *in vivo* assay to monitor the strength of kinetochore-microtubule attachments in the one-cell *C. elegans* embryo. This assay is based on a method we employed previously in which the kinetics of spindle pole separation following nuclear envelope breakdown (NEBD), measured by live imaging of embryos expressing GFP fusions to histone H2b and γ -tubulin (Fig. 1A), is used to infer the presence or absence of kinetochore-microtubule attachments (Desai et al., 2003; Oegema et al., 2001).

Spindle poles in embryos begin to separate ~50s after NEBD (Fig. 1B). The kinetics of spindle pole separation from this time point until anaphase onset (~200s after NEBD) is influenced by kinetochore-microtubule attachments, which resist spindle elongation forces. In control embryos, pole separation occurs at a rate of 2.0 μ m/min during the pre-anaphase interval (50s-200s after NEBD). In embryos depleted of NDC-80, chromosome segregation fails and spindle poles separate rapidly (4.5 μ m/min) during this interval (Fig. 1A,B & S1A, B). The 2.2-fold increase in pole separation velocity in *ndc-80(RNAi)* highlights the importance of NDC-80 -mediated attachments in restraining pre-anaphase spindle elongation. The effect of NDC-80 depletion also indicates that kinetochore-microtubule attachments on aligning chromosomes in wild type embryos are bearing load and are likely undergoing tension-induced microtubule polymerization to allow spindle elongation.

To determine whether the spindle pole separation assay could detect a reduction in affinity of the Ndc80 complex for microtubules, rather than absence of attachments, we engineered mutants in the NDC-80 CH domain that are predicted to reduce microtubule-binding affinity. Based on prior work (Ciferri et al., 2008; Tooley et al., 2011), we targeted 2 sets of residues for mutation. The first set includes 4 residues—K100, R125, K144 and H155—which form the 'Toe' that contacts the inter-tubulin interface (Fig. 1C) (Alushin et al., 2010). We refer to the second set of 2 residues, K66 and K96, as the 'Ankle' based on their positioning relative to the Toe residues (Fig. 1C).

To assess the effect of mutating CH domain residues on kinetochore microtubule attachments in vivo, we employed a previously described NDC-80 replacement system (Cheerambathur et al., 2013) to generate mutants in which the 'Toe', 'Ankle', or 'Toe +

Ankle' residues were mutated to alanine. In this approach, an RNA interference (RNAi)– resistant *ndc-80* transgene is inserted in single copy at a fixed genomic locus under the control of endogenous *ndc-80* regulatory sequences. A wildtype (WT) *ndc-80* transgene (encoding NDC-80^{WT}) fully rescued both NDC-80 depletion (Fig. 1B) and a *ndc-80* null mutant, and immunoblotting has confirmed that transgene-encoded WT and various engineered NDC-80 mutants express at comparable levels to endogenous NDC-80 (Cheerambathur et al., 2013).

To monitor phenotypes associated with the 'Toe', 'Ankle', or 'Toe + Ankle' affinity mutants, the transgenes were introduced into a strain expressing GFP::H2b and GFP:: γ -tubulin. Quantitative analysis of spindle pole separation kinetics after endogenous NDC-80 depletion revealed that the rate of pre-anaphase spindle pole separation for the CH^{Toe+Ankle} mutant (4.4 µm/min) was nearly identical to that for NDC-80 depletion ('No Transgene'; Fig. 1D). In contrast, the rates for the CH^{Toe} and CH^{Ankle} mutants were intermediate between the rates for NDC-80 depletion (Fig. 1D & S1A,B), which is consistent with prior work (Lampert et al., 2012; Tooley et al., 2011) suggesting that 'Toe' and 'Ankle' residues make independent contributions to the affinity of NDC-80 for microtubules. Notably, the CH^{Toe} mutation by itself resulted in a severe chromosome segregation defect, whereas the CH^{Ankle} mutation did not (Fig. 1E), indicating that, while both sets of residues are important for attachment strength/dynamics, the Toe residues are more important for biorientation and segregation than the Ankle residues.

Based on the analysis of these three NDC-80 CH domain mutants, we conclude that the spindle pole separation assay is sufficiently sensitive to detect reduced affinity of NDC-80 for microtubules *in vivo*, even when the effects are mild enough to not overtly disrupt chromosome segregation.

Deletion of the NDC-80 N-Tail Does Not Affect Load-Bearing Kinetochore-Microtubule Attachment Formation in One-Cell Embryos

Deletion of the basic NDC-80 N-tail (pI=11.6) substantially reduces the *in vitro* microtubule binding affinity of the reconstituted *C. elegans* NDC-80 complex, as also observed for Ndc80 complexes from other species (Cheerambathur et al., 2013; Ciferri et al., 2008; Lampert et al., 2012; Umbreit et al., 2012; Wei et al., 2007). Surprisingly, in prior work we did not observe a significant effect of deletion of the N-Tail (NDC-80^{Tail}) on kinetochore-microtubule attachments or on chromosome segregation in one-cell *C. elegans* embryos (Cheerambathur et al., 2013), which we confirmed in independent experiments (Fig. 2A).

To determine if there was a subtle kinetochore-microtubule attachment defect associated with absence of the NDC-80 N-Tail, we combined NDC-80 ^{Tail} with removal of kinetochore-localized dynein motor. Dynein is recruited to kinetochores by the Rod/Zwilch/ Zw10(RZZ)-Spindly module (ROD-1/ZWL-1/CZW-1 and SPDL-1 in *C. elegans*; Fig. 2B) and can be selectively removed from kinetochores by ROD-1 depletion. ROD-1 depletion leads to delayed formation of load-bearing attachments (reflected in premature pole separation that recovers prior to anaphase onset) and ~30% frequency of lagging anaphase chromatin in one-cell embryos (Fig. 2B,S2A). Even in the sensitized ROD-1 depletion, NDC-80 ^{Tail} behaved identically to NDC-80^{WT} (Fig. 2B,S2A). By contrast, the CH^{Ankle}

mutant of NDC-80 significantly enhanced the chromosome segregation and spindle pole separation phenotypes of ROD-1 depletion (Fig. 2B,S2A). We further assessed NDC-80 ^{Tail} by combining it with CH^{Toe} mutations in the same transgene. The combined Tail-CH^{Toe} NDC-80 mutant behaved identically to the CH^{Toe} mutant in the spindle pole separation assay (Fig. S2B,C), showing no enhancement. Again, this was in contrast to the enhancement observed with the combined CH^{Toe} and CH^{Ankle} mutations (Fig. 1D). Thus, following removal of kinetochore dynein, or in the presence of CH domain mutations, NDC-80 ^{Tail} behaves equivalently to NDC-80^{WT}.

We conclude that even with combinatorial inhibitions and a sensitive *in vivo* assay that is capable of identifying mutations that reduce the affinity of NDC-80 for microtubules, NDC-80 ^{Tail} behaves similarly to NDC-80^{WT}, suggesting that the N-tail does not make a significant contribution to the affinity of NDC-80 for microtubules at kinetochores in the one-cell *C. elegans* embryo.

Mutation of Aurora Phosphorylation Sites in the N-Tail Restrains Spindle Elongation Until Activation of the Ga Pathway

The C. elegans NDC-80 N-tail harbors four Aurora kinase phosphorylation sites that are conserved in related species (Fig S2D). Mutation of these four sites to alanine abrogates Aurora kinase-mediated phosphorylation of the reconstituted NDC-80 complex, and mutation of these sites to aspartate significantly reduces NDC-80 complex microtubule binding affinity in vitro (Cheerambathur et al., 2013; Cheeseman et al., 2006). To address the role of these Aurora target sites in vivo, we analyzed the consequences of preventing phosphorylation by changing them to alanines (NDC-80^{4A}). In contrast to NDC-80^{Tail} (Fig. 2A), NDC- 80^{4A} exhibited a striking phenotype in the spindle pole separation assay (Fig. 2C) ---pre-anaphase spindle pole separation was significantly inhibited, with almost no spindle elongation between 50s and 130s after NEBD. To determine which Aurora kinase phosphorylates these residues in vivo, we analyzed spindle pole separation in a temperaturesensitive mutant of Aurora B/AIR-2 (Severson et al., 2000; Lewellyn et al., 2011) or following depletion of the Aurora A/AIR-1 activator TPX2/TPXL-1 (Lewellyn et al., 2011; Ozlü et al., 2005; Severson et al., 2000). This analysis suggested that the N-tail sites are likely targeted by Aurora A rather than Aurora B in vivo (Fig. S3A-E). Aurora A has also been implicated in control of kinetochore-microtubule attachments and Ndc80 tail phosphorylation in other species (Chmátal et al., 2015; Ye et al., 2015).

Although spindle elongation was inhibited by NDC-80^{4A} for a substantial portion of the NEBD-anaphase onset interval, ~1 min prior to anaphase onset the spindle poles began to separate at a rate comparable to controls (Fig. 2C). Previous work has shown that during this pre-anaphase interval, the cell polarity-regulated G-protein (Ga) pathway activates cortical dynein to generate forces pulling on the asters that position the metaphase spindle and promote its elongation (Fig. 2D;(Gönczy and Rose, 2005; Labbé et al., 2004; Nguyen-Ngoc et al., 2007). Thus, we wondered if Ga-dependent pulling forces were responsible for the late spindle elongation observed in NDC-80^{4A}. To test this, we inhibited the Ga pathway by depletion of its essential regulators GPR-1 and GPR-2 (Fig. 2D; (Colombo et al., 2003)) which does not alter the NEBD-anaphase interval, relative to control embryos, or result in

chromosome segregation defects (Fig. S2E). In *gpr-1/2(RNAi)* embryos, the spindle elongation rate was initially similar to that in controls; however, the increase in the spindle elongation rate that occurs ~1 min prior to anaphase onset in controls, failed to occur, indicating that it is mediated by Ga pathway activation (Fig. 2D; S2E; *red shaded region*). In the NDC-80^{4A} mutant, inhibition of the Ga-dependent force, by depletion of GPR-1/2, abolished the spindle pole separation observed prior to anaphase onset (Fig. 2E; *compare to* Fig. 2D). Thus, the non-phosphorylatable NDC-80 N-tail restrains spindle elongation prior to Ga activation, during the interval when the chromosomes are aligning (50–130s after NEBD, Fig. 1A,B) but does not inhibit elongation once the Ga-dependent forces, by pulling on the asters, ramp up the tension at kinetochore-microtubule attachment sites in the spindle.

Next, to test if the suppression of spindle elongation prior to Ga activation by the nonphosphorylatable tail requires the ability of the CH domain to dock onto microtubules, we combined the non-phosphorylatable N-tail mutant with the CH^{Toe} mutant in the same *ndc-80* transgene (Fig. 2F). Comparison of CH^{Toe} and 4A-CH^{Toe} forms of NDC-80 revealed identical phenotypes (Fig. 2F,S2F), indicating that the inhibition of spindle elongation observed in NDC-80^{4A} is dependent on the docking of the CH Toe to the microtubule lattice.

The NDC-80^{4A} mutant on its own exhibited normal chromosome alignment and segregation and supported embryo viability (Fig. S2F; Cheerambathur et al., 2013). To test if the NDC-80^{4A} mutant exhibited chromosome segregation phenotypes in a sensitized background, we compromised chromosome alignment by reducing the activity of the kinesin-4 family chromokinesin KLP-19. In one-cell *C. elegans* embryos KLP-19 provides an important non-kinetochore force for positioning and orienting chromosomes on the spindle (Powers et al., 2004). Weak inhibition of KLP-19 (generated by combining *ndc-80* and *klp-19* dsRNAs in 24:1 ratio) resulted in modest (~30%) embryonic lethality in the presence of NDC-80^{WT} that was significantly elevated (to ~90%) in the presence of NDC-80^{4A} (Fig. 2G). Thus, NDC-80 N-tail phosphorylation promotes embryonic viability when the activity of the chromokinesin KLP-19 is reduced.

Suppression of Spindle Elongation by the Non-Phosphorylatable NDC-80 Tail Mutant Requires the SKA Complex

The results above suggest that the presence of the unphosphorylated NDC-80 N-tail restrains spindle elongation, likely by altering the properties of kinetochore-microtubule attachments. One possibility for how the unphosphorylated N-tail could accomplish this is by promoting the recruitment of an effector that suppresses attachment dynamics. A candidate for such an effector is the Ska complex, a kinetochore-localized microtubule-binding complex that is conserved across metazoans and is suggested to be a functional counterpart of the Dam1 complex in fungi (Gaitanos et al., 2009; Welburn et al., 2009). Inhibition of the Ska complex in cultured mammalian cells compromises kinetochore-microtubule interactions (Gaitanos et al., 2009; Raaijmakers et al., 2009; Welburn et al., 2009). Localization of the Ska complex to kinetochores is dependent on the Ndc80 complex and regulated by Aurora kinase activity (Chan et al., 2012). Although the two complexes do not interact significantly in solution, the Ska complex enhances the microtubule binding affinity and end-tracking ability of the Ndc80 complex *in vitro* (Schmidt et al., 2012). These observations have led to the model that

the Ndc80 complex recruits the Ska complex to kinetochores and their cooperation generates robust kinetochore-microtubule attachments.

In *C. elegans*, the SKA complex is heterotrimeric, consisting of two SKA-1 subunits and one SKA-3 subunit (Fig. 3A; Schmidt et al., 2012). To determine if the SKA complex is required for the non-phosphorylatable NDC-80 N-tail to prevent spindle elongation, we compared the effect of SKA-3 depletion on spindle pole elongation kinetics in NDC-80^{WT} versus NDC-80^{4A} (Fig. 3A). In the presence of NDC-80^{WT}, SKA-3 depleted embryos exhibited spindle pole separation dynamics similar to those in controls. In contrast, SKA complex depletion in the non-phosphorylatable NDC-80^{4A} mutant eliminated the inhibition of spindle pole separation (Fig. 3A). Thus, the ability of the non-phosphorylatable NDC-80 N-tail to inhibit spindle elongation is dependent on the SKA complex.

Dephosphorylation of the NDC-80 N-tail and Activation of the Ga Pathway Both Promote SKA Complex Recruitment

Given that the SKA complex is required for the dephosphorylated N-tail to restrain spindle elongation, we next assessed if there was a difference in SKA complex localization between NDC-80^{WT} and NDC-80^{4A} embryos. We first analyzed a strain co-expressing SKA-1 tagged at its endogenous locus with GFP and an mCherry fusion with the NDC-80 complex subunit Nuf2^{HIM-10} (Fig. 3B). The NDC-80 complex localized to the diffuse kinetochores of *C. elegans* throughout mitosis, as expected from prior work (Desai et al., 2003). By contrast, robust SKA-1 localization was only observed in a brief time window beginning ~60s prior to anaphase onset, after chromosomes were aligned at the metaphase plate (Fig. 3B; Movie S1).

SKA-1 localization to kinetochores required NDC-80 (Fig. 3C), as expected from prior analysis in mammalian cells (Gaitanos et al., 2009; Raaijmakers et al., 2009; Welburn et al., 2009). Notably, SKA-1 recruitment was abolished by mutation of the Toe residues of the NDC-80 CH domain; by contrast, SKA-1 recruitment was not significantly affected by the CH^{Ankle} mutant (Fig. 3C). Thus, SKA-1 localization is dependent on the NDC-80 CH domain Toe docking to the microtubule lattice.

The abrupt recruitment of the SKA complex (Fig. 3B) correlated with the activation of the Ga-dependent pulling forces that likely place bioriented kinetochore-microtubule interfaces under higher tension (Fig. 2D). Therefore, we next tested whether SKA-1 recruitment was affected by *gpr-1/2* depletion. Localization of SKA-1 was significantly diminished following inhibition of Ga-dependent forces (Fig. 3D), suggesting that force applied onto NDC-80 bound to microtubules via its CH domain promotes a conformation that is recognized by the SKA complex.

In the NDC-80^{4A} mutant, spindle elongation is restrained in a SKA complex-dependent manner (Fig. 3A) suggesting that tail dephosphorylation influences the recruitment of SKA. Consistent with this idea, SKA-1 localization was significantly elevated in the presence of NDC-80^{4A} compared to NDC-80^{WT} (Fig. 3E,F). However the presence of a non-phosphorylatable tail is not sufficient to recruit SKA-1, as SKA-1 localization was not observed in a combined non-phosphorylatable N-tail-CH^{Toe} mutant (Fig. 3C). These results

suggest that the non-phosphorylatable N-tail promotes adoption of a microtubule-bound conformation of NDC-80 recognized by the SKA complex. Additional experiments suggested that tail dephosphorylation modulates but is not essential for the rapid increase in SKA-1 recruitment that occurs concomitant with Ga activation. There is a decrease in the maximum intensity of SKA-1 recruited by NDC-80^{Tail} relative to NDC-80^{WT} (Fig. 3E,F & S4B), and a phosphomimetic 4D tail mutation, which behaves analogously to NDC-80^{WT} and NDC-80^{Tail} in the spindle pole separation assay (Fig. S4A), also reduced SKA recruitment (Fig. 3E,F & S4B).

Taken together, the above data suggest that in wild-type embryos N-tail dephosphorylation and increased pulling forces resulting from Ga activation both contribute to SKA-1 recruitment after chromosomes have aligned at the metaphase plate.

The SKA Complex Dampens Chromosome Movement on the Spindle

The functional effect of the non-phosphorylatable N-tail—suppression of spindle elongation —is mediated by the SKA complex, suggesting that recruitment of the SKA complex modulates the properties of the kinetochore-microtubule interface. To address the function of the SKA complex, we used CRISPR/Cas9 to delete *ska-1* (Fig. S5A) and also used RNAi to deplete SKA-3. We optimized imaging conditions to monitor chromosome dynamics at high temporal resolution (image stack acquisition every 2 seconds) and developed a method to quantify chromosome distribution over time.

In control embryos, chromosomes rapidly move (by ~100s after NEBD) to the vicinity of the metaphase plate but continue to exhibit dynamic movement. In the ~40 seconds prior to anaphase onset this motion is dampened (Fig. 4A). The tightly aligned, motion-dampened metaphase plate is then separated into two relatively evenly shaped chromosome masses in anaphase (Fig. 4A). Imaging of chromosome dynamics revealed that the dampening of movement that tightens the metaphase plate is significantly perturbed by both mutant (*ska-1*) and RNAi (*ska-3*) inhibition of the SKA complex (Fig. 4A). In both conditions, chromosomes continued to exhibit dynamic movements after reaching the spindle equator, underwent large excursions from the metaphase plate and were observed to separate as irregular-shaped chromosome masses in anaphase (Fig. 4A; Movie S2). In 8 of 15 embryos imaged at high time resolution, individual chromosomes were observed to abruptly detach from the main chromosome mass and realign (*green arrowhead* Fig. 4A; Movie S2), a behavior that was not observed in 13 control embryos imaged under equivalent conditions.

To quantify chromosome dynamics, we used a bounding box to encase the chromosomes in a maximal intensity projection of a background-subtracted image and plotted the width of the bounding box along the spindle axis over time. Both individual (Fig. 4B, S5B) and averaged (Fig. 4C) traces are shown. In both controls and SKA complex inhibitions, chromosomes became confined to a smaller spindle region over time. However, individual traces indicated broader chromosome distribution along the spindle axis and abrupt chromosome motions in the absence of the SKA complex (Fig. 4B), and the averaged traces indicated a significant difference in chromosome confinement on the spindle between the control and SKA complex inhibitions (Fig. 4C). We did not observe any effect of SKA complex inhibitions on the NEBD-anaphase onset interval (Fig. S5C), indicating that the

Page 10

role proposed for this complex in promoting anaphase onset in mammalian tissue culture cells (Sivakumar et al., 2016) is not conserved in *C. elegans. ska-1* worms are homozygous viable at 20°C but exhibit mi ld embryonic lethality (~3%) that was elevated 5-fold at 15°C, a temperature that des tabilizes microtubules (Fig 4D). In addition, *ska-1* worms were sensitive to reduction in activity of the chromokinesin KLP-19 (Fig. 4D), as also observed for NDC-80^{4A} (Fig 4D). Thus, while SKA complex-mediated dampening of chromosome motion after biorientation is not as essential for viability as NDC-80 complex-mediated attachments, it contributes to viability.

We conclude that the SKA complex is required for dampening chromosome motion after alignment to tightly confine bi-oriented chromosomes on the spindle and coordinate their segregation during anaphase. Based on the effect of NDC-80^{4A}, which requires the SKA complex, we suggest that this dampening is due to an alteration of the dynamic nature of NDC-80-mediated attachments. Moreover, this dampening after biorientation contributes to organismal viability, likely by ensuring accurate chromosome segregation.

The Microtubule-Binding Domain of SKA-1 Contributes to Its Localization and Function

The microtubule-binding activity of the Ska complex has been mapped to a conserved Cterminal domain of Ska1 that adopts a winged-helix fold (Abad et al., 2014; Schmidt et al., 2012). In vitro analysis of the C. elegans SKA complex has shown that deletion of this domain inhibits microtubule binding but does not disrupt complex formation (Schmidt et al., 2012); in addition the domain and the intact complex both bind microtubules with similar weak affinity (K_d \sim 6–10 μ M). To determine if this conserved domain contributes to SKA complex function in vivo, we employed a targeted transgene-based replacement system to generate strains expressing SKA-1::GFP (WT) and a comparable fusion lacking the microtubule-binding domain (MTBD) (Fig. 5A; Fig. S5D). In the presence of endogenous SKA-1, both the wildtype and MTBD SKA-1 localized to metaphase kinetochores (Fig. 5A). This observation suggests that MTBD SKA-1 is expressed and associates with SKA-3, which is in agreement with *in vitro* analysis (Schmidt et al., 2012). When the transgeneencoded SKA-1 variants were crossed into ska-1, wild-type SKA-1 localized robustly but localization of the MTBD mutant was significantly compromised (~30% relative to wildtype SKA-1 just prior to anaphase onset; Fig. 5B). This result suggests that the localization of MTBD SKA-1 observed in the presence of endogenous SKA-1 is likely due to its association with wildtype SKA-1 in the trimeric SKA complex. Consistent with the reduced localization of MTBD SKA-1, imaging of chromosome dynamics highlighted wider chromosome distribution with excursions away from the plate, phenotypes similar to ska-1 but less severe in magnitude (Fig. 5C; Fig. S5E). Thus, the microtubule-binding domain of SKA-1 contributes to NDC-80-dependent SKA complex localization on bioriented chromosomes and to the dampening of chromosome motion prior to anaphase onset.

DISCUSSION

Mechanisms that ensure chromosome biorientation on the spindle prior to anaphase onset, such as the spindle checkpoint or error correction pathways, have been the focus of many efforts to understand the high fidelity of chromosome segregation (Godek et al., 2014). Here,

we describe a mechanism that acts after chromosome biorientation and contributes to organismal viability, likely by ensuring accurate segregation. Specifically, our results suggest that both N-tail dephosphorylation and spindle-external force-induced tension promote a specific conformation of microtubule-bound NDC-80 that recruits the SKA complex to dampen motion of bioriented chromosomes and spatially constrain them prior to their segregation in anaphase (Fig. 5D). This dampening is functionally important as the absence of the SKA complex compromises viability.

It has been appreciated for over a decade that kinetochore-microtubule attachments are regulated by phosphorylation of the basic N-terminal tail of the Ndc80 subunit of the Ndc80 complex (Deluca et al., 2006). The predominant view on the molecular mechanism underlying this regulation, based on *in vitro* studies, is that phosphorylation counteracts electrostatic interactions of the basic Ndc80 N-tail with the acidic microtubule lattice (Alushin et al., 2012; Cheeseman et al., 2006; Ciferri et al., 2008; Wei et al., 2007; Zaytsev et al., 2015). In contrast to this view, we show that the NDC-80 N-tail does not make a detectable contribution to microtubule affinity at kinetochores in the one-cell C. elegans embryo. Nevertheless, analogous to what has been reported in vertebrate cells (DeLuca et al., 2011), a non-phosphorylatable mutant of the NDC-80 N-tail prematurely stabilizes kinetochore-microtubule attachments. As this effect of non-phosphorylatable NDC-80 is entirely dependent on the SKA complex, we suggest that in metazoan species where both complexes are present, Ndc80 tail phosphorylation modulates kinetochore-microtubule attachment dynamics either exclusively or in part via the Ska complex. In vertebrate cells, Ndc80 tail deletion or alteration of tail phosphorylation is known to perturb kinetochoremicrotubule interactions, and it should be feasible to test if the Ska complex contributes to these effects. Notably, work described in the accompanying manuscript (Janczyck et. al., 2017) indicates that a specific alteration of the N-tail of human Ndc80 perturbs Ska complex localization to kinetochores, revealing a link between the N-tail and the Ska complex in human cells.

In the *C. elegans* embryo, the action of the SKA complex on NDC-80-mediated attachments dampens chromosome motion prior to anaphase onset. Dampening of chromosome motion could be achieved by stabilizing a paused state of dynamic microtubules at the kinetochore or by promoting slow continual polymerization at bi-oriented sister kinetochores; the latter is likely the case in the *C. elegans* embryo, as spindle elongation continues to occur at a slow rate until anaphase onset. The abrupt detachment of individual chromosomes observed in the absence of the SKA complex also suggests that SKA helps strengthen NDC-80-mediated attachments. This idea should be testable in reconstitution experiments employing optical trapping, analogous to those conducted for budding yeast kinetochores (Tien et al., 2010).

Structural studies and *in vitro* microtubule-binding analysis have shown that the Ska1 subunit of the Ska complex directly binds to microtubules and synergizes with the Ndc80 complex (Abad et al., 2014; Schmidt et al., 2012). Our results show that SKA complex enrichment at the kinetochore-microtubule interface in the *C. elegans* embryo depends on the Toe residues in the NDC-80 CH domain and, to a significant extent, on the microtubule-binding domain of SKA-1. We propose that the SKA complex recognizes a specific microtubule-bound conformation of the NDC-80 complex and concomitant microtubule

binding by the conserved winged-helix domain of SKA-1 provides additional affinity to enable SKA complex recruitment and alteration of attachment dynamics at kinetochores. We note that the microtubule-binding domain deletion does not eliminate SKA complex localization and does not fully phenocopy *ska-1*. Thus, we suspect that the SKA complex lacking this domain is able to recognize and modulate NDC-80 complex-mediated attachments to a degree, an idea that will need to be explored in future work. The distinct microtubule lattice features recognized by the Ndc80 and Ska complexes (Abad et al., 2014; Alushin et al., 2012; Schmidt et al., 2012) are also likely to be important for their integrated action at the kinetochore-microtubule interface.

In human cells, deletion of the microtubule-binding domain of Ska1 does not prevent kinetochore localization, although it appears reduced (Redli et al., 2016; Schmidt et al., 2012). The human Ska complex has an additional subunit (Ska2), is hexameric (*C. elegans* SKA complex is trimeric), and the Ska3 subunits are significantly divergent. Thus, it is possible that the human Ska complex has intrinsic affinity for the Ndc80 complex in the absence of microtubule binding. While robust biochemical association of purified recombinant human Ska and Ndc80 complexes has not been reported to date, this possibility is supported by localization of Ska to kinetochores lacking microtubule attachments following Aurora B kinase inhibition in human cells (Chan et al., 2012). The microtubule-binding domain of human Ska1 has also been suggested to bind protein phosphatase 1 (Sivakumar et al., 2016) and to bind and activate Aurora B (Redli et al., 2016). We have not observed reduction in protein phosphatase 1 (GSP-2) localization at kinetochores or phenotypes associated with reduced Aurora B activity in *ska-1 (not shown*), suggesting that these proposed interactions are not conserved in *C. elegans*.

The abrupt timing of SKA complex recruitment observed in the *C. elegans* embryo is striking. We suggest that this timing is dictated primarily by the generation of a microtubulebound NDC-80 conformation that is promoted by tension at attachment sites (Fig. 5D). In this view, the dephosphorylated N-tail or other mechanisms, including spindle-external force generation pathways such as the Ga/LGN pathway or poleward flux of kinetochore microtubules observed in other species (Matos et al., 2009), act in a unified manner by promoting generation of a microtubule-bound Ndc80 conformation that recruits Ska to trigger the synergy between these two microtubule-binding complexes. We note that a role for the N-terminal tail in Ndc80 conformation control is supported by analysis employing a conformation-sensitive sensor engineered into Ndc80 in budding yeast, which lacks Ska but harbors the functionally analogous microtubule-binding Dam complex (Suzuki et al., 2016). One important unanswered question with respect to timing of SKA recruitment is the mechanism of Ndc80 tail dephosphorylation. While specific kinetochore-localized phosphatases are suggested as being involved in dephosphorylation of Aurora substrates at the kinetochore (Liu et al., 2010; Suijkerbuijk et al., 2012), the precise mechanisms controlling Ndc80 tail dephosphorylation and its relationship to attachment and tension at the kinetochore-microtubule interface remain to be clarified.

RNAi analysis of the Ska complex in human tissue culture cells has suggested a role in kinetochore-microtubule attachments as well as in promoting the metaphase-anaphase transition (Gaitanos et al., 2009; Schmidt et al., 2012; Sivakumar et al., 2016). Our analysis

of the *C. elegans* SKA complex using both a null *ska-1* mutant and RNAi-mediated depletions of both subunits affirmed a function for this complex in the regulation of attachment dynamics after biorientation but did not reveal a role in controlling the timing of the metaphase-anaphase transition. In contrast to mutants that disrupt docking of the NDC-80 CH domain docking onto the microtubule lattice, loss of SKA complex function did not lead to penetrant lethality, indicating that SKA activity is not essential for biorientation. Nonetheless, loss of the SKA complex compromised organismal viability, especially when microtubule stability was reduced or when a different pathway that aids chromosome segregation was weakened. Thus, we propose that SKA-mediated alteration of dynamics at the kinetochore-microtubule interface after biorientation functions as a safety mechanism to ensure that daughter cells inherit the correct genome complement. Misregulation of such a mechanism may contribute to premature kinetochore microtubule stability-linked chromosome segregation errors in cancer cells, a possibility that will be interesting to address in future work.

STAR Methods

Contact for Reagent and Resource Sharing

For further information, and requests regarding reagents should be directed to and will be fulfilled by the Lead Contact, Arshad Desai (abdesai@ucsd.edu).

Experimental Model and Subject Details

The genotypes of the *C. elegans* strains used in this study are described in **Reagents and Resources section** Strains were maintained at 20°C, except for temperature-sensitive strains, which were maintained at 16° C.

Method Details

C. elegans transgenic strain construction—Strains with integrated single-copy *ndc-80* transgenes, which are RNAi-resistant and *ska-1::gfp* transgenes were constructed using the MosSCI method (Frøkjaer-Jensen et al., 2008). The *ndc-80* transgene was described previously (Cheerambathur et al., 2013). In brief, the *ndc-80* genomic locus including introns, 2000 bp upstream of the start codon and 1000 bp downstream of the stop codon are present in the transgene. A 776 bp region spanning exons 3–5 was re-encoded in the transgene as described (Cheerambathur et al., 2013) to make the transgene-encoded product resistant to a dsRNA targeting this region of endogenous *ndc-80*. The engineering of the *ska-1::gfp* transgene is described in Fig. S5D. Single-copy insertions were confirmed by PCR analysis flanking the insertion site.

Endogenous GFP tagging and deletion of the *ska-1* locus were done using CRISPR/Cas9 (Dickinson et al., 2013). For *ska-1::gfp*, the repair template included homology arms flanking the linker sequence (GGRAGSG) and GFP (1667 bp right homology arm comprised of the *ska-1* coding sequence and 5' UTR and 2050 bp left homology arm comprised of the 3' UTR and the downstream gene locus Y106G6H.13). A *Cb-unc-119* selection marker flanked by *loxp* sequences was inserted between the GFP and the 3' homology arm of the repair template. For *ska-1* deletion, the repair template was generated

by inserting a Cb-unc-119 selection marker flanked by loxp sequences between the appropriate homology arms (2000 bp right homology arm comprised of the 5' UTR of ska-1 coding sequence and 2050 bp left homology arm comprised of the 3' UTR and the downstream gene locus Y106G6H.13). To modify the ska-1 locus using Cas9-triggered homologous recombination, the repair template ($50ng/\mu L$) was co-injected with a plasmid (40ng/ μ L), modified from pDD162, containing Cas9 and respective guide RNA sequences (5'-AAATTTTCATCGGTCTCTT-3' for ska-1::gfp and 5'-AATGGAATCGTTTATTGAT-3' for ska-1) and three plasmids encoding fluorescent markers for negative selection [pCFJ90 (Pmyo-2::mCherry, 2.5 ng/µL), pCFJ104 (Pmyo-3::mCherry, 5 ng/µL) and pGH8 (Prab-3::mCherry, 10 ng/µL)] into the strain HT1593. Recombinant strains were identified by selecting moving worms without fluorescent markers and were confirmed by PCR spanning both homology regions. The *Cb-unc-119* selection marker was later deleted using Cre recombinase-mediated excision. An injection mix containing pDD104 (Peft-3::Cre, 50 ng/µL) and coinjection markers [pCFJ90 (Pmyo-2::mCherry, 2.5 ng/µL) and pCFJ104 (Pmyo-3::mCherry, 5 ng/µL)] was injected into the gonads of unc-119(+) young adult *ska-1::gfp* or *ska-1* animals. Cre-excised animals were identified by selecting for the Unc phenotype and were confirmed by PCR. The unc-119 mutation was subsequently removed by mating with N2 males.

RNA-mediated Interference—Double-stranded RNA synthesis was performed by first using PCR to generate DNA templates using the primers and templates listed in Table S1 in the supplement. PCR reactions were cleaned using the QIAquick PCR Purification Kit (Qiagen). Individual T3 and T7 transcription reactions using MEGAscript® T7/T3 Transcription Kit (ThermoFisher Scientific) on each template were used to generate single-stranded RNAs. The two reactions were purified using the MEGAclearTM Kit (ThermoFisher Scientific), mixed and annealed at 37°C for 30 minutes to generate double- stranded RNAs (Oegema et al., 2001). L4 hermaphrodites were injected with dsRNA and incubated at 20°C for 36–46h before dissection and imaging of embryos. For *ndc-80&crod-1(RNAi)* and *ndc-80&gpr-1/2(RNAi)*, 1 mg/mL individual dsRNAs were mixed at a 1:1 ratio, except for the analysis of embryonic lethality following *ndc-80&klp-19* or control *hyls-1&klp-19* depletions, where a ratio of 24:1 was used to mildly reduce KLP-19 activity.

Embryonic lethality assays—For assaying embryonic lethality of *ska-1* strains at different temperatures, L4 worms were singled onto plates and allowed to lay eggs at 15° and 20°C. After 24 hours the mother was removed and the number of eggs and hatched larvae were scored 24 hours later. N2 worms were used as controls.

For embryonic lethality assays involving partial depletions of *klp-19*, L4 worms were injected with dsRNA mixes containing *klp-19 and ndc-80* dsRNAs or *klp-19 and hyls-1* dsRNAs and incubated at 20°C. The adult worms were singled onto plates after 36 hours and viability of embryos during the 36–48 hour interval was determined.

Live Imaging, Fluorescence Intensity Quantification and Image Analysis—For imaging one-cell embryos, hermaphrodites were dissected in M9 buffer and transferred to a freshly prepared 2% agarose pad and covered with a 22×22 mm coverslip. In experiments that involved imaging of the temperature-sensitive *air-2(or207ts)* strain, the dissection and

imaging were performed as described in (Lewellyn et al., 2011). In brief, for pre-meiotic upshift, worms were placed in a pre-heated microscope room (26.5°C) for 30–60 min prior to dissection and imaging. For post-meiotic upshift, worms were kept at ~16°C in the pre-heated room, dissected and imaged at the restrictive temperature.

For pole-tracking analysis of one cell embryos expressing GFP::Histone H2B and GFP:: γ -tubulin, 5×2 µm z-stacks were collected with 2×2 binning and a 60X 1.3 NA U-planApo objective (Olympus) at 10 second intervals and 150 ms exposure at 19°C on a DeltaVision microscope (Applied Precision) equipped with a CoolSnap camera (Roper Scientific). Maximum intensity projections of z-stacks were made using Image J (Fiji) and pole tracking profiles were generated by measuring spindle length (the distance between the two centrosomes in the projections) over time using Metamorph (Molecular Devices). Spindle pole separation was plotted relative to Nuclear Envelope Breakdown (NEBD). NEBD was scored as the frame where the diffuse intranuclear histone signal equilibrated with the cytoplasm and chromosomes exhibited abrupt rapid movement indicating interaction with microtubules. Anaphase onset was scored as the frame where two distinct masses of sister chromosomes were first visible. Pre-anaphase and anaphase spindle elongation rates were calculated using linear regression analysis on the spindle pole elongation profiles within time intervals 50–200s and 200–300s, respectively.

Images of one-cell embryos expressing GFP::H2B or mCherry::H2B for analysis of chromosome dynamics, SKA1::GFP, and GFP::β-tubulin;mCherry::H2B were collected using a confocal imaging system (Revolution XD Confocal System; Andor Technology) with a spinning disk confocal scanner unit (CSU-10; Yokogawa Corporation of America) mounted on an inverted microscope (TE2000-E; Nikon) equipped with solid-state 100-mW lasers, 60X, 1.4 NA Plan Apochromat (for imaging embryos expressing SKA1::GFP or GFP::β-tubulin;mCherry::H2B) and 100X, 1.4 NA Plan Apochromat lens (for chromosome dynamics analysis and SKA1::GFP imaging in Fig. 3B), and a back-illuminated EMCCD camera (binning 1×1; iXon DV887; Andor Technology).

To quantify SKA-1::GFP fluorescence, $5\times2 \mu m z$ -stacks were acquired at 20 second intervals and 500 ms exposure. Z-stacks were projected and fluorescence intensity was quantified using Image J (Fiji). A rectangular box was drawn around the SKA-1::GFP fluorescence and the integrated intensity in the box was recorded. The box was expanded by 5 pixels on each side, and the integrated intensity was measured. The signal and area difference between the expanded box and the original box were used to calculate the average background signal per pixel. The integrated chromosomal SKA-1::GFP intensity in the original box was then calculated by subtracting the background signal.

For minimum bounding box analysis used to quantify chromosome distribution, $5 \times 1.5 \ \mu m z$ stacks were acquired every 2 s in the case of GFP::H2B or 5 s in the case of mCherry::H2B. For each z-stack, a maximum intensity projection image was generated using Image J (Fiji), and rotated to position the spindle poles horizontally. Next, the intensity of all projection images of the timelapse z-series was normalized, converted to 8-bit, and both spindle poles cropped out. Subsequently, the background signal was subtracted, and to each processed

projection image a minimum bounding box was fitted (only including pixels with value > 0, i.e. histone signal from the chromosomes) and the width of this box was recorded.

For generating kymographs, a maximum intensity projection of each z-stack ($5\times2 \mu m$), was made using Image J (Fiji), and rotated to position the spindle poles horizontally. Next, a 30-pixel wide linescan was drawn through the spindle region and kymograph was generated using the kymograph plugin in Image J (Fiji).

Quantification and Statistical Analysis

Details of the methods employed to extract and quantify various parameters in microscopy datasets are described in the image analysis section. The statistical tests used to determine significance are described in the figure legends.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

- The Ndc80 tail is dispensable for microtubule affinity at *C. elegans* kinetochores
- Preventing Ndc80 tail phosphorylation nonetheless stabilizes microtubule attachments
- The Ska complex is required for stabilization by the dephosphorylated Ndc80 tail
- Ska complex recruitment dampens chromosome motion to promote accurate segregation

Cheerambathur et al.



Figure 1. Quantitative analysis of spindle elongation discriminates NDC-80 mutations *in vivo* that alter microtubule-binding affinity

(A) Stills from movies of control and *ndc-80(RNAi)* one-cell embryos expressing GFP::H2b and GFP:: γ -tubulin, to label chromosomes (*green arrow*) and spindle poles (*green arrowheads*), respectively. Time is in seconds after NEBD. Yellow arrowheads highlight premature spindle pole separation in *ndc-80(RNAi)*. Scale bar, 5 µm.

(**B**) Single copy transgene insertion-based NDC-80 replacement system. Spindle pole tracking analysis below shows rescue of *ndc-80(RNAi)* by transgene-encoded RNAi-resistant wild-type NDC-80. Error bars are the 95% confidence interval.

(C) Structure of the human Ndc80-Nuf2 head (*pink/blue*) bound to a tubulin dimer (*gray, labeled* α/β) in the microtubule lattice (PDB 3IZ0). The Toe and Ankle residues of the Ndc80 CH domain are colored in green and yellow, respectively. A rotated *en face* view of the microtubule-binding surface of the CH domain is on the right. Residue numbers are for human Ndc80 (*black*) and *C. elegans* NDC-80 (*green/yellow*).

(**D**) Spindle pole separation analysis for the indicated CH domain mutants following endogenous NDC-80 depletion. The 50s-200s interval after NEBD is plotted. Error bars are the 95% confidence interval. WT and No Transgene data are the same as in *Fig. 1B*.

(E) Chromosome segregation phenotypes for the indicated CH domain mutants highlighted by image stills from time-lapse movies 20 sec prior to and after anaphase onset; percentage of one-cell embryo anaphases with detectable lagging chromatin are indicated on the right. Scale bar, 5 μ m. See also Fig. S1.





(C) Spindle pole separation analysis of the indicated conditions. WT and Tail data are the same as in Fig. 1B and *Fig. 2A*, respectively. Error bars are the 95% confidence interval.
(D) Schematic of the Ga pathway that generates an asymmetric force for spindle displacement starting ~1 min prior to anaphase onset (*orange & red* arrows indicate forces; *black dashed arrow* indicates direction of motor; adapted from (Gönczy and Rose, 2005). Graph shows effect of Ga pathway inhibition by depletion of GPR-1/2 in the spindle pole separation assay. Red arrows mark anaphase onset, which is not affected by GPR-1/2 depletion. Error bars are the 95% confidence interval.

(E) & (F) Spindle pole separation analysis of the indicated conditions. Error bars are the 95% confidence interval. 4A and CH^{Toe} alone data is the same as in *Fig. 2C* and Fig. 1D, respectively.

(G) Embryonic lethality in WT and 4A NDC-80 following weak depletion of KLP-19. *n*=number of worms. See also Figs. S2 & S3.

Cheerambathur et al.



Figure 3. The non-phosphorylatable NDC-80 N-tail mutant suppresses spindle elongation via the SKA complex

(A) Schematics of *C. elegans* SKA complex subunits SKA-1 and SKA-3. Spindle pole separation analysis for the indicated conditions. WT and 4A alone data are same as in Fig. 1B and Fig. 2C, respectively. Error bars are the 95% confidence interval.

(B) Stills from a time-lapse movie of a one-cell embryo co-expressing mCherry-tagged NDC-80 complex and GFP-tagged SKA complex. Scale bar, 5 μ m.

(C) SKA-1::GFP localization in the indicated conditions. 10–12 embryos were imaged per condition. Scale bar, 2.5 μ m.

(**D**) (*top*) Stills from time-lapse movies of SKA-1::GFP localization in control and *gpr-1/2(RNAi)*. (*bottom*) Quantification of SKA-1::GFP chromosomal intensity relative to anaphase onset. Error bars are the 95% confidence interval. Measured fluorescence intensities were normalized to the averaged maximal intensity value in control.

(E) Stills from time-lapse movies of SKA-1::GFP chromosomal localization in the indicated conditions. Scale bar, $2.5 \mu m$.

(**F**) Quantification of SKA-1::GFP chromosomal intensity relative to anaphase onset for the indicated conditions, performed as in *Fig. 3D* above. See also Fig. S4.





(A) High temporal resolution analysis (6-plane z-stack at 1.5 μ m spacing acquired every 2s) of chromosome dynamics. Low time resolution (30s interval) stills from early prometaphase to just after anaphase onset are shown for control on top and higher time-resolution stills for the indicated conditions during the selected interval are shown below. Scale bar, 2.5 μ m. (B) Method employed to analyze chromosome distribution along the spindle axis using a bounding box. Individual control and *ska-1* traces are shown; stills below illustrate chromosome behavior. Scale bar, 2.5 μ m.

(C) Average plot of chromosome distribution along the spindle axis for the indicated conditions. Error bars are the 95% confidence interval.

(D) *(left)* Comparison of embryonic lethality in WT and *ska-1* at 20°C and 15°C. *(right)* Plot of embryonic lethality in control and *ska-1* following partial depletion of KLP-19. n=number of worms. See also Fig. S5.

Cheerambathur et al.



Dampened Motion

Figure 5. The microtubule-binding domain of SKA-1 contributes to SKA complex localization and function $% \left({{{\rm{SKA}}} \right) = 0.05} \right)$

(A) Schematic of transgene-expressed GFP fusions of SKA-1; the microtubule-binding domain deletion (MTBD) is identical to the one characterized *in vitro* (Schmidt et al., 2012). Images below show metaphase stage embryos in strains where endogenous SKA-1 is present. Scale bar, 5 μ m.

(**B**) Stills from time-lapse movies for the indicated conditions. The graph on the right plots background-subtracted chromosomal GFP fluorescence for the frame (–20s) prior to

anaphase onset. Measurements were normalized to the mean value of wild-type SKA-1::GFP. Error bars are the 95% confidence interval. Scale bar, 5 $\mu m.$

(C) Examples of bounding box width analysis of mCherry::H2b timelapse sequences,

performed as in Fig. 4B; stills below illustrate chromosome behavior for the two analyzed conditions. Scale bar, $2.5 \ \mu m$.

(**D**) Model for dampening of chromosome motion after biorientation. Boxes with dashed lines contain speculations discussed in the text. See also Fig. S5.