Aurora B-dependent Ndc80 degradation regulates kinetochore composition in meiosis

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The kinetochore complex is a conserved machinery that connects chromosomes to spindle microtubules. During meiosis, the kinetochore is restructured to accommodate a specialized chromosome segregation pattern. In budding yeast, meiotic kinetochore remodeling is mediated by the temporal changes in the abundance of a single subunit called Ndc80. We previously described the regulatory events that control the timely synthesis of Ndc80. Here, we report that Ndc80 turnover is also tightly regulated in meiosis: Ndc80 degradation is active in meiotic prophase, but not in metaphase I. Ndc80 degradation depends on the ubiquitin ligase APC^Ama1 and is mediated by the proteasome. Importantly, Aurora B-dependent Ndc80 phosphorylation, a mark that has been previously implicated in correcting erroneous microtubule–kinetochore attachments, is essential for Ndc80 degradation in a microtubule-independent manner. The N terminus of Ndc80, including a 27-residue sequence and Aurora B phosphorylation sites, is both necessary and sufficient for kinetochore protein degradation. Finally, defects in Ndc80 turnover predispose meiotic cells to chromosome mis-segregation. Our study elucidates the mechanism by which meiotic cells modulate their kinetochore composition through regulated Ndc80 degradation, and demonstrates that Aurora B-dependent regulation of kinetochores extends beyond altering microtubule attachments.

[Keywords: meiosis; kinetochore; Aurora B; Ndc80; chromosome; proteolysis; APC]

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Reproduction is a fundamental feature of life and depends on the accurate segregation of chromosomes from one generation to the next. In eukaryotes, a conserved protein complex known as the kinetochore mediates chromosome segregation. Research over the past three decades has identified at least 40 different proteins that constitute the core of this essential machinery [reviewed extensively in Biggins 2013]. While the function of individual kinetochore components has been well established, much less is understood about how the levels of specific subunits are regulated under varying cellular states and how these changes affect kinetochore function.

The kinetochore is composed of two distinct parts: inner and outer kinetochore. The inner kinetochore subunits associate with the chromosome at the centromere, while the outer kinetochore components interact with spindle microtubules. It has been shown that changes in either part can have a profound impact on kinetochore activity and genome inheritance, with potentially deleterious consequences. For example, overexpression of the centromeric histone CENP-A, a component of the inner kinetochore, in yeast, flies, and human cells causes chromosome mis-segregation and genomic instability [Heun et al. 2006; Au et al. 2008; Shrestha et al. 2017]. Additionally, overexpression of the outer kinetochore subunits, such as Hec1 [also known as Ndc80] or SKA1, has been observed in many types of cancers and implicated in tumorigenesis [Chen et al. 1997, 2018; Hayama et al. 2006; Li et al. 2014; Shen et al. 2016].

Aside from these pathological states, changes to kinetochore composition also occur in physiological contexts. In various organisms, the kinetochore undergoes...
extensive remodeling during meiotic differentiation (Asakawa et al. 2005; Sun et al. 2011; Miller et al. 2012; Kim et al. 2013; Meyer et al. 2015), which is the developmental program that generates reproductive cells through two consecutive nuclear divisions. Specifically in budding yeast, the Ndc80 complex disassembles in early meiosis and reassembles during the meiotic divisions, thereby restricting kinetochore activity in a temporal fashion (Asakawa et al. 2005; Miller et al. 2012; Meyer et al. 2015; Chen et al. 2017). This dynamic kinetochore behavior is driven by the fluctuating Ndc80 levels, which are barely detectable in meiotic prophase but become highly abundant during the meiotic divisions. Failure to temporally regulate Ndc80 protein levels and kinetochore activity causes defects in meiotic chromosome segregation and gamete inviability (Miller et al. 2012; Chen et al. 2017), highlighting the importance of Ndc80 regulation.

One way to regulate Ndc80 protein levels occurs through controlling Ndc80 synthesis. Ndc80 production is relatively high during the meiotic divisions, but is completely shut down in meiotic prophase (Fig. 1A; Chen et al. 2017; Chia et al. 2017). This repression in synthesis requires the expression of a meiosis-specific, 5′ extended mRNA expressed from an alternate NDC80 promoter. This transcript, called LUTI (long undecoded transcript isoform), is induced by the transcription factor complex Ime1–Ume6 after meiotic entry and cannot be translated into Ndc80 protein. Instead, NDC80LUTI expression acts to interfere with the transcription of the canonical, protein-coding NDC80 mRNA isoform.

As a result, in meiotic prophase, a stage when NDC80LUTI is highly expressed, Ndc80 protein synthesis is turned off. After cells exit from meiotic prophase, transcription of the coding NDC80 isoform is induced by another transcription factor called Ndt80, leading to reexpression of Ndc80 and kinetochore activation (Chen et al. 2017). Thus, the developmentally coordinated toggling between these two functionally distinct mRNA isoforms controls Ndc80 production in meiosis.

The LUTI-based regulation explains how meiotic cells can effectively repress Ndc80 protein synthesis. However, since Ndc80 is clearly detected at meiotic entry (Asakawa et al. 2005; Miller et al. 2012; Meyer et al. 2015; Chen et al. 2017), regulated Ndc80 synthesis alone cannot fully explain kinetochore inactivation in meiotic prophase. Additional mechanisms must be in place to clear the existing pool of Ndc80 such that the kinetochores can disassemble in a timely manner. Interestingly, the human homolog of Ndc80, Hec1, undergoes degradation in a cell-cycle-dependent manner, but the turn-over mechanism remains elusive (Ferretti et al. 2010). More generally, little is known about the factors that

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**Figure 1.** Ndc80 degradation is temporally regulated during meiosis. (A) LUTI-based regulation of Ndc80 protein synthesis in budding yeast meiosis. In meiotic prophase, the Ime1–Ume6 transcription factor complex induces a long undecoded transcript isoform of NDC80 called NDC80LUTI, which cannot produce Ndc80 protein due to the upstream open reading frames in its 5′ extension. NDC80LUTI represses transcription of a protein-coding isoform of NDC80, NDC80ORF. This combination of act of transcriptional and translational repression, NDC80LUTI inhibits Ndc80 protein synthesis. In the meiotic divisions, NDC80ORF is induced by a second transcriptional factor, Ndt80. By this combination, Ndc80 degradation can be temporally regulated.

(B) The lexO-LUTI system induces NDC80LUTI expression upon β-estradiol addition, thus conditionally inhibiting NDC80ORF expression and Ndc80 protein synthesis. (Top) Regulatory elements of the NDC80 gene. (Bottom) The lexO-LUTI system. msc, a mutant MSE site defective in Ndt80 binding. (C) Ndc80 turnover in early or late meiotic prophase. The strain carrying the lexO-LUTI (UB14883) was transferred to the sporulation medium (SPO) at 0 h to induce meiosis, and β-estradiol was added at either 1.5 or 4 h after meiosis induction. The strain was halted in meiotic prophase using an ndt80-Δ block. Here and throughout, Ndc80 levels were determined by anti-V5 immunoblot. Hsk2, loading control. Unless specified, the numbers below the immunoblots were calculated by first normalizing Ndc80 levels to Hsk2 levels in each lane, and then dividing the ratio to the 0-h time point. All the experiments in this study were performed at least twice, and one representative biological replicate is shown. (D) Induction levels of NDC80LUTI mRNA for the experiment in C, measured by reverse transcription followed by quantitative PCR (RT-qPCR). For all RT-qPCR experiments, NDC80LUTI signals were normalized to that of PFY1. [a.u.] Arbitrary unit. The mean from three independent experiments, along with the standard error of the mean, is displayed. The P-values were calculated by a two-tailed Student’s t-test. (E) Ndc80 turnover in late meiotic prophase or in metaphase I arrest. The nrd80 Δ [UB19616] and the nrd80 Δ cdc20 mn [UB19618] strains were cultured in SPO for 4 h before β-estradiol addition. Both strains were halted in meiotic prophase with an nrd80Δ block. The cdc20 meiotic null mutant [cdc20 mn, UB19678] was cultured in SPO for 5 h before β-estradiol addition and subsequently halted in metaphase I for 3 h.
mediate kinetochore subunit degradation in a developmental context.

Here, we describe the mechanism by which Ndc80 degradation is controlled in budding yeast meiosis. We found that the degradation of Ndc80 is temporally regulated. Its proteolysis in meiotic prophase requires Aurora B/Ipl1 kinase-dependent phosphorylation, which has been previously linked to correcting erroneous microtubule–kinetochore attachments (for review, see Higgins 2013). The N terminus of Ndc80, including a 27-residue sequence and Ipl1 phosphorylation sites, is both necessary and sufficient for kinetochore protein degradation. In addition to phosphorylation, Ndc80 degradation depends on the ubiquitin ligase APC

We used a strain in which the endogenous NDC80 promoter was replaced with an inducible promoter control by an array of eight lex operons ([8lexO]) (Fig. 1B). The same strain carries a chimeric IlexA-B112 transcription factor fused to an estradiol-binding domain [lexA-B112-ER], which allows inducible transcription from the 8lexO promoter in the presence of β-estradiol (Ottoz et al. 2014). Without β-estradiol [uninduced], the coding NDC80 transcript [hereafter referred to as NDC80

In meiotic prophase, the residual Ndc80 protein from the premeiotic cell cycle is turned over by an unknown mechanism (Chen et al. 2017). To study Ndc80 degradation without a confounding effect from its synthesis regulation, we took advantage of a previously established method, which allowed us to turn off Ndc80 synthesis in a conditional manner (Chia et al. 2017). Specifically, we used a strain in which the endogenous NDC80LUTI promoter was replaced with an inducible promoter controlled by an array of eight lex operators ([8lexO]) (Fig. 1B). The same strain carries a chimeric IlexA-B112 transcription factor fused to an estradiol-binding domain [lexA-B112-ER], which allows inducible transcription from the 8lexO promoter in the presence of β-estradiol (Ottoz et al. 2014). Without β-estradiol [uninduced], the coding NDC80 transcript [hereafter referred to as NDC80

We determined whether Ndc80 is degraded beyond meiotic prophase, we monitored Ndc80 levels during a metaphase I arrest induced by Cdc20 depletion [cdc20-mn]. Cdc20 is an activator of the anaphase-promoting complex, APC/C, a ubiquitin ligase necessary for metaphase-to-anaphase transition (Visintin et al. 1997; Hwang et al. 1998; Yu 2007). For this experiment, we mutated the Ndt80 binding site [also known as mid-sporelement [MSE]] at the NDC80 promoter (Chen et al. 2017). This alteration is required because the second burst of Ndc80 synthesis, which depends on the MSE site, occurs after cells exit meiotic prophase. Mutating this site ensures that Ndc80 synthesis can be repressed by β-estradiol addition even after meiotic prophase. We found that while Ndc80 was degraded in meiotic prophase, it remained remarkably stable during the metaphase I arrest induced by cdc20-mn [Fig. 1E]. The level of NDC80LUTI induction was ~40% lower in cdc20-mn cells than in wild type [Supplemental Fig. S1B]. In principle, this reduction of NDC80LUTI could cause an increase in Ndc80 synthesis, leading to higher protein levels. To exclude this possibility, we used cycloheximide to globally inhibit protein synthesis. Ndc80 was still stable during the metaphase I arrest and degraded in late prophase I under these conditions [Supplemental Fig. S1C], suggesting that the stability of Ndc80 protein differed between the two states. While it is possible that APC

Using this system, we examined Ndc80 turnover at different stages of meiosis to determine the specific time window of Ndc80 degradation. We treated cells with β-estradiol either close to meiotic entry (1.5 h after meiotic induction) or later (4 h after meiotic induction). Meanwhile, the cells were held in meiotic prophase by deletion of NDT80, which encodes a transcription factor required for meiotic progression. We found that Ndc80 was degraded with similar kinetics in either condition [Fig. 1C]. The levels of NDC80LUTI induction were also similar, as measured by reverse transcription followed by quantitative polymerase chain reaction (RT-qPCR) [Fig. 1D], suggesting that Ndc80 synthesis was successfully repressed. This result suggests that Ndc80 turnover can occur throughout meiotic prophase.

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To identify the regulators of Ndc80 degradation, we surveyed the key cellular and proteolytic events in meiotic prophase. We found that synopsis, recombination, and DNA replication were all dispensable for Ndc80 degradation. Ndc80 degradation was normal in spo11Δ cells despite the lack of recombination and synopsis [Fig. 2A; Giroux et al. 1989; Cao et al. 1990; Keeney et al. 1997]. Ndc80 degradation also occurred normally in cells depleted of the DNA replication factor Cdc6 [cdc6-mn] (Hochwagen et al. 2005; Brar et al. 2009; Blitzblau et al. 2012), consistent with a previous report [Fig. 2A; Meyer et al. 2015]. These results suggest that Ndc80 degradation, and thus kinetochore remodeling, is independent of major meiosis-specific changes to chromosomes.

Next, we tested the ubiquitin-proteasome system, a key protein degradation pathway in the cell [Finley et al. 2012]. In meiotic prophase, proteasomes are localized to chromosomes (Ahuja et al. 2017) and could mediate Ndc80 degradation. We treated meiotic prophase cells with the proteasome inhibitor MG132. Compared with
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∼did not change significantly, but it had decreased by 
realized within 1 h of MG132 treatment (Supplemental Fig. 
the vehicle control, Ndc80 levels were modestly stabi-
ized within 1 h of MG132 treatment (Supplemental Fig. 
the differences in Ndc80 synthesis, we performed a cyclo-
heximide addition (3.5 h). (For either condition (DMSO or MG132), all of the time points 
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For either condition (DMSO or MG132), all of the time points 
This result indicates that the proteasome contributes, at least partially, 
to Ndc80 degradation.

Previously, it has been shown that the decline of Ndc80 levels in meiotic prophase requires the kinase Aurora B/ 
Ipl1 (Meyer et al. 2015), raising the possibility that Ipl1 
regulates Ndc80 degradation. We confirmed that Ndc80 
abundance was increased upon meiotic depletion of Ipl1 
(ipl1-mn) (Fig. 2C). This increase was not the result of 
elevated Ndc80 synthesis, as shown by three observa-
tions. First, the level of NDC80 
 mRNA was not altered in 
ipl1-mn mutants (Supplemental Fig. S2C). Second, Ipl1 depletion increased Ndc80 levels additively 
with a mutant that fails to repress Ndc80 synthesis 
(ΔNDC80LUTI) (Fig. 2C). Finally, the expression of the protein-coding NDC80 
isoform did not significantly 
change in the double mutant (ΔNDC80LUTI ipl1-mn) 
compared with the single mutant (ΔNDC80LUTI), as shown by single molecule RNA fluorescence in situ hybridization 
(smFISH) (Fig. 2D, Supplemental Fig. S2D). Based on these 
data, we conclude that Ipl1 regulates Ndc80 turnover 
rather than synthesis.

Ndc80 degradation requires Ipl1-mediated phosphorylation

How does Ipl1 regulate Ndc80 abundance mechanistical-
ly? In one model, Ipl1 depletion may alter Ndc80 abun-
dance indirectly by affecting microtubule behavior in 
meiotic prophase. At this meiotic stage, the yeast centro-
somes, known as the spindle pole bodies (SPBs), are duplic-
ated but prevented from separating to form spindle 
microtubules. Meanwhile, the kinetochores become 
dispersed from the SPBs, presumably due to the lack of 
microtubule–kinetochore interactions (Kim et al. 2013; 
Meyer et al. 2013, 2015). Both kinetochore dispersion 
and inhibition of spindle formation in meiotic prophase 
require Aurora B/Ipl1 activity (Kim et al. 2013; Meyer 
et al. 2013, 2015). ipl1-mn mutants prematurely separate 
the duplicated SPBs and form long microtubules (Shirk 
et al. 2011; Kim et al. 2013). These microtubules interact 
with the kinetochores, leading to kinetochore reclustering 
(Meyer et al. 2013, 2015). In this context, microtubules 
could shield Ndc80 from degradation. Accordingly, the 
effect of ipl1-mn would depend on the presence of micro-
tubules. However, when we depolymerized microtubules 
in ipl1-mn cells using a microtubule poison cocktail 
[benomyl and nocodazole], Ndc80 remained stable in 
meiotic prophase (Supplemental Fig. S3A–C). Thus, it is 
unlikely that Ipl1 indirectly regulates Ndc80 degradation 
by affecting microtubule behavior.

Since Ipl1 is a kinase, it may promote Ndc80 turnover 
by phosphorylating factors that turn over Ndc80, or by phosphorylating Ndc80 itself. Previous work has shown 
that Ndc80 is a direct substrate of Ipl1 (Cheeseman et al. 
2002; Akiyoshi et al. 2009). Ndc80 has seven known Ipl1 
consensus sites ([KR]-X-[ST]-[ILVST]) (Cheeseman et al. 
2002), which are required for Ipl1 to phosphorylate 
Ndc80 in vitro (Akiyoshi et al. 2009). To test whether 
Ndc80 is degraded through a phosphorylation-dependent 
mechanism, we first asked whether Ndc80 is phosphory-
lated at the time of its degradation. We immunoprecipit-
eted Ndc80 from meiotic prophase cells 1 h after MG132 
treatment and analyzed the posttranslational modificati-
ons by mass spectrometry. We found two recurring, 
high-confidence phosphorylation sites T54 and T248 on 
Ndc80 (Fig. 3A). The first site has been previously

the vehicle control, Ndc80 levels were modestly stabi-
ized within 1 h of MG132 treatment (Supplemental Fig. 
Supplemental Fig. S2A). During this time, the expression of NDC80LUTI did not change significantly, but it had decreased by 
~50% by the second hour of treatment (Supplemental Fig. S2B). To exclude the confounding effects from the 
differences in Ndc80 synthesis, we performed a cyclo-
heximide chase experiment. The levels of Ndc80 were 
stable in the first hour of cycloheximide treatment 
though they eventually declined (Fig. 2B). This result 
indicates that the proteasome contributes, at least partially, 
to Ndc80 degradation.

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Ipl1 (Meyer et al. 2015), raising the possibility that Ipl1 
regulates Ndc80 degradation. We confirmed that Ndc80

Figure 2. Proteasome and the Aurora B/Ipl1 kinase regulate Ndc80 degradation. (A) Dependency of Ndc80 protein degradation on recombination, synopsis, or DNA replication. The wild-type (UB1338), spo11A (UB11793), and cdc6 meiotic null (cdc6-mn, UB13656) strains were transferred to SPO at 0 h and 
halted in meiotic prophase using the pGAL-NDT80 GAL4.ER 
system until 6 h in SPO. The numbers below the immunoblots were 
calculated by first normalizing Ndc80 levels to Hxk2 levels 
in each lane, and then dividing the ratio to the 0-h time 
point. (B) Dependency of Ndc80 degradation on active protea-
somes. Cells (pdr5Δ, UB2405) were induced to sporulate at 0 h. After 3 h in SPO, cells were split and treated with either 
DMSO or the proteasome inhibitor MG132 (100 μM). Thirty minutes later, cycloheximide [CHX] was added (0.2 mg/mL). For either condition (DMSO or MG132), all of the time points 
were normalized to the time point immediately before the 
cycloheximide addition (3.5 h). (C) The effects on Ndc80 levels 
when the meiotic depletion of Ipl1 [ipl1-mn] was combined 
with a mutant that fails to repress Ndc80 synthesis 
(ΔNDC80LUTI). The wild-type [UB1338], ipl1-mn [UB1013], 
ΔNDC80LUTI [UB2932], and ΔNDC80LUTI ipl1-mn [UB3948] 
cells were sporulated as in A. (D) smFISH quantification of the 
NDC80LUTI and NDC80LUTI mRNA levels in the ΔNDC80LUTI 
cells (UB2932) and ΔNDC80LUTI ipl1-mn (UB3948) cells in mei-

totic prophase. Samples were taken at 5 h in SPO. The relative 
frequence histograms of the cells with a given number of 
NDC80LUTI and NDC80LUTI transcripts per cell were graphed 
using the data pooled from two independent experiments. 
(Dashed line) The median number of transcripts per cell. A total of 
218 cells was counted for ΔNDC80LUTI and 208 cells for 
ΔNDC80LUTI ipl1-mn. Two-tailed Wilcoxon rank sum test was 
performed.
characterized and contains an Ipl1 consensus site (Cheeseman et al. 2002; Akiyoshi et al. 2009), whereas the second site lacks the Ipl1 consensus sequence.

Next, we mutated the serine or threonine in all seven Ipl1 consensus sites, generating the allele (NDC80-7A) known to greatly reduce Ndc80 phosphorylation by Ipl1 in vitro (Akiyoshi et al. 2009). The mutations did not affect NDC80LUTI expression (Supplemental Fig. S3D), ruling out the possibility that the synthesis repression of Ndc80 was disrupted. We found that Ndc80-7A was highly stable in meiotic prophase (Fig. 3B). Mutating additional serine and threonine residues around T54 and T248 (NDC80-14A) did not enhance Ndc80 stabilization (Fig. 3B). Thus, we conclude that the seven Ipl1 consensus sites at the N-terminal region of Ndc80 are required for its turnover in meiotic prophase.

We further asked whether the stabilized Ndc80-7A proteins localized to the kinetochores and affected kinetochore behavior. We fused Ndc80-7A to the enhanced green fluorescent protein (eGFP), and generated a mCherry-tagged allele for the inner kinetochore protein Mtw1, which remains chromosome-associated throughout meiotic prophase. For both the wild-type and NDC80-7A mutant, over 85% of the cells had clustered kinetochores at meiotic entry (0 h in SPO), and all of the kinetochore clusters had Ndc80-eGFP signal (Fig. 3C,D). When the wild-type cells progressed into meiotic prophase (3–5 h in SPO), the kinetochores became dispersed in >70% of the cells, and none of the dispersed kinetochores had Ndc80-eGFP signal (Fig. 3D). In contrast, fewer than 30% of the NDC80-7A cells had dispersed kinetochores after 3 h in SPO (Fig. 3D). Instead, ~40% of the cells had partially clustered kinetochores (Fig. 3C, partial and inset), all of which had Ndc80-7A-eGFP on them. By 5 h, >50% of the NDC80-7A cells had dispersed kinetochores (Fig. 3D), and about half of these cells had Ndc80-7A-eGFP on at least one kinetochore (Fig. 3E). These results demonstrate that the stabilized Ndc80-7A proteins localize to the kinetochore and affect the timing of kinetochore dispersion in meiotic prophase.

Ndc80 degradation requires a specific sequence at its N terminus

Since the seven Ipl1 consensus sites are located in the 112-residue N-terminal tail of Ndc80, we systematically truncated this region to narrow down the sites necessary for Ndc80 degradation. We found residues 2–28 to be necessary for the decline of Ndc80 levels (Fig. 4A, top panel). Within this segment, residues 11–19 were the most critical (Fig. 4A, bottom panel). To our surprise, the decline of Ndc80 levels was not altered when the 11 serines and threonines in the first 30 residues of Ndc80 were mutated (NDC80-11A) (underlined in Fig. 4B), indicating that Ndc80 degradation does not depend on the potential phosphorylation sites within this region. Ndc80 turnover was also normal when the four histidines in the 11–19 region [green letters in Fig. 4B] were mutated to either alanines (NDC80-4A) or leucines (NDC80-4L) (Supplemental Fig. S4A).

How does the 2–28 region regulate Ndc80 abundance? We first confirmed that the 2–28 region regulates Ndc80 degradation rather than Ndc80 synthesis. The wild-type, Δ2–28, and Δ11–19 cells had similar NDC80LUTI levels...
in meiotic prophase [Supplemental Fig. S4B]. Also, the Ndc80 protein levels were additively increased when the Δ11–19 mutant was combined with the mutant that fails to inhibit Ndc80 synthesis (ΔNDC80LUTI) [Fig. 4C].

Despite the differences in protein levels, the expression of the coding NDC80 ORF mRNA was not significantly different between the double mutant (ΔNDC80LUTI Δ11–19) and the single mutant (ΔNDC80LUTI) [Supplemental Fig. S4C,D], suggesting that the 2–28 residues regulate Ndc80 stability.

Since Ndc80 degradation requires Ipl1-dependent phosphorylation, we next asked whether such phosphorylation depends on the 2–28 residues. We immunoprecipitated Ndc80(Δ2–28) protein from meiotic prophase and performed mass spectrometry. Both phosphorylation sites [T54 and T248] observed in the wild-type Ndc80 protein were detected [Fig. 3A]. In addition, we performed an in vitro kinase assay using a recombinant Ipl1 protein variant [AurB∗] (de Regt et al. 2018) and found no noticeable difference in the degree of Ndc80 phosphorylation between the wild-type and Ndc80(Δ2–28) protein [Fig. 4D]. These results strongly suggest that the Ipl1-dependent Ndc80 phosphorylation occurs normally in the Δ2–28 mutant. Thus, we conclude that the 2–28 region is required for Ndc80 degradation at a step downstream from or in parallel to Ipl1-dependent phosphorylation.

We attempted to understand the role of the 2–28 residues in Ndc80 degradation by looking for changes in the binding partners of the wild-type or Ndc80(Δ2–28) protein during meiotic prophase. By quantitative mass spectrometry using tandem mass tags, we found that Sis1, a J domain protein that regulates heat-shock protein activity [Kampinga and Craig 2010], interacted with the wild-type but not the Ndc80(Δ2–28) protein during meiotic prophase. By quantitative mass spectrometry using tandem mass tags, we found that Sis1, a J domain protein that regulates heat-shock protein activity [Kampinga and Craig 2010], interacted with the wild-type but not the Ndc80(Δ2–28) protein during meiotic prophase. By quantitative mass spectrometry using tandem mass tags, we found that Sis1, a J domain protein that regulates heat-shock protein activity [Kampinga and Craig 2010], interacted with the wild-type but not the Ndc80(Δ2–28) protein during meiotic prophase. By quantitative mass spectrometry using tandem mass tags, we found that Sis1, a J domain protein that regulates heat-shock protein activity [Kampinga and Craig 2010], interacted with the wild-type but not the Ndc80(Δ2–28) protein during meiotic prophase.

Despite acting through an unknown mechanism, the 2–28 residues of Ndc80 are evidently important for clearing Ndc80 from the kinetochores and facilitating kinetochore dispersion in meiotic prophase. For the Δ2–28 mutant, we observed an increased proportion of cells with partially clustered kinetochores after 3 h in SPO
tor of the APC/C, Ama1, to be required for Ndc80 turnover [Fig. 6A]. APC\textsuperscript{Ama1} regulates Ndc80 degradation rather than synthesis, because the levels of \textit{NDC80\textsuperscript{LUTI}} were not significantly altered in the \textit{ama1} mutant [Supplemental Fig. S6A]. A cycloheximide chase experiment further supported that APC\textsuperscript{Ama1} acts at a posttranslational step in regulating Ndc80 levels [Fig. 6C,D]. In addition to Ndc80, the Ame1-tail fusion was also stabilized in an APC\textsuperscript{Ama1}-dependent manner [Fig. 6B].

It has been shown previously that \textit{AME1} deletion causes a mitotic-like cellular state in meiotic prophase [Okaz et al. 2012]. This occurs due to stabilization of two APC\textsuperscript{Ama1} substrates: Ndd1, a transcription factor required for the mitotic expression of B-type cyclins and Clb4, a B-type cyclin itself. Therefore, it is possible that such a mitotic-like state in the \textit{ama1} mutant could mimic a condition in which Ndc80 is stable, such as metaphase I. However, we found no evidence in support of this model. In our strains, premature spindle formation, a phenotype associated with elevated cyclin/CDK activity, was not apparent in the \textit{ama1} mutant [Supplemental Fig. S6B]. As an additional test, we removed Ndd1 and Clb4 from the \textit{ama1} mutant and found that Ndc80 was still stable [Fig. 6C,D, Supplemental Fig. S6C], suggesting that \textit{AME1} deletion does not stabilize Ndc80 levels through creating an alternative cellular state in meiotic prophase.

Next, we tested whether Aurora B/Ipl1 activity is disrupted in the \textit{ama1} mutant, as Ipl1 down-regulation would lead to Ndc80 stabilization. We found that the wild-type and \textit{ama1} mutant cells had comparable levels of serine 10 phosphorylated histone H3 [Fig. 6E], which is an established Aurora B/Ipl1 substrate [Hsu et al. 2000]. Thus, Ipl1 activity is unaffected by \textit{AME1} deletion. These observations led us to conclude that APC\textsuperscript{Ama1} acts downstream from Ipl1-dependent phosphorylation to mediate Ndc80 degradation.

\textbf{Defects in Ndc80 degradation predispose meiotic cells to chromosome segregation errors}

How does Ndc80 degradation impact kinetochore activity and function? We used the \textit{\Delta2–28} mutant to address this question. We found that of the APC/C, Ama1, to be required for Ndc80 turnover [Fig. 6A]. APC\textsuperscript{Ama1} regulates Ndc80 degradation rather than synthesis, because the levels of \textit{NDC80\textsuperscript{LUTI}} were not significantly altered in the \textit{ama1} mutant [Supplemental Fig. S6A]. A cycloheximide chase experiment further supported that APC\textsuperscript{Ama1} acts at a posttranslational step in regulating Ndc80 levels [Fig. 6C,D]. In addition to Ndc80, the Ame1-tail fusion was also stabilized in an APC\textsuperscript{Ama1}-dependent manner [Fig. 6B].

It has been shown previously that \textit{AME1} deletion causes a mitotic-like cellular state in meiotic prophase [Okaz et al. 2012]. This occurs due to stabilization of two APC\textsuperscript{Ama1} substrates: Ndd1, a transcription factor required for the mitotic expression of B-type cyclins and Clb4, a B-type cyclin itself. Therefore, it is possible that such a mitotic-like state in the \textit{ama1} mutant could mimic a condition in which Ndc80 is stable, such as metaphase I. However, we found no evidence in support of this model. In our strains, premature spindle formation, a phenotype associated with elevated cyclin/CDK activity, was not apparent in the \textit{ama1} mutant [Supplemental Fig. S6B]. As an additional test, we removed Ndd1 and Clb4 from the \textit{ama1} mutant and found that Ndc80 was still stable [Fig. 6C,D, Supplemental Fig. S6C], suggesting that \textit{AME1} deletion does not stabilize Ndc80 levels through creating an alternative cellular state in meiotic prophase.

Next, we tested whether Aurora B/Ipl1 activity is disrupted in the \textit{ama1} mutant, as Ipl1 down-regulation would lead to Ndc80 stabilization. We found that the wild-type and \textit{ama1} mutant cells had comparable levels of serine 10 phosphorylated histone H3 [Fig. 6E], which is an established Aurora B/Ipl1 substrate [Hsu et al. 2000]. Thus, Ipl1 activity is unaffected by \textit{AME1} deletion. These observations led us to conclude that APC\textsuperscript{Ama1} acts downstream from Ipl1-dependent phosphorylation to mediate Ndc80 degradation.

Defects in Ndc80 degradation predispose meiotic cells to chromosome segregation errors

How does Ndc80 degradation impact kinetochore activity and function? We used the \textit{\Delta2–28} mutant to address this question. We found that...
question. We first tested whether this mutant had growth defects at a higher temperature (37°C) or on plates containing the microtubule depolymerizing drug benomyl. These two phenotypes are often exhibited by mutants with defects in kinetochore function (Spencer et al. 1990; Wigge et al. 1998; Hyland et al. 1999) and spindle assembly checkpoint (e.g., mad2Δ) (Li and Murray 1991). We found that the Δ2–28 mutant grew similarly to the wild-type cells in both conditions (Fig. 7A), demonstrating that these amino acids in Ndc80 are dispensable for normal kinetochore function in mitotic cells.

To examine whether Ndc80 degradation affects kinetochore function in meiosis, we monitored chromosome movements using the centromeric TetR/TetO GFP dot assay [Michaelis et al. 1997], where both homologs of chromosome V contained TetO arrays that were integrated near the centromere and were bound by the TetR-GFP fusion protein [CENV-GFP]. We performed our experiment in the spo11Δ background because (1) these cells have long spindles that help resolve chromosome movements [Shonn et al. 2000] and (2) the homologous chromosomes fail to recombine [Klapholz et al. 1985], which renders the spo11Δ cells unable to establish biorientation. As a result, multiple rounds of microtubule–kinetochore attachments and detachments occur for an extended period, as part of the error correction mechanism. Ipl1 inactivation is known to cause defects in error correction, which is manifested by a reduced frequency of kinetochore detachments in the spo11Δ cells [Meyer et al. 2013]. We found a similar distribution of kinetochore detachments between the wild-type and Δ2–28 cells (Fig. 7B,C), suggesting that the frequency of error correction [and, by inference, the kinetochore function] is unaltered upon Ndc80 stabilization. Consistently, >95% of the wild-type and the Δ2–28 tetrads correctly segregated their chromosomes, as shown by tracking the segregation of the homozygous CENV-GFP dots (Fig. 7D, Supplemental Fig. S7A). Both the wild-type and Δ2–28 cells had >95% sporulation efficiency (Fig. 7E) and >96% spore viability (Supplemental Fig. S7B,C), a metric that would be lowered if chromosomes missegregated. These results suggest Ndc80 degradation is dispensable for correcting erroneous microtubule–kinetochore interactions in meiosis.

Instead, we found that Ndc80 degradation modulated kinetochore activity by changing kinetochore composition. Remodeling kinetochore composition is crucial for proper meiosis I. In meiotic prophase, the outer kinetochore dissociates from the inner kinetochore to down-regulate kinetochore activity, which prevents premature microtubule–kinetochore interactions and is crucial to establish a meiosis I-specific chromosome segregation pattern [Asakawa et al. 2005; Miller et al. 2012; Meyer et al. 2015; Chen et al. 2017]. When the Δ2–28 or Δ11–19 mutant were combined with a mutant that prematurely forms spindle microtubules in meiotic prophase (CUP-CLB3), we observed that sister chromatids, instead of homologous chromosomes, segregated in meiosis I (Fig. 7E). In this sensitized background, the NDC80-7A mutant also caused precocious sister chromatid segregation, albeit modestly (Fig. 7E), likely because the level of Ndc80-7A eventually reduced in prolonged prophase arrest (Fig. 3B, 6 h). The same phenotype occurs when Ndc80 is over-expressed in meiotic prophase or when the LUTI-based repression of Ndc80 synthesis is disrupted [Miller et al. 2012; Chen et al. 2017]. Therefore, the lack of Ndc80 turnover in meiotic prophase prematurely activates the kinetochore by providing a high abundance of Ndc80, which is the limiting subunit of the kinetochore activity in meiotic prophase, and predisposes cells to meiotic chromosome

Figure 6. APCAma1 regulates Ndc80 degradation. (A) Dependency of Ndc80 levels on Ama1 in meiotic prophase. Sporulation of the wild-type (UB6598) and ama1Δ (UB22499) cells was synchronized with the pCUP-IME1/pCUP-IME4 system. Meiotic entry was induced by CuSO4 addition after cells were incubated in SPO for 2 h. The numbers below the immunoblots were calculated by first normalizing Ndc80 levels to Hxk2 levels in each lane, and then dividing the ratio to the 0-h time point. (B) Dependency of Ame1-tail fusion levels on Ama1 in meiotic prophase. The AME1-tail AMA1 (UB20354) and AME1-tail ama1Δ (UB23001) strains were sporulated as in Figure 2A. Quantification of Ndc80 and Ame1 were performed as in Figure 5B. [rel] Relative. (C) Assessing the requirement of Ndd1 and Clb4 in the Ndc80 stabilization that was induced by ama1Δ. The ama1Δ (UB20223) and pCLB2-NDD1 (ndd1-mn) clb4Δ ama1Δ (UB22850) cells were cultured in SPO for 3 h before a final concentration of 0.2 mg/ml cycloheximide was added. During the entire CHX experiment, both strains were halted in meiotic prophase. (D) The Ndc80 protein levels relative to the time point immediately before CHX addition. The average and the standard errors of the mean for five independent experiments are graphed. (E) Ipl1 activity in the ama1Δ mutant. The wild-type (UB3954), ipl1-mn (UB1013), and ama1Δ (UB20223) cells were sporulated as in Figure 2A. The levels of H3 serine 10 phosphorylation (H3S10-P) were detected by a phospho-specific antibody. The number below each lane was first normalized to Hxk2 levels and then to the 0-h time point of the wild-type strain.

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kinetochore dispersion, causes premature activation of Ndc80 in meiotic prophase alters the kinetics of drive the degradation of kinetochore proteins. The failure of the Ipl1 consensus sites, is both necessary and sufficient to fication known for correcting erroneous microtubule phosphorylation of Ndc80, a posttranslational modi-

Ndc80 degradation is coupled to Aurora B/Ipl1-mediated gase APCAma1 and proteasome activity. Furthermore, Ndc80 degradation is independent of major chro-

mosome remodeling events, but requires the ubiquitin li-
gase APCAn1 and proteasome activity. Moreover, Ndc80 degradation is coupled to Aurora B/Ipl1-mediated phosphorylation of Ndc80, a posttranslational modi-

fication known for correcting erroneous microtubule–kinetochore attachments. The N terminus of Ndc80, which includes a 27-residue sequence and the Aurora B/ Ipl1 consensus sites, is both necessary and sufficient to drive the degradation of kinetochore proteins. The failure to degrade Ndc80 in meiotic prophase alters the kinetics of kinetochore dispersion, causes premature activation of ki-

netochores, and predisposes meiotic cells to chromosome segregation errors. Lastly, disrupting both Ndc80 synthe-
sis repression (ΔLUTT) and Ndc80 degradation (Δ11–19) synergistically enhanced this segregation phenotype [Sup-

plemental Fig. S7D], consistent with the idea that both modes of regulation contribute to kinetochore inactiva-
tion in meiotic prophase.

Discussion

In this study, we showed that Ndc80 degradation in mei-

osis is a temporally regulated process. The proteolysis of Ndc80 in meiotic prophase is independent of major chro-

mosome remodeling events, but requires the ubiquitin li-
gase APCAn1 and proteasome activity. Furthermore, Ndc80 degradation is coupled to Aurora B/Ipl1-mediated phosphorylation of Ndc80, a posttranslational modi-

fication known for correcting erroneous microtubule–kinetochore attachments. The N terminus of Ndc80, which includes a 27-residue sequence and the Aurora B/ Ipl1 consensus sites, is both necessary and sufficient to drive the degradation of kinetochore proteins. The failure to degrade Ndc80 in meiotic prophase alters the kinetics of kinetochore dispersion, causes premature activation of ki-

Figure 7. Ndc80 degradation regulates kinetochore composition in meiosis. [A] Growth phenotype associated with the various truncations of the Ndc80 N-terminal tail. Tested strains included the temperature-sensitive NDC80 allele (ndc80-1, UB494), the spindle assembly checkpoint mutant Δmad2 (UB700), wild-type NDC80 (UB3262), and the NDC80 mutants that carry the following deletion: residues 2–112 (Δ2–112, UB3275), residues 29–56 (Δ29–56, UB4699), and residues 2–28 (Δ2–28, UB5015). Cells were serially diluted and grown on plates containing nutrient-rich medium (YPD) at 30°C or 37°C, as well as on a benomyl plate (15 µg/mL) at 33°C. [B] Frequency distribution of the cells with a given number of kinetochore detachments and re-attachments for the experiment in C. The mean percentage and the standard deviation for three biological replicates are displayed. [C] Kinetochore at-

tachments and detachments for the NDC80 (UB15905) and Δ2–28 (UB15873) strains in the spo11Δ background. Each strain carries the TetR-GFP fusion protein, and both homologs of chromo-

some V have the centromeric TetO array [homozygo-
gous CENV-GFP, marked in green]. [Magenta] Spec42 fused with mCherry, which marks the spindle pole body (SPB). White arrowhead marks the kinetochore that would undergo detachment. The time when the SPBs were split is marked as 0. Scale bar, 5 µm. [D] The percentage of cells correctly segregated chromosome V for the wild-type (UB21757) and Δ2–28 (UB21758) strains. These strains carry the TetR-GFP fusion proteins, and both homologs of chromosome V are marked by the centromeric TetO re-

peats (CENV-GFP). Strains were sporulated for 7 h before formaldehyde fixation. At least 90 cells were counted for each strain. For D and E, the mean and the range of two biological replicates are graphed. [E] Spore formation for the meiosis I of wild-type (UB21756) and Δ2–28 (UB21758) strains. [F] Sister chromatid segregation in wild-type (UB2942), pCUP-CLB3 NDC80 (UB877), pCUP-CLB3 pCUP-NDC80 (UB880), pCUP-CLB3 NDC80Δ11-19 (UB16561), pCUP-CLB3 NDC80Δ2-28 (UB16565), and pCUP-CLB3 NDC80Δ7A (UB16872). All strains carry pGAL-NDT80 and GAL4-ER, which allows reversible arrest in meiotic prophase. A pair of the chromosome V sister chromatids is marked by the centromeric TetO repeats. A schematic for normal and abnormal meiosis I is shown at the right. In an abnormal meiosis I, two separated GFP dots were observed in binucleates, one in each nucleus. Cells were sporulated for 5 h before addition of CuSO4 to induce cyclin Clb3 expression. Immediately after the induction, cells were released from the prophase arrest using β-estradiol, which induces NDT80. Samples were taken 1 h 45 min after the release. The average fraction of binucleates that displayed sister segre-

gation in meiosis I, as well as the range of two biological replicates, is graphed. One-hundred cells were counted per strain per experiment.

Phosphorylation-mediated Ndc80 degradation in meiotic prophase

Multiple lines of evidence support a model in which Au-

ora B/Ipl1-dependent phosphorylation of Ndc80 triggers its degradation in meiotic prophase. First, Ipl1 is required for Ndc80 degradation. Second, phosphorylation of an Ipl1 consensus site on Ndc80 is detected by mass spectrometry in meiotic prophase, a stage when Ndc80 degradation occurs. Third, mutating the seven Ipl1 consensus sites leads to Ndc80 stabilization in meiotic prophase, suggest-

ing that these sites are necessary for Ndc80 degradation. Altogether, these results are consistent with a phosphory-

lation-dependent degradation mechanism.

Given that Ndc80 phosphorylation is not disrupted when the 2–28 residues are deleted, we posit that this 2–28 segment may act in parallel to or downstream from the Ipl1-dependent phosphorylation to promote Ndc80 degradation. These residues may serve as the binding site for protein factors that mediate Ndc80 proteolysis.
It is possible that such factors are recruited by Ndc80 phosphorylation, or that phosphorylation alters the local conformation around the 2–28 residues to expose this segment for protein factor binding.

Our data are consistent with the idea that the ubiquitin ligase APC\(^{\text{Ama1}}\) and the proteasome act downstream from the Ipl1-dependent phosphorylation to mediate Ndc80 proteolysis. We excluded the possibility that the \textit{ama1}Δ mutation stabilizes Ndc80 by increasing cyclin/CDK activity or by down-regulating Ipl1 activity [Fig. 6C–E]. It is currently unknown whether APC\(^{\text{Ama1}}\) directly ubiquitinates Ndc80 or regulates other player(s) in the degradation pathway to promote Ndc80 turnover. Interestingly, the APC/C degron repository [Davey laboratory and Morgan laboratory] predicts that Ndc80 has two D-box motifs and three KEN motifs, which are known recognition sites for the APC/C [Glotzer et al. 1991]. However, these sites are located outside of the 112-residue N-terminal region, which is both necessary and sufficient for Ndc80 degradation in meiotic prophase. Furthermore, mutating the D-boxes does not block Ndc80 degradation [Supplemental Fig. S7E]. The D-box motif is also not required for the APC\(^{\text{Ama1}}\)-dependent proteolysis of Ssp1 [Maier et al. 2007; Diamond et al. 2009]. Future work to characterize the substrates of APC\(^{\text{Ama1}}\), as well as the mechanism of substrate recognition, will be important to fully grasp the meiotic regulation of Ndc80 turnover.

**Interplay between error correction and Ndc80 degradation**

It is well established that Ndc80 phosphorylation by Aurora B/Ipl1 helps correct microtubule–kinetochore attachments that do not result in tension [Biggins 2013]. Ndc80 phosphorylation is thought to weaken microtubule binding, resulting in microtubule detachment and thus providing the opportunity for the kinetochore to reattach in the correct orientation. How is the error correction process interrelated with Ndc80 degradation given that both processes require Ndc80 phosphorylation by Aurora B? We found that disrupting Ndc80 degradation alone \(\Delta\text{2–28}\) does not appear to affect meiotic chromosome segregation [Fig. 7B–E; Supplemental Fig. S7A–C]. This observation suggests that Ndc80 degradation is not required for error correction.

Instead, we propose that the completion of error correction may repress Ndc80 degradation in metaphase I, a stage when Ndc80 is stable [Fig. 1E; Supplemental Fig. S1B,C]. Once the chromosomes attach to spindles properly, Ndc80 phosphorylation is removed by phosphatases to stabilize the microtubule attachments [Biggins 2013]. Thus, it is possible that the Ndc80 levels become stable after the removal of Ndc80 phosphorylation in metaphase I. We note that it is unlikely that microtubule binding directly prevents Ndc80 degradation through steric effect because, at least in meiotic prophase, microtubule depolymerization cannot destabilize Ndc80 in \textit{ipl1–mn} cells.

Our model presents a puzzle, however. As the major microtubule-binding site, Ndc80 is required to build new attachments during error correction. In our model, Ndc80 phosphorylation would cause its degradation at a time when the presence of Ndc80 is essential. This problem could be reconciled by the fact that Ndc80 protein is upregulated as cells exit from meiotic prophase [Chen et al. 2017], which may provide extra proteins to compensate for the loss due to degradation. In addition, the activity of the ubiquitin ligase APC\(^{\text{Ama1}}\) is repressed by ClbCDK activity, which rises in prometaphase I [Oeschlaegel et al. 2005; Tsuchiya et al. 2011]. Since APC\(^{\text{Ama1}}\) is required for Ndc80 degradation, the degradation mechanism of Ndc80 could be turned off after meiotic prophase, even before error correction is completed.

**Protein turnover as a mechanism to create meiosis-specific kinetochores**

In vegetative growth, the yeast kinetochores transiently disassemble during DNA replication when the centromeric DNA replicates, but they remain bound to centromeres for the rest of the cell cycle. It has been shown that the subunit stoichiometry of the kinetochore is regulated during the cell cycle [Dhatchinamoorthy et al. 2017, 2019]. However, it is unclear whether the protein levels of the kinetochore subunits are regulated in vegetative growth. Degradation of a few kinetochore proteins has been reported in yeast. These include Dsn1 [Akiyoshi et al. 2013], Cbf13 [Kaplan et al. 1997], and the centromeric histone Cse4 in budding yeast [Herrero and Thorpe 2016; Ohkuni et al. 2016; Cheng et al. 2017], as well as Spc7 in fission yeast [Kriegenburg et al. 2014]. The degradation pathways of these subunits have been proposed to be quality controls that remove nonfunctional or excess proteins, rather than direct means to alter kinetochore function.

In contrast, regulating protein abundance is the key mechanism of controlling kinetochore activity in meiosis. The outer kinetochore disassembles in meiotic prophase and reassembles in prometaphase I. This disassembly is triggered by a reduction in Ndc80 levels, which occurs as a result of two separate mechanisms: LUTI-based repression of new Ndc80 synthesis and degradation of the existing Ndc80 pool. In meiosis, Ndc80 degradation is not a passive consequence of DNA replication but a targeted process mediated by Aurora B/Ipl1, which has also been implicated in preventing untimely spindle formation in meiotic prophase [Shirk et al. 2011; Kim et al. 2013]. We propose that Ipl1 prevents premature microtubule–kinetochore interactions through two independent pathways: one by altering microtubule behavior, and the other by triggering Ndc80 proteolysis. This dual mechanism ensures that the kinetochore interacts with spindles only after meiotic prophase. This delay in microtubule–kinetochore interaction is required for proper meiotic chromosome segregation [Miller et al. 2012].

**Cells customize kinetochore activity by controlling kinetochore assembly or disassembly**

Examples from various organisms indicate that regulating kinetochore assembly or disassembly is a common way to
customise kinetochore activity based on the specific cell cycle stage or cell type. During mitotic exit in human cells, the outer kinetochore components, known as the KMN network, dissociate from the inner kinetochore in every cell cycle. A failure to do so causes chromosome missegregation in the next cell cycle (Gascoigne and Cheeseman 2013). In C. elegans oocytes, kinetochores facilitate homolog biorientation and error sensing in meiosis I, but they do not segregate the chromosomes. Instead, the ring-complex and microtubule motors, such as dynein, segregate the chromosomes, while the kinetochore proteins disappear from the chromosomes in anaphase I (Dumont et al. 2010; Muscat et al. 2015; Davis-Roca et al. 2017; Laband et al. 2017). Lastly, in C. elegans, Drosophila, and iPSc-derived human motor neurons, some or all of the core kinetochore proteins localize to the regions of synaptic neuropil and axons that are devoid of nuclei. There they control axon outgrowth and dendritic extension in the developing sensory nervous system (Cheerambathur et al. 2019; Zhao et al. 2019). All of these examples highlight that the timing and location of kinetochore assembly are tuned to the cell cycle and cell type.

Based on our finding that the Aurora B kinase regulates the turnover of the kinetochore subunit Ndc80, it will be interesting to test whether other kinetochore restructuring events also depend on Aurora B. For example, the protein levels of Ndc80/Hec1 decline at the mitotic exit in human cells (Ferretti et al. 2010), but the mechanism is unknown. We posit that Aurora B and/or Aurora A may trigger Ndc80/Hec1 degradation at the mitotic exit in human cells, contributing to the KMN disassociation. In addition, the C. elegans Aurora B kinase, AIR-2, localizes to spindle microtubules during anaphase I (Schumacher et al. 1998; Speijiers et al. 2000; Romano et al. 2003). Thus, it is possible that AIR-2 phosphorylation causes the dissociation or degradation of one or more kinetochore subunits, leading to kinetochore disassembly at that time. Our discovery that Aurora B phosphorylation can cause the turnover of kinetochore proteins provides a new means by which Aurora B can influence kinetochore composition and activity: Aurora B regulates not only the construction but also the destruction of the kinetochore.

Materials and methods

Yeast strains and plasmids

All strains used in this study were derivatives of SK1 unless specified. The centromeric TetR/TetO GFP dot assay was first described in Michaelis et al. (1997), the ndc80Δ temperature-sensitive mutant was first described in Wigge et al. (1998), pCUP-NDC80 and pCUP-CLB3 were first described in Lee and Amon (2003); Miller et al. (2012), the meiotic deletion alleles pCLB2-3HA-CDC6 were first described in Lee and Amon (2003), pSCEC1-3HA-1Pl1, ndt80Δ, spo11Δ, pdr5Δ, mad2Δ, mad3Δ, ama1Δ, pCLB2-3HA-NDD1, cdc4Δ, NDC80-eGFP, MTW1-mCherry, and ΔNDC80ΔUTT (deletion of the 400–600 bp upstream of the NDC80 coding region). The V5 tagging plasmid was kindly provided by Vincent Guacci (University of California at Berkeley).

Previously, an NDC80 single integration plasmid was generated [Chen et al. 2017]. This plasmid includes the SK1 genomic sequence spanning from 1000 bp upstream of to 357 bp downstream from the NDC80 coding region, as well as a C-terminal fusion of the V5 epitope to NDC80 referred to as NDC80-V5 [Chen et al. 2017]. All the single integration plasmids used in this study were derived from this NDC80-V5 parent vector.

We cloned the following mutant NDC80 alleles by Gibson assembly (Gibson et al. 2009) using GBlocks gene fragments (IDT): NDC80-7A-3V5, NDC80-1A-3V5, NDC80(4H to A)-3V5, NDC80(4H to L)-3V5, NDC80-11A-3V5, and NDC80-4D-box-3V5. The 8lexO-LUTI-NDC80-3V5 construct, also cloned by Gibson assembly, has eight lex operators (lexO) and the promoter from CYCI inserted at 536 bp upstream of the translation start site of NDC80 ORF. This insertion allows lexA-B112-ER to control the expression of NDC80(1127) mRNA in a dose-dependent manner (Chia et al. 2017). The 8lexO-LUTI-mse-NDC80-3V5 construct has a mutation in the Ndt80 binding site (mse), which diminishes the Ndt80-dependent expression of NDC80ORF mRNA [Chen et al. 2017]. The pNDC80-AME1-3V5 and pNDC80-AME1-tail-3V5 (referred to as AME1-tail in the Results) constructs are controlled by the NDC80 promoter [1000 bp upstream of the NDC80 coding region], which is sufficient to repress protein synthesis in the coding region in meiotic prophage [Chen et al. 2017]. For the pNDC80-AME1-tail-3V5 construct, the 2–12 residues of Ndc80 were fused to the C terminus of Ame1.

In addition, systematic deletions of the N-terminal residues of Ndc80 [2–28, 29–56, 2–10, 11–19, 20–28, and 57–112] were made on the NDC80-3V5 parent vector using the QS site-directed mutagenesis kit (New England Biolabs). All of the single integration plasmids described above were digested with Pmel, transformed into yeast, and selected for integration at the LEU2 locus.

To generate the Δ2–28, Δ11–19, and NDC80-7A alleles at the endogenous locus of NDC80, the CRISPR/Cas9 method was used [Anand et al. 2017]. Oligonucleotides encoding the guide RNA (5′-TACATCACATGGACCCCTACCCGG-3′) were cloned into a centromeric plasmid carrying a URA3 marker and pPGK1-Cas9 (a gift from Gavin Schlissel, University of California at Berkeley). This plasmid was cotransformed into yeast along with the repair templates amplified from the respective LEU2 single integration plasmids by PCR (Δ2–28, Δ11–19, and NDC80-7A plasmids). For NDC80-7A, the following primers were used to amplify the repair template while introducing synonymous mutations to the PAM sequence to prevent re-editing: 5′-ACATGTCGCTACATCACATGGACCCCTACCCGGTTGCTT CaCAAATACCAACTGCAACATC-3′ and 5′-CTTCTTGAAT ACGCTTTGGAAGTTTTGCTTCTTATGGTCTGGATC TCTATTGCTCAG-3′. The lowercase letters correspond to the synonymous mutations that abolish the PAM site: a C-to-A mutation at the 66th nucleotide from the translation start site of Ndc80, as well as a G-to-A at the 351st nucleotide.

Yeast growth condition, synchronous sporulation, and media

To prepare for sporulation, diploid cells were grown in YPD (1% yeast extract, 2% peptone, 2% glucose, and supplemented with 22.4 mg/L uracil and 80 mg/L tryptophan) for 20–24 h at room temperature or 30°C. For optimal aeration, the total volume of the flask exceeded the volume of the medium by 10-fold. Subsequently, cells were transferred to BYTA (1% yeast extract, 2%
hacto tryptone, 1% potassium acetate, 50 mM potassium phthalate at 0.25–0.3 OD600 cells/mL and grown for another 15–17 h at 30°C. The cells were then pelleted, washed with sterile milliQ water, and resuspended at 1.85 OD600 cells/mL in sporulation (SPO) medium (0.5% [w/v] potassium acetate at pH 7, 0.02% [w/v] raffinose) at 30°C unless specified.

Two methods were used to synchronize sporulation in this study. The pCUP1-IME1 pCUP1-IME4 method was described in Berchowitz et al. (2013), in which the endogenous promoters of IME1 and IME4 were replaced with the inducible CUP1 promoter. To initiate synchronous sporulation, the expression of IME1 and IME4 was induced by adding 50 µM copper (II) sulphate 2 h after cells were transferred to SPO. The pGAL-NDT80 GAL4.ER system was used to allow cells to undergo synchronized meiotic divisions (Carlile and Amon 2008). After incubating the sporulating diploids for 5–6 h in SPO (see the figure legend for the specific arrest duration for each experiment), cells were released from pachytene arrest by inducing NDT80 expression with the addition of 1 µM β-estradiol.

RNA extraction and RT-qPCR

A previously described RNA extraction protocol was modified as described below (Koster and Timmers 2015). RNA was extracted with acid phenol:chloroform:isoamyl alcohol (125:24:1, pH 4.7) and then precipitated by isopropanol. Briefly, 1.85–3.7 OD600 of cells were harvested by centrifugation at 21,000 rcf for 1 min, and frozen in liquid nitrogen. Pellets were thawed on ice and vigorously shaken for 30 min at 65°C in 400 µL of TES buffer (10 mM Tris at pH 7.5, 10 mM EDTA, 0.5% SDS), 200 µL of acid-washed glass beads [Sigma], and 400 µL of acid phenol. The lysates were clarified by centrifugation at 21,000 rcf for 10 min at 4°C. The aqueous phase was transferred to 300 µL of chloroform, vortexed, and centrifuged at 21,000 rcf for 5 min at room temperature. To precipitate the RNA, the aqueous phase was added to 400 µL isopropanol supplemented with 0.33 M NaOAc, incubated for >2 h at −20°C, and centrifuged at 21,000 rcf for 20 min at 4°C. The precipitated RNA pellets were washed with 80% ethanol, air-dried, and resuspended in nuclease-free water.

A total of 5 µg of RNA was treated with DNase using the TURBO DNA-free kit [Thermo Fisher] and then reverse-transcribed into cDNA using SuperScript II or III [Thermo Fisher] following the standard protocol. The single-stranded cDNA was quantified by 7500 Fast real-time PCR machine [Thermo Fisher] using SYBR Green mix [Thermo Fisher]. The primers used for the RT-qPCR reactions were NDC80ORF (GGTTGAGAGCCCCGTTAAGT and ACGGTTGGTGGATAATGAGC). The Ct value for each sample was determined by taking the average Ct of triplicate reactions, and the levels of the NDC80ORF transcript were normalized to those of the FFY1 transcript.

Single-molecule RNA fluorescence in situ hybridization (smFISH)

Single-molecule RNA FISH was performed exactly as described previously (Chen et al. 2017). To quantify smFISH spots, the maximum-intensity projection of Z-stacks was generated in Fiji (Schindelin et al. 2012), different channels were split, and the images were analyzed by the StarSearch Program (Raj et al. 2008). Cell boundaries were hand-drawn. The thresholds were adjusted such that they fell within “plateaus,” as described previously (Raj et al. 2008). The same threshold was used for a given probe set in each experimental replicate. An NDC80ORF transcript was identified as a colocalized spot when the two spots found in either channel have pixels overlapped for >50%. The number of NDC80ORF transcripts in each cell was calculated by subtracting the number of CF 590 (LUIT-only) spots from the total number of spots with the Q 670 (common region) signal. To compare between different strains, data were pooled from two biological replicates, and the nonparametric Wilcoxon Rank Sum test was performed with Prism 6 [Graphpad]. The relative frequency histograms were normalized so that the maximum bin height is the same across all the histograms in the same figure.

Fluorescence microscopy

To fix cells, formaldehyde was added to approximately one OD600 cells to a final concentration of 3.7%, incubated for 15 min at room temperature, washed once with KPi/sorbitol buffer (100 mM potassium phosphate at pH 7.5, 1.2 M sorbitol), and resuspended in the VectaShield antifade mounting medium with DAPI overnight before imaging [Vector Laboratories]. The anti-fade medium preserved the samples from photobleaching during imaging. All images were acquired as Z-stacks of 20 slices (0.3-µm step size) using a DeltaVision Elite wide-field fluorescence microscope, a PCO Edge sCMOS camera [DeltaVision, GE Healthcare], a 100×/1.40 oil immersion objective, and the following filters: DAPI [EX390/18 and EM435/48], GFP/FITC [EX475/28 and EM525/48], and mCherry [EX575/25 and EM625/45]. Images were deconvolved in softWoRx software [softWoRx, GE Healthcare] with a 3D iterative constrained deconvolution algorithm [enhanced ratio] with 15 iterations.

Kinetochore classification and quantification

The three classes of kinetochores were classified as follows: The “clustered” class was defined by the presence of a single Mtw1-mCherry focus in a cell, or the presence of less than three distinct Mtw1-mCherry foci. For this class, Ndc80 signal was always detected and colocalized with the Mtw1-mCherry signal in the cells, and the DAPI signal was often compacted and round, without thread-like structures. The “dispersed” class was defined by more than six distinct Mtw1-mCherry foci located throughout the projected DAPI area of the cell. For this class, the DAPI signal was almost always thread-like, suggesting a pachytene state. The “partially clustered” class was defined by the presence of at least three distinct Mtw1-mCherry foci and all foci were located within the same half of the projected DAPI boundary of the cell.

Live-cell imaging

Time-lapse microscopy was performed in an environmental chamber heated to 30°C. Strains were induced to sporulate by transferring to SPO and halted in meiotic prophase using the pGAL-NDT80 GAL4.ER block for 5 h in SPO. Cells were then released with 1 µM β-estradiol and, 30 min later, immobilized on concanavalin A-coated, glass-bottom 96-well plates [Corning] and cultured in 100 µL of SPO [with 1 µM β-estradiol] medium in the prewarmed environmental chamber. Z-stacks of five slices (1-µm step size) were acquired using a 60×/NA1.42 oil immersion plan apochromat objective, and mCherry filter (10% T, 25-msec exposure) and FITC filter (10% T, 25-msec exposure), at a frequency of 45 sec for 90 min.

The following pipeline was used to quantify kinetochore detachment events. First, the movie was maximum-projected using Fiji (Schindelin et al. 2012). Next, the time of spindle pole body [SPB] separation and each kinetochore detachment event were tracked. The time of SPB separation is defined as the first frame when two distinct mCherry foci were detected. Kinetochore attachment is defined as a CENV-GFP focus staying in
the proximity of one SPB (less than a third of the pole-to-pole distance) for at least three consecutive time frames. A detachment event occurred when a CENV-GFP focus switched SPB or crossed the midline between the two poles (the CENV-GFP focus often stayed in the midzone for more than three consecutive frames before being recaptured by the original SPB). The total number of detachment events occurred after the splitting of the SPBs was quantified for each cell that maintained in the focal plane for the entire movie. The percentage of cells that underwent a given number of detachment events was calculated and graphed.

Denaturing immunoprecipitation (IP) and mass spectrometry (MS)

Denatured protein extracts were prepared as described in Sawyer et al. (2019), with the following modifications. After cells were harvested and dried completely, 150 µL of zirconia beads and 150 µL of lysis buffer (50 mM Tris at pH 7.5, 1 mM EDTA, 2.75 mM DTT, 1x PhosSTOP [Roche], 0.48 µg/mL Pefabloc SC [Sigma], 2 mM pepstatin A, 2x Complete EDTA Free [Roche]) were added to each tube. Cells were lysed mechanically on a Mini-Beadbeater-96 (BioSpec Products) for 5 min at room temperature. SDS was added to a final concentration of 1%, the extracts were boiled for 5 min at 95°C, and NP-40 lysis buffer (50 mM Tris at pH 7.5, 150 mM NaCl, 1% NP-40, 5% glycerol, 1x PhosSTOP, 0.64 µg/mL Pefabloc SC [Sigma], 2.67 mM pepstatin A, 3x Complete EDTA Free) was added to a final volume of 1.5 mL (i.e., diluting SDS to 0.1%). Lysates were clarified by centrifuging at 15,000 rpm for 15 min at 4°C. The total cleared lysate for each sample was pooled from three tubes (3825 µL total), added to 80 µL of pre-equilibrated anti-V5 agrose beads [Sigma], and incubated in loBind tubes (Eppendorf) for 2 h at 4°C with rotation. Using loBind tubes and low-adhesion tubes [USA Scientific] improved the amount of protein recovered from the IP. In each MS experiment, the cleared lysates were pooled from six tubes for each sample; therefore, two IP reactions were set up in parallel. The beads of the parallel IP reactions were washed once with the 3 mL of high-salt wash buffer (50 mM Tris at pH 7.5, 0.5 M NaCl, 1 mM EDTA, 1% NP-40), combined into one low-adhesion tube (i.e., 160 µL of beads final), and then washed once with 950 µL of high-salt wash buffer. Next, the beads were washed twice with each of the following: (1) 950 µL of buffer 2 (50 mM Tris at pH 7.5, 150 mM NaCl, 10 mM MgCl₂, 0.05% NP-40, 5% glycerol) and (2) 950 µL of buffer 3 (50 mM Tris at pH 7.5, 150 mM NaCl, 10 mM MgCl₂, 5% glycerol). After the last wash, the wash buffer was aspirated completely, and the beads were resuspended in 80 µL of trypsin buffer (2 M urea, 50 mM Tris at pH 7.5, 5 µg/mL trypsin) to digest the bound proteins for 1 h at 37°C with agitation. The beads were centrifuged at 100 rcf for 30 sec, and the digested peptides (the supernatant) were collected. The beads were then washed twice with 60 µL of urea buffer [2 M urea, 50 mM Tris at pH 7.5]. The supernatant of both washes was collected and combined with the digested peptides (i.e., the final volume is 200 µL). After brief centrifugation, the combined digested peptides were cleared from residual beads and frozen in liquid nitrogen.

MS was performed by the Vincent J. Coates Proteomics/Mass Spectrometry Laboratory at the University of California at Berkeley, as described in Sawyer et al. (2019). The mudPIT method was used. A nano LC column (packed in a 100-µm inner diameter glass capillary with an emitter tip) was composed of 10 cm of Polaris c185-µm packing material [Varian], followed by 4 cm of Partisphere 5 SCX [Whatman]. The column was loaded with a pressure bomb, washed extensively with buffer A, and then directly coupled to an electrospray ionization source mounted on a LTQ XL linear ion trap mass spectrometer (Thermo Fisher). Chromatography was performed on an Agilent 1200 HPLC equipped with a split line to deliver a flow rate of 300 nL/min. Peptides were eluted using either a four-step or an eight-step mudPIT procedure (Washburn et al. 2001). Buffer A was 5% acetonitrile and 0.02% heptfluorobutyric acid (HFBA). Buffer B was 80% acetonitrile and 0.02% HFBA. Buffer C was 250 mM ammonium acetate, 5% acetonitrile, and 0.02% HFBA. Buffer D was the same as buffer C, but with 500 mM ammonium acetate. Proteins were identified with the Integrated Proteomics Pipeline software [IP2, Integrated Proteomics Applications] using the parameters and cutoffs exactly as described by Sawyer et al. (2019). Information about the Ndc80 phosphorylation sites is provided in Supplemental Table S2.

Native IP, co-IP, and TMT quantitative MS

For the TMT quantitative MS and co-IP experiments, cells were sporulated in 2% SPO [2% [w/v] potassium acetate at pH 7, 0.02% [w/v] raffinose, supplemented with 40 mg/L uracil, 20 mg/L histidine, 20 mg/L leucine, 20 mg/L tryptophan] at 30°C. Increasing the concentration of acetate and supplementing amino acids improved sporulation efficiency. Four hours after the cells were transferred to 2% SPO, 200 mM PMSF [dissolved in ethanol] was added to cultures to a final concentration of 2 mM. For the MS and the co-IP experiments, 462 or 92 OD₆₀₀ of cells, respectively, were harvested by centrifuging at 1700 rcf, 4°C, for 2 min. The supernatant was discarded, and the cell pellets were transferred to 3-mL tubes, centrifuged at 21,000 rpm for 1 min at 4°C. The supernatant was aspirated completely, and the pellets were snap-frozen in liquid nitrogen.

To extract proteins, pellets were thawed on ice, and 500 µL of NP-40 native IP buffer was added (50 mM Tris at pH 7.5, 150 mM NaCl, 2 mM MgCl₂, 1% NP-40, supplemented with PhosSTOP tablet [one tablet used in 10 mL of buffer] [Sigma], Complete ULTRA protease inhibitor cocktail [one tablet used in 20 mL of buffer] [Sigma], 0.64 µg/mL Pefabloc SC [Sigma], 2.7 µM pepstatin A]. Cells were lysed with 500 µL of zirconia beads [BioSpec] in a Fast-Prep-24 5G (MP Biomedicals LLC) using the “S. cerevisiae” setting for 40 sec at room temperature. Next, the cell lysates were chilled in an ice water bath for 2 min. The tubes were punctured with a 20.5-gauge needle, and the lysates were collected in 15-mL tubes (precooled to −20°C) by centrifuging at 3000 rpm for 20–40 sec. Lysates were clarified by centrifugation at 21,000 rcf for 20 min at 4°C and snap-frozen in liquid nitrogen. The protein concentration of each lysate was determined by Bradford assay [Bio-Rad].

Ndc80-3V5 IP was performed at a protein concentration of 10 mg/mL in NP-40 native IP buffer. For the TMT quantitative MS experiment, a total of 38–40 mg of protein, pooled from 10 tubes of lysate, was added to 167 µL of pre-equilibrated anti-V5 agrose beads [Sigma] in a total volume of 4 mL [i.e., beads constituted 1/24 of the total IP volume], incubated for 2.5 h at 4°C with rotation, and then washed with each of the following: (1) once with 3.5 mL of NP-40 native IP buffer, (2) once with 950 µL of NP-40 native IP buffer, (3) twice with 950 µL of buffer 2, and (4) twice with 950 µL of buffer 3. After the last wash, the proteins on the beads were digested in trypsin buffer, washed with urea buffer, and frozen in liquid nitrogen as described in the “Denaturating immunoprecipitation [IP] and mass spectrometry [MS]”. The TMT quantitative MS was performed as described in Cheng et al. (2018). The raw data can be accessed at MassIVE MSV00084664 [https://massive.ucsd.edu/ProteoSAFe/static/massive.jsp].

The co-IP experiments were performed similarly, except for the following changes: A total of 4 mg of protein was incubated with 27 µL of anti-V5 agrose beads in a total volume of 400 µL and washed twice with 950 µL of NP-40 native IP buffer before washing with buffer 2 and buffer 3. After the last wash, the wash buffer
was aspirated until 100 µL was left, and 50 µL of 3x SDS sample buffer (187.5 mM Tris at pH 6.8, 6% mercaptoethanol, 30% glycerol, 9% SDS, 0.05% bromophenol blue) was added to the beads before boiling for 5 min. After brief centrifugation, the eluted proteins were separated by SDS-PAGE as described in the immunoblotting section. A detailed protocol for the IP/TMT mass spectrometry is provided in Supplemental Table S3.

**Purification of the Ndc80 complex**

Native Ndc80 complex was purified from asynchronously growing *S. cerevisiae* cells (UB16284 and UB19957) through anti-Flag immunoprecipitation of Spc24-6His-3Flag as described in Miller et al. (2016). Cells were grown in YPD. Protein lysates were prepared by lysing cells in a freezer/Mill [SPEX SamplePrep] submerged in liquid nitrogen (Sarangapani et al. 2014). Lysed cells were resuspended in buffer H [BH] (25 mM HEPES at pH 8.0, 2 mM MgCl₂, 0.1 mM EDTA, 0.5 mM EGTA, 0.1% NP-40, 15% glycerol, 750 mM KCl), supplemented with the protease inhibitors [20 µg/mL leupeptin, 20 µg/mL pepstatin A, 20 µg/mL chymostatin, 200 µM phenylmethylsulfonyl fluoride) and the phosphatase inhibitors [0.1 mM Na-orthovanadate, 0.2 µM microcystin, 2 mM β-glycerophosphate, 1 mM Na pyrophosphate, 5 mM NaF]. The resuspended cell lysates were ultracentrifuged at 98,500g for 90 min at 4°C. Dynabeads conjugated with anti-Flag antibody (Sigma) were incubated with extract for 3 h at 4°C with constant rotation, followed by three washes with BH containing the protease inhibitors, phosphatase inhibitors, 2 mM DTT, and 1 M KCl. The beads were further washed twice with BH containing 150 mM KCl and the protease inhibitors. Associated proteins were eluted from the beads by gently agitating the beads in elution buffer (0.5 mg/mL 3Flag peptide in BH with 150 mM KCl and the protease inhibitors) for 30 min at room temperature.

**In vitro kinase assay**

The kinase assay was performed using a recombinant variant of Ipl1 [AurB⁺] that was created by fusing the C-activation box of Slits to Ipl1 [a kind gift from S. Biggins's laboratory, the Fred Hutchinson Cancer Research Center]. The kinase-dead (KD) mutant of AurB⁺ contains the K133R mutation. These two constructs were first reported in de Regt et al. (2018). To begin the reaction, in a volume of 6 µL, 6.63 ng of the purified WT-Ndc80 or Ndc80Δ(2-28) was incubated with 1 µM AurB⁺ or KD in BH0.15 [same ingredients as BH in the previous section except with 150 mM KCl] for 4 min at room temperature. Next, 6 µL of hot ATP mix [1:1 µL of 100 mM ATP, 3 µL of 6000 Ci/mmol 10 mCi/mL γ-32P ATP, 275 µL of BH0.15] was added to the kinase/substrate mix. After 1 or 5 min of incubation at 25°C [see the figure legend for details], reactions were stopped by adding 6 µL of 3x SDS sample buffer, and boiling for 5–10 min at 95°C. Proteins were separated by SDS-PAGE as described in “Immunoblotting.” The gel was fixed and silver stained based on the manufacturer’s instructions [Thermo Scientific]. After the staining was stopped by incubating the gel in 5% acetic acid for 10 min, the gel was dried by vacuum for 1 h at 80°C. The dried gel was exposed to a phosphor storage screen for over 24 h. Screens were imaged with a Typhoon Scanner [GE Healthcare].

**Immunoblotting**

To extract proteins from cells, approximately four OD₆₀₀ units of cells were treated with 5% trichloroacetic acid for at least 15 min at 4°C. After an acetone wash, the cell pellet was air dried, lysed with acid-washed glass beads [Sigma] in 100 µL of lysis buffer [50 mM Tris at pH 7.5, 1 mM EDTA, 2.75 mM DTT, cComplete EDTA Free protease inhibitor cocktail [Roche] using a Mini-Beadbeater-96 [Biospec Products]. Next, 3x SDS sample buffer [187.5 mM Tris at pH 6.8, 6% 2-mercaptoethanol, 30% glycerol, 9% SDS, 0.05% bromophenol blue] was added and the lysate was boiled for 5 min. Proteins were separated by SDS-PAGE using 4%–12% Bis-Tris Bolt gels [Thermo Fisher] and transferred onto 0.45-µm nitrocellulose membranes [Bio-Rad] using a semidy transfer apparatus [Trans-Blot Turbo Transfer System, Bio-Rad]. The membranes were blocked for >30 min with Odyssey blocking buffer [PBS] [LI-COR Biosciences] and incubated in primary antibodies overnight at 4°C. Membranes were washed with PBST (phosphate-buffered saline with 0.01% Tween-20) and incubated with an antismouse secondary antibody conjugated to IRDye 800CW at a 1:15,000 dilution [LI-COR Biosciences 926-32212] and/or an antirabbit antibody conjugated to IRDye 680RD at a 1:15,000 dilution [LI-COR Biosciences 926-68071] for >1 h at room temperature. Immunoblots were imaged and quantified using the Odyssey system [LI-COR Biosciences]. Antibodies used in this study were mouse anti-V5 antibody [Thermo Fisher R960-25] used at a 1:2,000 dilution, rabbit anti-hexokinase antibody [US Biological H2035] used at a 1:20,000 dilution, rabbit anti-Sis1 antibody [kind gift from E. Craig] used at a 1:5,000 dilution, and a rabbit anti-H4 serine 10 phosphorylation antibody [Sigma 06-570] used at 1:1,000 dilution. The levels of each protein of interest were typically normalized to Hxk2, except for in Figure 6F. For the quantification in Figure 6F, Ndc80 levels were not normalized to that of Hxk2 because Hxk2 levels declined after cycloheximide treatment for the amaΔ strain. To reflect the experimental variations, we performed five independent biological replicates and graphed the mean and the standard errors of the mean for all five replicates.

**Spot growth assay**

Cells were grown on YPG [2% glycerol + YEP] plates overnight, resuspended in milliQ water, and then diluted to an OD₆₀₀ of 0.1 or 0.2. Fivefold serial dilutions were performed. Cells were spotted onto YPD plates or YPD plates supplemented with 15 µg/mL benomyl [benomyl plates]. The cells on the YPD plates were incubated for 1–2 d at 30°C, and those on benomyl plates were incubated for 2 d at 23°C.

**Reproducibility**

All of the experiments in this study were performed at least twice independently, and one representative result was shown for all Western blots. All of the RT-qPCR experiments were performed three times independently, with the mean and standard errors of the mean graphed.

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**Author contributions** J.C. conceived the study, performed the data collection, formal analysis, investigation, methodology,
validation, and visualization; and wrote the draft of the manuscript. A.L., E.N.P., and H.L. performed the investigation, formal analysis, validation, and visualization and edited the manuscript. L.A.K. performed the investigation, data curation, methodology, and formal analysis for the denaturing IP and edited the manuscript. R.E. provided key resources (purifying the Ndc80 complex) and edited the manuscript. R.M.H. performed the investigation for the truncation analysis of Ndc80 and edited the manuscript. J.K.K. performed the investigation and formal analysis for quantitative mass spectrometry. M.J. supervised the study and was responsible for the project administration; and wrote the draft of the manuscript. Akiyoshi B, Nelson CR, Ranish JA, Biggins S. 2009. Analysis of the multiple roles of cohesin in Saccharomyces cerevisiae. Mol Biol Cell 20: 1243–1253. doi:10.1093/mcb/jcp1286.8

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