A role for the Smc3 hinge domain in the maintenance of sister chromatid cohesion

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ABSTRACT Cohesin is a conserved protein complex required for sister chromatid cohesion, chromosome condensation, DNA damage repair, and regulation of transcription. Although cohesin functions to tether DNA duplexes, the contribution of its individual domains to this activity remains poorly understood. We interrogated the Smc3p subunit of cohesin by random insertion mutagenesis. Analysis of a mutant in the Smc3p hinge revealed an unexpected role for this domain in cohesion maintenance and condensation. Further investigation revealed that the Smc3p hinge functions at a step following cohesin's stable binding to chromosomes and independently of Smc3p's regulation by the Eco1p acetyltransferase. Hinge mutant phenotypes resemble loss of Pds5p, which binds opposite the hinge near Smc3p's head domain. We propose that a specific conformation of the Smc3p hinge and Pds5p cooperate to promote cohesion maintenance and condensation.

INTRODUCTION
Cohesin is a conserved protein complex required for sister chromatid cohesion, chromosome condensation, DNA damage repair, and regulation of transcription (Onn et al., 2008). To accomplish these functions, chromosome-bound cohesin tethers two distinct DNA duplexes or two sites on a single DNA duplex. A remarkable feature of cohesin-mediated tethers is that they must persist for long periods. For example, once generated, cohesion between sister chromatids must be maintained for up to several hours until cells progress through mitosis. Cohesion maintenance is essential for a successful mitosis since it ensures bipolar attachment and proper segregation of chromosomes. This process is crucial in mammalian oocytes since cohesion must be maintained from its establishment during meiotic prophase I, which occurs during fetal development, until the egg is fertilized in adulthood. Failure to maintain this cohesion can lead to aneuploidy and may cause infertility and birth defects in humans (Duncan et al., 2012). However, despite its critical function, the mechanism and regulation of cohesion maintenance remains poorly understood.

Cohesin is a large multisubunit complex with an elaborate molecular architecture. In the budding yeast Saccharomyces cerevisiae, core cohesin subunits are Smc1p, Smc3p, Mcd1p (also called Scc1p), and Scc3p (Onn et al., 2008). The Structural Maintenance of Chromosome (Smc) proteins fold back on themselves to form large dumbbell-shaped structures with two globular domains, referred to as the head and hinge, separated by an ∼45-nm-long coiled coil (Onn et al., 2008). Cohesin or purified Smc1p-Smc3p heterodimers have been visualized by electron microscopy, atomic-force microscopy, and scanning-force microscopy (Haering et al., 2002; Sakai et al., 2003; Kulemzina et al., 2016). These studies revealed that Smc1p and Smc3p dimerize by an interaction between their heads and a separate interaction between their hinges. Dimerization of the heads is further stabilized by the kleisin subunit Mcd1p which binds through its N-terminus to Smc3p and its C-terminus to Smc1p (Haering et al., 2002). The existence of two dimerization interfaces allows cohesin to form large rings. This ring structure likely explains cohesin’s ability to bind DNA by topological entrapment. In addition to these ring structures, more complex conformations have also been observed (Sakai et al., 2003). Evidence supporting the biological significance of these other conformations has been lacking.

Sister chromatid cohesion is established in S phase and then maintained until anaphase onset. Cohesion establishment is a multistep process. In budding yeast, the Scc2p/Scc4p complex...
(Ciosk et al., 2000) loads cohesin onto DNA at centromeres and along chromosome arms at cohesin-associated regions (CARs) in early S phase (Megee et al., 1999; Laloraya et al., 2000; Glynn et al., 2004). During S phase, DNA-bound cohesin is converted into a form that tethers sister chromatids by the Eco1p acetyltransferase, which acetylates Smc3p at lysines 112 and 113 (Skibbens et al., 1999; Toth et al., 1999; Rolef Ben-Shahar et al., 2008; Unal et al., 2008; Zhang et al., 2008). Once cohesion is established in S phase, the cohesin-associated regulator Pds5p is required to maintain cohesion until anaphase onset (Hartman et al., 2000; Panizza et al., 2000; Stead et al., 2003).

The mechanism of cohesion maintenance is only partially understood. Pds5p colocalizes with cohesin on chromosomes and, when mutated, causes a decrease in cohesin binding to chromosomes, a reduction in cellular Mcd1p levels, and a cohesion maintenance defect (Hartman et al., 2000; Panizza et al., 2000). This maintenance defect can be suppressed by preventing premature Mcd1p degradation via a polySUMO-dependent pathway or preserving Smc3p acetylation by deleting the HOS1 deacetylase (Stead et al., 2003; Chan et al., 2013; D’Ambrosio and Lavoie, 2014). Thus, Pds5p may function to protect cohesin complex from factors that could dissolve cohesin. However, cohesion maintenance is a more complex process. The cohesin mutant Mcd1-ROCC is defective for cohesion maintenance yet Mcd1p levels are not reduced and Pds5p recruitment to cohesin and chromosomes is unaffected (Eng et al., 2014). These observations suggest that an additional step beyond Mcd1p stabilization or Pds5p recruitment is required for cohesion maintenance.

A clue for this additional step comes from imaging and biochemical studies of cohesin and Pds5p. Biochemical studies indicate Pds5p binds to Mcd1p, placing Pds5p adjacent to the Smc head domains (Chan et al., 2013; Lee et al., 2016; Muir et al., 2016; Ouyang et al., 2016). The functional significance of this interaction is supported by mutations in budding yeast Mcd1p that mimic the cohesion maintenance defects upon Pds5p depletion (Eng et al., 2014). However, cross-linking has shown human Pds5Bp interacts with all cohesin subunits, implying that its association with cohesin is more extensive and/or dynamic (Huis in t Veld et al., 2014). This proximity was supported by the observation that purified hinge domains are capable of binding to each other and forming structures that are similar to those formed by cohesin (Stead et al., 2003; Kurze et al., 2006; Ünal et al., 2006). Our screen identified three RIDs that mapped to the North hinge interface and six mapped near the South interface. Of the six RIDs near the South interface, five were located at or immediately adjacent to conserved glycine amino acids known to be necessary for SMG hinge dimerization in vitro (Figure 1B) (Hirano et al., 2007). The sixth RID, encoding an insertion of five amino acids (AAAAA) following D667, maps to a hairpin loop extending from the top of a beta-sheet that contributes to the South hinge interface. We hypothesized that the unusual position of the D667 RID might reveal a novel function of the hinge in cohesin function.

The RID screen utilizes overexpression to generate a dominant phenotype. We wanted to determine whether smc3-D667 could support viability when expressed at native levels. For this purpose, we transformed a haploid strain bearing SMC3-3V5-AID as the sole SMC3, henceforth abbreviated SMC3-AID, with either an integrating smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus. We then compared growth of the SMC3-AID parent alone to derivatives containing either smc3-D667 or SMC3 wild-type allele under native expression at the LEU2 locus.

### RESULTS

The D667 region of the Smc3p hinge enhances but is not essential for cohesin binding at centromeres and cohesin-associated regions

We used a random insertion dominant (RID) screen to identify partial loss of function alleles of SMC3 (Milutinovich et al., 2007; Eng et al., 2014, Orgil et al., 2015). We expected to obtain RID screen mutations at the interfaces between Smc3p and Smc1p or Mcd1p. These mutations would be expected to prevent assembly and subsequent loading of cohesin onto chromosomes. In addition to assembly mutants, we predicted that mutations that preserved cohesin assembly would be found. We reasoned that if Smc3p function is modulated after cohesin assemblies and binds chromosomes to maintain cohesion, then mutants of Smc3p could be found that impair this step.

Mutant SMC3 alleles were generated by in vitro transposon-mediated mutagenesis, which produced a library encoding random five-amino-acid insertions (Supplemental Figure 1; see Materials and Methods). In this library, SMC3 was placed under control of the conditional pGAL1 promoter. The library was transformed into both wild-type haploid yeast and the temperature-sensitive smc3-42 strain. Transformants were obtained on dextrose-containing media to repress RID library pGAL1-SMC3 expression. Colonies were then screened for impaired growth on plates containing galactose as the carbon source to drive pGAL1-mediated overexpression of mutant SMC3 alleles. The location of insertions within SMC3 that impaired growth of wild-type (Supplemental Table 1) or smc3-42 cells (Supplemental Table 2) when overexpressed were then determined by sequencing.

In the course of mapping RID mutations, we found 10 RIDs within the Smc3p hinge domain (Figure 1A). Nine of these RIDs were located near interfaces with Smc1p. Dimerization of the Smc1p and Smc3p hinges forms a toroidal structure with two interfaces termed “North” and “South” (Mishra et al., 2010). Mutations that disrupt the hinge interfaces or that neutralize the positively charged amino acids in the central channel have been studied previously (Kurze et al., 2011). Our screen identified three RIDs that mapped to the North hinge interface and six mapped near the South interface. Of the six RIDs near the South interface, five were located at or immediately adjacent to conserved glycine amino acids known to be necessary for SMG hinge dimerization in vitro (Figure 1B) (Hirano et al., 2001). The sixth RID, encoding an insertion of five amino acids (AAAAA) following D667, maps to a hairpin loop extending from the top of a beta-sheet that contributes to the South hinge interface. We hypothesized that the unusual position of the D667 RID might reveal a novel function of the hinge in cohesin function.

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media containing auxin. The fact that smc3-D667p SMCS-AID cells grew well in the absence of auxin indicated that smc3-D667p is recessive unless overexpressed. The smc3-D667p mutant may fail to support viability on auxin because it disrupts the sequence of charged amino acids that follow D667. Therefore, we assessed the viability of two SMC3 alleles in which D667 and nearby charged amino acids had been mutated to alanine. Unlike the smc3-D667 RID mutant, the smc3-D667A single mutant and the smc3-D667A,K668A,R669A triple mutant supported growth of SMC3-AID cells on auxin (Supplemental Figure 1B). Therefore, the smc3-D667 RID likely disrupted SMC3p function independent of its effect on nearby charged amino acids. In summary, the smc3-D667 RID allele was unable to support one or more essential cohesin functions.

The inviability of smc3-D667 cells could be due to a failure of cohesin to bind DNA or a failure to perform an essential cohesin function after binding DNA. To distinguish between these possibilities, we first assessed whether smc3-D667p cohesin binds DNA. Strains containing SMC3-AID alone or also a second SMC3, either wild-type SMC3 or smc3-D667, were arrested in G1 phase and treated with auxin to deplete SMC3-AIDp. Cells were then synchronously released from G1 into YPD media containing auxin and nocodazole to rearrest them in mid-M phase while maintaining SMC3-AIDp depletion (Figure 2A and Materials and Methods). To assess qualitatively whether smc3-D667p supported binding of cohesin to chromosomes, we processed mid-M phase–arrested cells for chromosome spreads and assessed chromosomal binding of the cohesin subunit Mcd1p by immunofluorescence. Mcd1p is a marker for the cohesin complex, since Mcd1p cannot bind chromosomes unless it is part of the four-subunit complex (Toth et al., 1999). As expected, robust Mcd1p signal was observed on chromosome spreads from cells with SMC3p (SMC3 SMC3-AID) but not from cells without it (SMC3-AID) (Figure 2B). In smc3-D667 SMC3-AID cells, Mcd1p bound to chromosomes at levels similar to wild-type cells. This result indicated that smc3-D667p supports both cohesin complex assembly and binding to chromosomes.

We used chromatin immunoprecipitation (ChIP) to assess whether the cohesin chromosomal binding observed via spreads reflected specific binding to CARs and centromeres. Mid-M phase cells prepared as described for chromosome spreads (Figure 2A) were fixed and processed for ChIP (Figure 2A and Materials and Methods). Cohesin binding was assessed using anti-Mcd1p antibodies. As expected, Mcd1p binding to CARs was robust in cells with SMC3p (SMC3 SMC3-AID) and absent in those without it (SMC3-AID) (Figure 2C). Mcd1p binding in smc3-D667 cells was similar to wild type at centromeres (Figure 2C, right) and at the pericentromeric CARC1 peak (Figure 2C, left) but somewhat reduced at centromere-distal TRM1 and CAR1 peaks (Figure 2C, center, and Supplemental Figure 2A). These results indicated that smc3-D667p cohesin localizes to CARs and centromeres. To corroborate further the DNA binding of smc3-D667p, we generated strains bearing SMC3p and smc3-D667p
using anti-HA to directly monitor the Smc3p cohesin subunit. As was observed in the Mcd1p ChIP, smc3-6HA-D667p (SMC3-AID) bound to CARs and centromeres, albeit at levels 50% reduced relative to wild-type Smc3p (Figure 2D and Supplemental Figure 3).

FIGURE 2: Cohesin containing smc3-D667p binds to chromosomes in mid-M phase–arrested cells. (A) Regimen used to prepare cells synchronously arrested in mid–M phase. Cultures were grown to mid–log phase at 23°C and treated with alpha factor for 3 h to arrest cells in G1 phase and then auxin was added and cells were incubated an additional hour in G1 to deplete Smc3-3V5-AIDp. Cells were synchronously released from G1 arrest into YPD media containing auxin and nocodazole to re-arrest in mid–M phase (Materials and Methods). (B) Chromosome spreads showing that smc3-D667p cohesin binds chromosomes at levels similar to wild type. Haploid SMC3 SMC3-AID (BRY474), SMC3-AID (VG3651-3D), and smc3-D667 SMC3-AID (BRY482) cells were grown as described in A. Aliquots of mid–M phase–arrested cells were fixed and processed for chromosome spreads. Bulk chromosomal DNA (DAPI) and cohesin binding (α–Mcd1) are shown. (C, D) ChIP showing that smc3-D667 cohesin binds to CARs and centromeres. (C) Haploid strains in B were arrested in mid–M phase as described in A and then fixed and processed for ChIP as described under Materials and Methods. ChIP of Mcd1p binding at CARC1 (left) and TRM1 (middle) and at two centromeres (right). Wild-type strain SMC3 (dotted lines and white bars), smc3-D667 strain (black lines and black bars), and SMC3-AID alone (gray lines and gray bars). (D) ChIP of HA epitope tagged Smc3p and smc3-D667p at CARC1 (left) and TRM1 (middle) and at two centromeres (right). Haploid strains SMC3-6HA SMC3-AID (BRY604; dotted lines and white bars), smc3-6HA-D667 SMC3-AID (BRY602; black lines and black bars) and SMC3-AID only (VG3651-3D; gray lines and gray bars) were arrested and processed for ChIP as described in C.

tagged with a 6HA epitope in the SMC3-AID background. Mid–M phase auxin-treated cells were prepared (Figure 2A) and the presence of smc3-6HA-D667 and Smc3-6HAp were confirmed by Western blotting (Supplemental Figure 3). We then performed ChIP using anti-HA to directly monitor the Smc3p cohesin subunit. As was observed in the Mcd1p ChIP, smc3-6HA-D667p (SMC3-6HA-D667 SMC3-AID) bound to CARs and centromeres, albeit at levels 50% reduced relative to wild-type Smc3p (Figure 2D and Supplemental
Figure 2B). These data, using two different cohesin subunits, show that smc3-D667p cohesin complex binds to CARs and centromeres at ~50% the levels of wild type.

The D667 region of the Smc3p hinge is required to maintain cohesion

Smc3-D667p cohesin binds chromosomes, so we assayed whether it can perform cohesin's function of tethering sister chromatids. Therefore, we assessed sister chromatid cohesion at centromere-proximal (TRP1) or centromere-distal (LYS4) loci by integrating tandem LacO repeats in strains that express a GFP-LacI fusion (Figure 3A and Materials and Methods). Strains bearing SMC3-AID alone or also containing either wild-type SMC3 or smc3-D667 were arrested in G1, treated with auxin to degrade Smc3-AIDp, and then synchronously released from G1 into media containing auxin and nocodazole to allow progression through S phase and arrest in mid–M phase (Figure 2A). Nearly all G1 cells in all strains contained a single GFP focus, indicating no preexisting aneuploidy (Figure 3B). As expected, only a small fraction of mid–M phase–arrested cells with Smc3p (SMC3 SMC3-AID) lost cohesion at TRP1 or LYS4, whereas cells lacking Smc3p (SMC3-AID) had almost complete loss of cohesin. Nearly two-thirds of cells expressing only smc3-D667 (smc3-D667 SMC3-AID) also had lost cohesion at these two loci. This result suggested that the D667 region of the hinge was required for either robust establishment and/or maintenance of cohesin.

These two possibilities can be distinguished by kinetic analysis of cohesion in populations of cells synchronously progressing through the cell cycle. Mutants that compromise cohesion establishment like those defective in core subunits of cohesin MCD1, SMC3, and SMC1 exhibit sister chromatid separation immediately after DNA replication (Guacci et al., 1997; Michaelis et al., 1997). Mutants that compromise cohesion maintenance like those defective in the cohesin regulator PD55 also lose cohesion but significantly later in the cell cycle than establishment mutants (Tanaka et al., 2001; Stead et al., 2003; Noble et al., 2006; Eng et al., 2014). Using the same strains as described above along with a PD55-AID strain, we assessed when cohesion was lost in smc3-D667. Strains were arrested in G1 and treated with auxin to degrade Smc3-AIDp and then released from G1 in the presence of auxin and nocodazole to allow cells to progress through S phase and arrest in mid–M. After release from G1, aliquots of cells were removed every 15 min to assess DNA content and cohesion at TRP1 and LYS4 (Figure 3C).

From analysis of the DNA content, all strains exhibited nearly identical kinetics of progression through S phase and subsequent arrest in mid–M (Supplemental Figure 4A). As expected for cells expressing Smc3p (SMC3 SMC3-AID), sister chromatids were paired through mid–M arrest, so few cells with separated sisters were detected. In contrast, both strains lacking Smc3p (SMC3-AID) and Pds5p (PD55-AID) lost cohesion. However, the cohesion loss in the PD55-AID cells was delayed by ~20 min, as published previously (Eng et al., 2014). Cells expressing only smc3-D667p (smc3-D667 SMC3-AID) resembled PD55-AID cells, with delayed cohesion loss at the LYS4 locus and a more pronounced delay in cohesion loss at the TRP1 locus. This delay in cohesion loss in cells with smc3-D667p demonstrated that smc3-D667 cells, like Pds5p-deficient cells, could establish but not maintain cohesion. Thus, the D667 region of the Smc3p hinge is important specifically for efficient maintenance of cohesion at both CEN-proximal and CEN-distal loci.

Cohesin is required to recruit the maintenance factor Pds5p to chromosomes (Hartman et al., 2000; Panizza et al., 2000). Since cells expressing smc3-D667p displayed a cohesion maintenance defect identical to cells depleted of Pds5p, we tested whether smc3-D667p cohesin was able to recruit Pds5p to chromosomes. To address this possibility, we first analyzed whether smc3-D667p supported Pds5p binding to chromosomes by ChIP using a Pds5p antibody (Figure 3D and Supplemental Figure 4B). The ratio of Pds5p bound to CARs and centromeres in cells with smc3-D667p (smc3-D667 SMC3-AID) to Smc3p was very similar to that seen for Mcd1p or smc3-6HA-D667p. These results indicate that cohesin with smc3-D667p can bind Pds5p and recruit it to chromosomes. The ability of Pds5p to bind cohesin with smc3-D667p was then tested by coimmunoprecipitation (Figure 3E). Cells expressing FLAG-tagged Scc3p and HA-tagged Smc3p or smc3-D667p were arrested in M phase after auxin-mediated depletion of Smc3-AIDp. Scc3p was immunoprecipitated using anti-FLAG antibody and cohesin subunits detected in the precipitates by Western blot. As expected, no Pds5p was detected in the FLAG immunoprecipitate from cells lacking Smc3p or when Scc3p was untagged (first and second lanes), while Pds5p and Smc3-6HAp were detected in the immunoprecipitate from cells expressing Smc3-6HAp (third lane). Importantly, similar Pds5p levels were observed in the immunoprecipitate from cells expressing smc3-D667-6HAp (fourth lane). Thus, smc3-D667p cohesin binds Pds5p and recruits it to DNA.

The D667 region of the Smc3p hinge is not required for its stable binding to chromosomes

Cohesin is known to convert from a DNA-bound, untethered state to a tethered state in S phase (Rolf Ben-Shahar et al., 2008; Ūnäl et al., 2008). We envisioned two models by which cohesion that had been established in S phase by smc3-D667p could fail to be maintained as cells progressed into M phase. In one model, cohesin reverts back to its untethered state without perturbing cohesin binding to DNA. Precedence for this phenotype comes from the cohesin mutant mcd1-ROCC which, like smc3-D667, is defective for cohesion maintenance (Eng et al., 2014). Alternatively, the smc3-D667p is less stably bound so dissociates from DNA. In this model, following cohesion establishment, cohesin dissociation from chromosomes could manifest as a cohesion maintenance defect. Detecting putative cohesin dissociation is difficult, because the Scc2p/Scc4p complex continues loading cohesin onto chromosomes in mid–M phase creating a pool of bound cohesin that does not contribute to cohesion (Lengronne et al., 2006). Therefore, the Scc2p/Scc4p complex must be inactivated to allow detection of cohesin dissociation.

To distinguish between these two models, we examined the stability of smc3-6HA-D667p binding to DNA under conditions where additional loading was prevented by depletion of the cohesin loader subunit Scc2p. This loader depletion approach revealed that in wild-type cells, cohesin (Mcd1p) binds stably to CARs but exhibits reduced stability at centromeres (Eng et al., 2014). Therefore, we replaced SCC2 with SCC2-3FLAG-AID in SMC3-AID strains bearing either wild-type Smc3-6HAp or smc3-D667-6HAp. Cultures of these strains were grown to mid–log phase and arrested in mid–M phase by addition of nocodazole. Cultures were then split and either auxin or vehicle (DMSO) was added, and then they were incubated for 1 h. The aliquot containing auxin will deplete both Scc2-3FLAG-AIDp and Smc3-3V5-AIDp. Samples were collected and either fixed for ChIP or processed for Western blot analysis (Figure 4A). Depletion of Scc2-3FLAG-AIDp and Smc3-3V5-AIDp was confirmed by Western blot (Figure 4B).

ChIP of Smc3-6HAp showed no difference in binding to CAR peaks TRM1 and CAR1 after Scc2-3FLAG-AIDp depletion (Figure 4C, left). The persistence of high ChIP levels even after 1 h indicated that cohesin remained very stably bound to DNA. Similarly, smc3-6HA-D667p ChIP at TRM1 and CAR1 peaks was unchanged by
FIGURE 3: The smc3-D667 mutant exhibits a cohesion maintenance defect. (A) Schematic of cohesion loss assay using loci tagged with GFP-LacI. After replication, cells with cohesion have a single GFP focus, whereas cells where cohesion is lost have two GFP foci. (B) Cohesion loss at CEN-proximal TRP1 and CEN-distal LYS4 loci in mid–M phase–arrested cells. Haploid strains were arrested in G1, depleted of Smc3p-AID, and then synchronously released from G1 and rearrested in mid–M phase under depletion conditions as described in Figure 2A. LacO arrays integrated at TRP1 (left) in haploid SMC3-AID yeast alone (BRY676) or also containing wild-type SMC3 (BRY678) or smc3-D667 (BRY680). LacO arrays integrated at LYS4 (right) in SMC3-AID yeast alone (VG3651-3D) or containing wild-type SMC3 (BRY474) or smc3-D667 (BRY482). Samples were collected from G1-arrested auxin-treated cells and mid–M phase–arrested cells and scored for cohesion. The percentage of cells with two GFP foci (sister separation) were averaged from two independent experiments and plotted. Cells (100–200) were scored per sample at each time point. Error bars represent SD. *p < 0.05 when tested against SMC3 SMC3-AID % sister separation in mid–M (t test, two-tailed). (C) Time course to assess the kinetics of cohesion loss. Haploid strains were arrested in G1, treated with auxin, and synchronously released into mid–M phase arrest in auxin containing media as described in Figure 2A. Samples were collected in G1 and every
Scc2-3FLAG-AIDp depletion (Figure 4C, right). At centromeres, Smc3-6HAp shows somewhat reduced binding after Scc2-3FLAG-AIDp depletion, confirming that this cohesion is less stably bound. Similarly, somewhat reduced binding of sccm3-6HA-D667p to centromeres was observed. These results demonstrated that smc3-6HA-D667p was as stably bound to chromosomes as wild-type Smc3-6HAp. Importantly, our results indicated that in mid–M phase–arrested smc3-D667 cells, when most sister chromatid cohesion is lost (Figure 3), smc3-D667p cohesion is stably bound to chromosomes. Thus, the D667 region of the Smc3p hinge performs a function in maintaining cohesion other than ensuring stable binding to DNA.

The D667 region of the Smc3p hinge modulates cohesion and supports viability by a mechanism independent of Eco1p-dependent acetylation

Eco1p is necessary for establishing cohesion during S phase through its acetylation of Smc3p at lysines K112 and K113. Although cohesion establishment occurs during S phase, Smc3p acetylation remains until anaphase onset, suggesting it may be required to maintain cohesion (Beckouet et al., 2010). Since smc3-D667p supported cohesion establishment, we predicted that it would be acetylated by Eco1p. Therefore, we used an antibody that specifically recognizes acetylated Smc3p-K113 to test the acetylation of smc3-D667p in cells arrested in mid–M. Cells were arrested in mid–M after auxin depletion (Figure 5A). As expected, in cells depleted of Eco1-AIDp or Smc3-AIDp, no acetylated Smc3p was detected (Figure 5B). While wild-type Smc3p showed a strong acetylation signal, acetylation signal for smc3-D667p was reduced. A reduction in acetylation signal was expected because cohesin was known to be acetylated only after binding to DNA, and less cohesin with smc3-D667p was bound to DNA than wild-type cohesin (Figure 2). Direct comparison of acetylation levels is possible when signal from the acetylation-recognizing antibody is linear across the observed range. However, we found that signal from the acetylation antibody was nonlinear (Supplemental Figure S5), making it possible that smc3-D667p acetylation levels were closer to Smc3p than Figure 5B suggested.

To assess whether the reduced amount of smc3-D667p acetylation was responsible for the cohesion maintenance defect, we first asked whether a change in acetylation levels correlated with the appearance of the cohesion defect. Reduced smc3-D667p acetylation may have resulted from a failure to acetylate it in S phase or to maintain it after S phase. To distinguish between these possibilities, we immunoprecipitated smc3-6HA-D667p from cells progressing synchronously through S phase following release from G1 arrest (Figure 5C). As expected, wild-type Smc3-6HA acetylation began to appear during S phase and then increased and remained high through M phase arrest (Figure 5D). While acetylation of smc3-6HA-D667p was lower than WT in early S phase, it increased as cells progressed into M phase. Therefore, smc3-D667 cells establish cohesion with low smc3-D667p acetylation levels, but its failure to maintain cohesion is not due to a subsequent decrease in acetylation levels.

We further examined the correlation between Smc3p acetylation levels and cohesion function by asking whether low levels of Smc3p acetylation always led to loss of essential cohesion function. Temperature-sensitive eco1 mutants (eco1-203 and eco1-1) establish and maintain cohesion at permissive temperature, yet eco1-1 has greatly reduced acetylation (Toth et al., 1999; Heidinger-Pauli et al., 2009; Rowland et al., 2009). We therefore compared Smc3p acetylation levels of the eco1-203 mutant grown at the permissive temperature 23°C to the smc3-D667 mutant. The level of Smc3p acetylation in eco1-203 cells was very similar to smc3-D667 cells (Figure 5E). This result suggested that the level of smc3-D667p acetylation was sufficient to support cohesion function. However, we could not rule out that the acetylation level of smc3-D667p was below a critical threshold too subtle to distinguish by Western blot.

We sought additional support for the idea that the lower smc3-D667p acetylation level is not responsible for its mutant phenotype. For this purpose, we assayed the smc3-D667 mutant in the SMC1-D1164E mutant background, as this SMC1 allele completely bypasses the need for Smc3p acetylation in both cohesion and viability (Çamdere et al., 2015; Elbatsh et al., 2016). In the presence of auxin, smc3-D667 SMC3-AID and SMC1-D1164E smc3-D667 SMC3-AID cells were inviable (Figure 6A). Therefore, the viability defect of smc3-D667 is distinct from eco1-ts and deletion mutants, which are bypassed by SMC1-D1164E. We next asked whether SMC1-D1164E restored cohesion to smc3-D667 cells as was observed for the eco1Δ wp1Δ mutant and eco1Δ cells (Çamdere et al., 2015). As expected, SMC1-D1164E restored cohesion at the LYS4 locus in the eco1Δ wp1Δ mutant (Figure 6B and Çamdere et al., 2015). However, SMC1-D1164E failed to restore cohesion to the smc3-D667 SMC3-AID mutant in the presence of auxin (Figure 6C). These results supported the idea that the viability and cohesion defects of smc3-D667 cells were independent of reduced levels of Smc3p acetylation.

The D667 region of the Smc3p hinge is required for rDNA condensation and viability even in the absence of antagonism by Wp1p

In addition to sister chromatid cohesion, cohesion and its regulators Pds5p and Eco1p are required for the proper mitotic condensation...
is located on the periphery of the primary chromosome mass. In interphase, the rDNA can be seen as a diffuse puff while in M phase it condenses into a loop (Guacci et al., 1994). Chromosome spreads of the SMC3-AID and PDS5-AID strains were prepared from cells of chromatids in budding yeast (Guacci et al., 1997; Skibbens et al., 1999; Hartman et al., 2000). We addressed whether smc3-D667 cells supported condensation by examining the morphology of the rDNA locus on chromosome XII. In chromosome spreads the rDNA locus on chromosome XII. In chromosome spreads the rDNA
arrested in mid–M phase (Figure 7A). The rDNA morphology was scored as 1) a tight, fully condensed loop; 2) a wide, decondensed loop; or 3) diffuse, with no apparent loop. In cells with wild-type Smc3p, the rDNA formed tight loops in almost all chromosome masses, indicative of chromosome condensation. In cells lacking Smc3p (SMC3-AID), the rDNA was almost always present as a diffuse mass, recapitulating the established role of Smc3p and cohesion in condensation. Cells expressing only smc3-D667p or depleted of Pds5p (PDS5-AID) exhibited very similar condensation defects and tight loops were rarely observed (Figure 7A). Thus, the D667 region of the Smc3p hinge is needed for two M-phase functions of cohesin, the maintenance of cohesion and condensation.

We next asked whether the condensation defect and inviability of smc3-D667 cells was due to antagonism by Wpl1p. Deletion of WPL1 restores viability to eco1 temperature-sensitive or eco1Δ strains that have impaired or absent acetylation (Rowland et al., 2009; Guacci and Koshland 2012). If the defect of smc3-D667 can be attributed to a loss of Eco1p function, then wpl1Δ would restore condensation and viability to smc3-D667 cells. To test this idea, we characterized the consequences of WPL1 deletion in the smc3-D667 strain. wpl1Δ failed to restore viability to smc3-D667 SMC3-AID cells on media containing auxin (Figure 7B). Consistent with smc3-D667 representing a defect distinct from cells lacking Smc3p acetylation, wpl1Δ failed to restore condensation of the rDNA or cohesion to smc3-D667 cells (Figure 7, C and D, respectively). Altogether, our observations confirmed that the critical defects in smc3-D667 cells were independent of Smc3p acetylation or antagonism by Wpl1p.

The D667 region is necessary for interallelic complementation

Interallelic complementation between alleles of SMC3 or MCD1 revealed the ability of two separate cohesin complexes to share activities to restore cohesin functions. Additional evidence suggests that this communication between cohesins might reflect direct cohesin–cohesin interaction on chromosomes (Eng et al., 2015). We wondered whether the D667 region of the hinge was needed for cohesin–cohesin communication. To test this idea, we asked whether smc3-D667 could partner with the temperature-sensitive smc3-42 allele to exhibit interallelic complementation. The temperature-sensitive smc3-42 strain cannot grow at its restrictive temperature of 34°C. Previously it had been shown that the smc3-K113R allele cannot support viability as the sole copy of SMC3. However, a strain in which both smc3-K113R and smc3-42 alleles are present exhibits robust growth at 34°C, a condition in which neither single mutant can grow (a summary of complementation relationships is provided in Figure 8B). With this knowledge, we asked whether smc3-D667 could substitute for smc3-K113R and complement smc3-42. As a metric for the extent of interallelic complementation, we repeated the previous experiment with smc3-42 and smc3-K113R. As expected, at 34°C neither smc3-42 nor smc3-K113R single mutants were viable, while the smc3-42 smc3-K113R double mutant showed robust growth similar to wild type (Figure 8A). As expected, the smc3-D667 single mutant failed to grow. The double smc3-42 smc3-D667 mutant resembled the growth of smc3-42 alone. Thus, the property of interallelic complementation observed between smc3-42 and smc3-K113R was not observed between smc3-42 and smc3-D667. Therefore, smc3-D667 lacks the activity necessary for interallelic complementation. This result suggested that the D667 region of the hinge is necessary for cohesin–cohesin communication.

DISCUSSION

Cohesin has a complex architecture with a heterodimeric ATPase domain and a hinge domain connected by a long coiled coil. The roles of these domains in cohesin’s activity on chromosomes is poorly understood. Here, we identified and characterized smc3-D667, a mutant in the Smc3p hinge domain that blocks cohesin function in M phase. Kinetic analyses of cohesion during the cell cycle reveal that this mutation allows cohesin establishment but impairs subsequent maintenance of cohesion. We also show that this mutation impairs mitotic chromosome condensation of the rDNA. However, this mutation does not perturb the stable association of cohesin with chromosomes as measured by the persistence of this association even after loader inactivation. Together, our results support a function of cohesin’s hinge domain in cohesion maintenance and condensation independent of cohesin’s stable binding to chromosomes.

The cohesion maintenance and condensation functions of the hinge domain revealed by smc3-D667 have not been reported previously. Two mutations that impact the North and South interfaces of the hinge dimer revealed a role of the hinge in cohesin binding to chromosomes, as expected given the role of the hinge dimer in maintaining the topological integrity of cohesin (Mishra et al., 2010). The novel phenotypes of smc3-D667 are consistent with D667 localization, determined by alignment to Smc3p homologues, within a loop not expected to impact the dimer interface. One study designed a cluster of mutations in SMC1 and SMC3 that neutralize the positive charges in a central channel formed by hinge dimerization (Kurze et al., 2011). This cluster of mutations (charge neutralization alleles) caused defects in cohesin and Smc3p acetylation but did not impair stable binding of cohesin to chromosomes, all phenotypes similar to the smc3-D667 allele. However, unlike our study of smc3-D667, the charge neutralization alleles were not analyzed for establishment and maintenance of cohesion, the functional significance of the reduced Smc3p acetylation, or condensation. If these alleles had the same cohesion and condensation defects as the smc3-D667 allele, as we predict, then these results would imply that changes to two distinct regions of the hinge dimer contribute to a common function needed for cohesion maintenance and condensation. The potential cooperation of the D667 region of the Smc3p hinge and the hinge channel could reflect a previously unrecognized conformational change of the hinge dimer needed for cohesin function. Indeed, in addition to the strict toroidal structures seen by crystalization of the cohesin or TmSMC hinge dimers, a recently published structure of the related GsSMC hinge dimer revealed that hinge dimers may adopt an asymmetric, relaxed conformation resembling a spring washer (Haering et al., 2002; Kurze et al., 2011; Kamada et al., 2017). Surprisingly, while both hinge interfaces remained intact in this structure, the relaxed face of the GsSMC hinge dimer involved a break in the beta-sheet connected by a loop homologous to the D667 loop of Smc3p. Together with our results, further investigation of hinge structural flexibility on conformations and functions of cohesin seem worthwhile.

The unusual phenotypes of smc3-D667 are also strikingly similar to those described for Pds5p depletion and mcd1 alleles (Chan et al., 2013, Eng et al., 2014). They all allow stable cohesin binding to DNA but cause defects in cohesion maintenance and condensation. The smc3-D667 mutant reduces cohesin chromosome binding by 50%. Previous work demonstrated that reduction of cellular Mcd1p reduced cohesin binding to CARs by 50%, yet cohesion was established and maintained at nearly wild-type levels (Heidinger-Pauli et al., 2010). This shows that merely reducing cohesin binding on chromosomes is not sufficient to produce a maintenance defect.
FIGURE 5: smc3-D667p has reduced acetylation at K113. (A) Regimen used to assess Smc3-K113 acetylation in mid–M phase–arrested cells. Early log phase cultures were treated with 0.75 mM auxin for 1 h to deplete Smc3-3V5-AIDp, and then nocodazole was added, and cultures were incubated 3 h to arrest cells in mid–M phase. (B) Reduced K113 acetylation of smc3-D667p. Haploid ECO1-AID (VG3633-2D), SMC3-AID (VG3651-3D), SMC3 SMC3-AID (BRY474), and smc3-D667 SMC3-AID (BRY482) cultures grown as described in A. Protein extracts were made and subjected to SDS–PAGE and then analyzed by Western blot. Antibodies against Smc3-K113 acetylation (Smc3-ac) are shown as short and long exposures; anti-Mcd1p antibodies (Mcd1p) serve as control for cohesin levels and antibodies against tubulin (Tub1) for a loading control. (C) Regimen used to determine the kinetics of Smc3-K113 acetylation establishment within a single cell cycle. Log phase cultures grown in YPD at 23°C were arrested in G1 using alpha factor, treated with auxin to deplete Smc3p-AID in G1, and then released into fresh YPD containing auxin and nocodazole to synchronously arrest cells in mid–M phase (Materials and Methods). (D) smc3-6HA-D667p has reduced acetylation in S phase but acetylation remains in mid–M phase. Haploid SMC3-AID cells expressing Smc3-6HAp (BRY604, left) or smc3-6HA-D667p (BRY602, right) were grown as described in C. Aliquots were taken at the indicated time points, and protein
Our observation that the Smc3p hinge also functions in cohesion maintenance suggests that it may cooperate alongside Pds5p and Mcd1p to promote a common molecular function. Indeed, this common function provides a biological explanation for in vivo FRET studies that suggest the formation of a complex involving the head, hinge, and Pds5p (Mc Intyre et al., 2007), and recent biochemical experiments that detected a supramolecular complex between the Schizosaccharomyces pombe hinge dimer and Psc3p (Sc3p orthologue) that binds to the head-associated Rad21p (Mcd1p orthologue) and Pds5p. Altogether these biochemical results along with our study support the idea that the hinge, Mcd1p, and Pds5p cooperate in a structural conformation required to promote cohesion maintenance and condensation.

Potential insight into the molecular function of this complex conformation comes from several additional observations. One possibility was the protection of Eco1p acetylation of Smc3p. Here, we show that while the level of smc3-D667 acetylation is lower than wild type, it is equal to that of the eco1-203 mutant at its permissive temperature, which supports both viability and sister chromatin cohesion. Furthermore, we show that SMC1-D1164E and wpl1Δ, two different mutations previously shown to bypass the absence of Eco1p acetylation in viability, cohesion (only smc1-D1164E) and condensation (only wpl1Δ) are unable to restore these functions to the smc3-D667 mutant. Finally, while Pds5p depletion also shows reduced Smc3p acetylation, the mcd1-

Therefore, the reduced chromosomal binding of cohesin in the smc3-D667 mutant is unlikely to be the major cause of its maintenance defect. Rather, an additional activity of cohesin must be impaired (see below). Interestingly, the mcd1-ROCC mutant displays a cohesion maintenance defect yet cohesin binds at wild-type levels (Eng et al., 2014). This result suggests that smc3-D667, mcd1-ROCC, and pds5 mutants may render cohesin sensitive to a cohesion antagonizing pathway active late in the cell cycle, but our results suggest that at least for smc3-D667, this putative pathway would have to be independent of Wpl1p function or Mcd3p-K112,K113 acetylation. Further investigation will be necessary to better understand the mechanistic underpinnings of cohesion maintenance.

extracts were made. A small portion was reserved for total protein, and then anti-HA antibody was added to immunoprecipitate SMC3-6HAp or smc3-6HA-D667p (Materials and Methods). Samples were subjected to SDS–PAGE and then analyzed by Western blot. Antibodies against SMC3-K113 acetylation (Smc3-ac) and both short and long exposures are shown for better comparison. Antibodies were used to monitor levels of the Smc3p (HA) and Mcd1p (Mcd1) cohesin subunits and anti-Tubulin antibodies (Tub1) used as a loading control. Samples were also collected to assess DNA content by flow cytometry (right side). (E) Similar levels of K113 acetylation in smc3-D667 and eco1-203 at permissive temperature. Haploid strains SMC3-AID (VG351-3D), SMC3 SMC3-AID (BRY474), smc3-D667 SMC3-AID (BRY833), and SMC3 SMC3-AID SMC1-D1164E (BRY833) were grown according the regimen in Figure 2A and processed to assess cohesion loss at the CEN-distal LYS4 locus as described in B. For both B and C, the percentage of cells with two GFP foci (sister separation) were derived from two independent experiments. An amount of 100–200 cells was scored per sample at each time point. Error bars represent SD. NS = not significant (p = 0.74; t test, two-tailed).
The smc3-D667 mutant is defective in condensation and cohesion even in the absence of cohesin antagonist Wpl1p. (A) Condensation of the rDNA locus in smc3-D667 cells. Percentage of chromosome masses displaying tight loop, wide loop, or diffuse masses; chi-squared. Data were collected from two independent experiments. (B) Cohesion loss in smc3-D667 wpl1Δ. Haploid strains SMC3-AID (BRY474), SMC3-AID (VG3651-3D), smc3-D667 SMC3-AID (BRY482), and PD55-AID (TE228) were grown and treated as in Figure 2A and then assessed as if for in situ hybridization (see Materials and Methods). Chromosome masses were scored for rDNA locus morphology after staining with DAPI. \( n = 200 \) for each genotype. * \( p < 0.05 \) (when compared with SMC3 SMC3-AID masses; chi-squared). Data were collected from two independent experiments. (B) wpl1Δ fails to restore viability to smc3-D667 cells. Haploid SMC3-AID strain derivatives with SMC3 (BRY474), SMC3 wpl1Δ (BRY716), smc3-D667 (BRY482), smc3-D667 wpl1Δ (BRY718), or SMC3-AID alone (VG3651-3D) were grown and plated as described in Figure 1C. (C) Quantification of condensed rDNA masses from mid-M phase–arrested cells. Haploid strains SMC3-AID (VG3651-3D), SMC3 SMC3-AID (BRY474), smc3-D667 SMC3-AID (BRY482), smc3-D667 SMC3-AID wpl1Δ (BRY718), and wpl1Δ (DK5561) were treated and processed as in A. The percentage of chromosome masses displaying a tight rDNA loop is shown. (D) Cohesion loss in smc3-D667 wpl1Δ cells. Haploid wpl1Δ (DK5561) and SMC3-AID strain derivatives with SMC3 (BRY474), SMC3-AID alone (VG3651-3D), smc3-D667 (BRY482), and smc3-D667 wpl1Δ (BRY718) were treated as in Figure 2A, and the percentages of separated sisters at the LYS4 locus were plotted. * \( p < 0.05 \) when tested against SMC3 SMC3-AID (t test, two-tailed). NS = not significant (\( p = 0.25 \)). Error bars represent the SD.

**FIGURE 7:**

MATERIALS AND METHODS

Random insertion screen of SMC3

Plasmid pBR25 containing pGAL-SMC3 URA3 ARS/CEN was subject to in vitro transposition according to the protocol recommended by the MuA transposase MGS Kit (ThermoFisher Cat. F701). After transforming into TOP10 cells (Thermo), 5756 AmpR KanR colonies were pooled, and plasmids were harvested by Midi Prep (Qiagen). The pooled library was digested with NotI to excise the KanR marker, gel extracted, and religated. Ligation products were transformed once again into TOP10 cells and confirmed to have lost KanR by replica plating. More than 30,000 colonies were pooled, and plasmids were harvested by Midi Prep to obtain a library of pGAL-SMC3 plasmids with 15 extra smc3-42p binding to DNA (Eng et al., 2015). Similar phenotypic and molecular interallelic complementation for mcd1 alleles was also observed (Eng et al., 2015). These observations led us to suggest that interallelic complementation of cohesin mutants reflected cohesin communication likely by the physical interaction between cohesin complexes. The importance of SMC complex oligomerization in their function is gaining traction. The inability of smc3-D667 to complement smc3-42 is consistent with the idea that the D667 region of the hinge is necessary for the physical interaction between cohesins, and this physical interaction is necessary for maintaining cohesion and condensation.

We propose a working model in which cohesin oligomerizes by forming inverted dimers such that the hinge of one cohesin binds to the head of the other cohesin possibly through binding to Scc3p and that this hinge-head interaction is stabilized by Pds5p. As suggested previously, we can imagine two ways in which hinge-dependent oligomerization might be critical for maintenance of tethering (Eng et al., 2015). We previously showed that mere binding of cohesin to DNA is insufficient to generate tethering, implying that tethering requires an additional activity (Eng et al., 2014). In one model (intramolecular handcuff), two DNA binding activities reside in the same cohesin. In this case, oligomerization may inhibit (possibly by physical occlusion) factors that destabilize one of these binding activities. In a second model (intermolecular handcuff) tethering is achieved directly by hinge-dependent oligomerization of two cohesins each of which has a single DNA binding activity. Resolving these models awaits direct biochemical assays for cohesin oligomerization.
nucleotides randomly inserted. Library depth was calculated by multiplying the fraction of pBR25 coding for SMC3 (3,693 base pairs of 10,083 base pairs total) by the number of AmpR KanR colonies (5756) to obtain 2118 plasmids expected to have an insertion in SMC3. From this calculation, we expect plasmids represented in the library harboring insertions every ∼1.7 base pairs along SMC3. The library was transformed into wild-type (3349-1B) and smc3-42 (3358-3B) strains, which were incubated at 23°C for 3 d to select for transformants on synthetic complete media lacking uracil (SC –URA) with 2% dextrose supplied as the carbon source. Wild-type colonies (3382) and smc-42 colonies (1811) were screened. Transformation colonies were replica plated onto SC –URA 2% galactose and SC –URA 2% dextrose plates as a control and incubated overnight at 23°C. Colonies that were slow growing or inviable on galactose plates were then grown overnight in liquid YPD and plated in 10-fold serial dilutions on 1) galactose plates to confirm slow growth and 2) 5-fluoroorotic acid (FOA) plates with 2% galactose to confirm linkage of slow growth to presence of the RID library plasmid. Insertion mutations were identified by PCR and sequencing across the entire SMC3 open reading frame (ORF).

**Yeast strains, media, and growth**

All strains used are in the A364A background, and their genotypes can be found in Table 1. Yeast extract/peptone/dextrose media and synthetic dropout media were prepared as previously described (Guacci et al., 1997). Conditional AID degron strains were grown in YPD and auxin (3-indoleacetic acid; Sigma Aldrich Cat. I3750) added to a final concentration of 0.75 mM to deplete AID-tagged proteins. YPD agar plates supplemented with auxin were made by cooling molten YPD 2% agar to 55°C prior to addition of auxin.

**Cohesion assays**

Sister chromatid cohesion was assessed at either the centromere-distal LYS4 locus or the centromere-proximal TRP1 locus on chromosome IV in which LacO arrays had been integrated. The GFP-LacI fusion allele integrated at HIS3 allows fluorescence microscopic visualization of LacO arrays. Cohesion was scored by growing cells to mid–log phase (OD600 ∼0.3) and arresting them in G1 using alpha factor at 10⁻⁸ M (Sigma-Aldrich T6901-5MG). After arresting for 3 h, auxin was added to a final concentration of 0.75 mM to deplete Smc3-AIDp for 1 h. Cells were released from G1 arrest by washing in YPD containing auxin and 0.1 mg/ml Pronase E (Sigma Aldrich). Cultures were incubated at 23°C and samples fixed either 1) periodically for assessing S-phase cohesion establishment or 2) after 3 h in which >95% of cells had arrested in G2/M. In addition to fixation for microscopy, samples were taken in parallel to assess DNA content by flow cytometry. Cohesion was scored by counting the GFP-LacI foci in the nucleus by fluorescence microscopy of fixed cells.

**Monitoring condensation at the rDNA locus**

Cells were grown as if for assessing cohesion by arresting in YPD containing auxin and nocodazole following release from G1. Cells were fixed, spheroplasted, and lysed to allow binding of chromosomes to slides as described previously (Guacci et al., 1994). Briefly, 1 ml of mid–M phase–arrested cells were fixed 2 h in 100 μl of 37% formaldehyde, washed twice in water, and spheroplasted for 1 h. Triton X-100 was added to 0.5% for 5 min, and then cells were pelleted and resuspended in water. Cells were then added to poly-Lysine-coated slides for 10 min. SDS (0.5%) was added for 10 min to solubilize membranes and release DNA masses and then removed. Slides were fixed in 3:1 methanol:acetic acid for 5 min and allowed
TABLE 1: Strains.

| Strain       | Genotype                                                                                      | Reference                  |
|--------------|----------------------------------------------------------------------------------------------|----------------------------|
| BRY467       | MATa smc3-D667-LEU2::leu2-3,112 smc3Δ::HPH lys4::LacO(DK)-NAT bar1 pHIS3-GFP LacI-Trp1::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT Bar1 pHIS3-GFP LacI-Trp1 his3-11,15 trp1-1 ura3-52 + pEU42 (SMC3 CEN URA3) | This study                 |
| BRY474       | MATa SMC3-LEU2::leu2-3,112 SMC3-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 SMC3-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY482       | MATa smc3-D667-LEU2::leu2-3,112 SMC3-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY602       | MATa smc3-6HA608-D667-URA3::ura3-52 SMC3-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY604       | MATa SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT leu2-3,112 pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY607       | MATa SMC3-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY621       | MATa SMC3-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY625       | MATa SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY647       | MATa SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY648       | MATa SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY649       | MATa SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY650       | MATa SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY676       | MATa SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY678       | MATa SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY680       | MATa SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY714       | MATa rad61Δ::HPMX SMC3-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY716       | MATa rad61Δ::HPMX SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY718       | MATa rad61Δ::HPMX SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY720       | MATa rad61Δ::HPMX SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY721       | MATa rad61Δ::HPMX SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY723       | MATa rad61Δ::HPMX SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY724       | MATa rad61Δ::HPMX SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY756       | MATa rad61Δ::HPMX SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |
| BRY815       | MATa rad61Δ::HPMX SMC3-CDC20-3V5-AID2-KANMX trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-Trp1 his3-11,15 ura3-52 bar1 | This study                 |

This study

Guacci et al. (2015)
| Strain   | Genotype                                                                 | Reference                  |
|----------|---------------------------------------------------------------------------|----------------------------|
| BRY832   | MATa smc1-D1164E SMC3-LEU2::leu2-2,3,112 SMC3-3V5-AID608 trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15, ura3-52 bar1 | This study                 |
| BRY833   | MATa SMC1-D1164E smc3-D667-LEU2::leu2-2,3,112 SMC3-3V5-AID608 trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15, ura3-52 bar1 | This study                 |
| BRY840   | MATa SCC2-3FLAG-AID2-HPHMX SMC3-608-STOP-TRP1 lys4::LacO(DK)-NAT leu2-2,3,112 pHIS3-GFP LacI-HIS3::his3-11,15 bar1 | This study                 |
| BRY842   | MATa SCC2-3FLAG-AID2-HPHMX smc3-3V5-AID608 trp1Δ::OsTIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15, ura3-52 bar1 | This study                 |
| BRY865   | MATa smc3-D667A-LEU2::leu2-2,3,112 SMC3-3V5-AID608 trp1Δ::pGPD1-TIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15, ura3-52 bar1 | This study                 |
| BRY866   | MATa smc3-D667A, K668A, R669A-LEU2::leu2-2,3,112 SMC3-3V5-AID608 trp1Δ::pGPD1-TIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15, ura3-52 bar1 | This study                 |
| DK5535   | MATa mcd1-Q266-3FLAG-URA3::ura3-52 MCD1-AID-KANMX pHIS3-GFP LacI-HIS3::his3-11,15 bar1 | Eng et al. (2014)          |
| DK5542   | MATa MCD1-AID-KANMX6 ADH1-OsTIR1-URA3::ura3-52 lys4::LacO(DK)-NAT trp1-1 GFP LacI-HIS3::his3-11,15 bar1 | Eng et al. (2014)          |
| DK5561   | MATa rad61Δ::HPHMX pADH1-TIR1-URA3::ura3-42 lys4::LacO(DK)-NAT trp1-1 GFP LacI-HIS3::his3-11,15 bar1 | Eng et al. (2014)          |
| TE228    | MATa PDS5-3V5-AID2-KANMX6 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15 trp1-1 ura3-52 | Eng et al. (2014)          |
| TE576    | MATa smc3-42 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15, leu2-2,3,112 bar1 trp1-1 + pEU42 (SMC3 CEN URA3) | Eng et al. (2015)          |
| TE578    | MATa smc3-42 smc3-K113R-URA2::ura3-2,3,112 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15, trp1-1 ura3-52 | Eng et al. (2015)          |
| VG3349-1B| MATa lys4::LacO(DK)-NAT trp1-1 GFP LacI-HIS3::his3-11,15 bar1 leu2-2,3,112 ura3-52 | Guacci and Koshland (2012) |
| VG3358-3B| MATa smc3-42 lys4::LacO(DK)-NAT trp1-1 pHIS3-GFP LacI-HIS3::his3-11,15 bra1 leu2-2,3,112 ura3-52 | Guacci and Koshland (2012) |
| VG3464-16C| MATa smc3Δ::HPH lys4::LacO(DK)-NAT bar1 pHIS3-GFP LacI-HIS3::his3-11,15 leu2-2,3,112 ura3-52 + pEU42 (SMC3 CEN URA3) | Guacci and Koshland (2012) |
| VG3486   | MATa smc3Δ::HPH lys4::LacO(DK)-NAT bar1 pHIS3-GFP LacI-HIS3::his3-11,15 leu2-2,3,112 ura3-52 + pEU42 (SMC3 CEN URA3) + pEU41 (SMC3 CEN LEU2) | Eng et al. (2015)          |
| VG3486-K113R| MATa smc3Δ::HPH lys4::LacO(DK)-NAT bar1 pHIS3-GFP LacI-HIS3::his3-11,15 leu2-2,3,112 ura3-52 + pEU42 (SMC3 CEN URA3) + pEU41-K113R (smc3-K113R LEU2 CEN) | Eng et al. (2015)          |
| VG3503-4A| MATa rad61Δ::HPHM X pADH1-TIR1-URA3::ura3-42 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15, leu2-2,3,112 bar1 | Çamdere et al. (2015)      |
| VG3506-5D| MATa eco1-203 LacO-NAT:10kb-CEN4 trp1-1 pHIS3-GFP LacI-HIS3::his3-11,15 leu2-2,3,112 ura3-52 | This study                 |
| VG3575-2C| MATa smc1-D1164E rad61Δ::HPHM X eco1Δ::G418 lys4::LacO(DK)-NAT GFP LacI-HIS3::his3-11,15 trp1-1 leu2-2,3,112 ura3-52 bar1 | Çamdere et al. (2015)      |
| VG3575-1A| MATa smc3Δ::HPHM rad61Δ::KANMX leu2-2,3,112 lys4::LacO(DK)-NAT ura3-52 bar1 pHIS3-GFP LacI-HIS3::his3-11,15 trp1-1 + pEU42 (SMC3 CEN URA3) | Guacci et al. (2015)       |
| VG3620-4C| MATa trp1Δ::pGPD1-TIR1-CaTRP1 lys4::LacO(DK)-NAT leu2-2,3,112 pHIS3-GFP LacI-HIS3::his3-11,15, ura3-52 bar1 | Çamdere et al. (2015)      |
| VG3633-2D| MATa eco1-3V5-AID2-KANMX trp1Δ::pGPD1-TIR1-CaTRP1 lys4::LacO(DK)-NAT leu2-2,3,112 pHIS3-GFP LacI-HIS3::his3-11,15, ura3-52 bar1 | Çamdere et al. (2015)      |
| VG3651-3D| MATa SMC3-3V5-AID608 trp1Δ::pGPD1-TIR1-CaTRP1 lys4::LacO(DK)-NAT pHIS3-GFP LacI-HIS3::his3-11,15, leu2-2,3,112 ura3-52 bar1 | Çamdere et al. (2015)      |

**TABLE 1:** Strains. Continued
to dry. Cells on slides were treated with RNase A and Proteinase K and subject to a series of short 70, 80, 90, and 100% ethanol washes. After drying, DNA masses were visualized with 4',6-diamidino-2-phenylindole (DAPI) and rDNA morphology scored.

**Chromatin immunoprecipitation**

Cells were grown as if for assessing cohesion by arresting at mid-M phase in yeast extract, peptone, and dextrose (YPD) containing auxin and nocodazole following release from G1 arrest. ChiP was performed as described previously (Wahba et al., 2013; Eng et al., 2014), except that chromatin shearing was performed on a Bioruptor Pico machine (Diagenode, Denville, NJ) for 5 min (30 s on/off cycling). Immunoprecipitation was performed using monoclonal mouse anti-HA (Roche), monoclonal mouse anti-V5 (ThermoFisher), polyclonal rabbit anti-Pds5p (Covance Biosciences, Princeton, NJ), or polyclonal rabbit anti-Mcd1p (Covance) antibodies. A no antibody control was always included to assess specificity of chromatin recovery.

**Detection of Smc3-K113 acetylation by Western blotting**

Cells were grown to OD$_{600}$ = 0.5 in YPD at 23°C before addition of auxin to 0.75 mM and incubation for 1 h. Nocodazole was added to a final concentration of 15 μg/ml to arrest cells in mid-M phase. Cells were pelleted and resuspended in lysis buffer consisting of 25 mM HEPES, pH 8.0, 2 mM MgCl$_2$, 100 μM EDTA, 500 μM egtazic acid (EGTA), 1% NP-40, 150 mM KCl, 15% glycerol, Complete-Mini EDTA-free protease inhibitor cocktail (Roche), 10 mM sodium butyrate, and 20 mM beta-glycerophosphate. Cells were incubated in buffer for 30 min on ice, and then glass beads were added to a 1:1 volume ratio before bead beating for 3 min. Lysates were pelleted at 14,000 rpm for 30 min on ice, and then glass beads were added to a 1:1 volume ratio. Lysates were boiled in 120 mM HEPES, pH 7.0, containing 1% SDS at 95°C for 5 min and then diluted 1:1 in 2× Laemmli sample buffer. Smc3-K113 acetylation was detected by blotting with monoclonal mouse antibody (a gift from K. Shirahige, University of Tokyo) at a concentration of 1:1000 in 5% milk-PBST.

**Chromosomes spreads and microscopy**

Cells were grown as if for assessing cohesion by arresting in mid-M phase in YPD containing auxin and nocodazole following release from G1 arrest. Chromosome spreads were prepared as described previously (Wahba et al., 2013). Slides were incubated with 1:5000 rabbit polyclonal anti-Mcd1p and 1:5000 mouse anti-V5 antibody (Life Technologies). Antibodies were diluted in blocking buffer (5% BSA, 0.2% milk, 1× phosphate-buffered saline, 0.2% Triton X-100). Secondary Alexa Fluor 488–conjugated chicken anti-mouse and Alexa Fluor 568–conjugated donkey anti-rabbit (ThermoFisher Cats. A21200 and A10042) antibodies were diluted 1:5000 in blocking buffer. Indirect immunofluorescence was detected on an Axio-plan2 microscope (Zeiss, Thornwood, NY) using the 100× objective (numerical aperture 1.40) which is equipped with a Quantix charge-coupled camera (Photometrics).

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