Smc3 Deacetylation by Hos1 Facilitates Efficient Dissolution of Sister Chromatid Cohesion during Early Anaphase
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**Smc3 Deacetylation by Hos1 Facilitates Efficient Dissolution of Sister Chromatid Cohesion during Early Anaphase**

**Graphical Abstract**

Smc3 deacetylation by Hos1 facilitates removal of cohesins from chromosomes in early anaphase, without changing efficiency of Scc1 cleavage by separase.

**Highlights**

- Depletion of Hos1 causes a significant delay in completing chromosome segregation
- Smc3 deacetylation by Hos1 facilitates cohesin removal from chromosomes in anaphase
- Depletion of Hos1 does not change the efficiency of Scc1 cleavage by separase
- Smc3 deacetylation by Hos1 probably leads to disengagement of Smc1-Smc3 heads

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**In Brief**

Sister chromatid cohesion relies on cohesins and is dissolved when cells enter anaphase. Li et al. show that, in addition to cleavage of cohesin Scc1 by separase, deacetylation of cohesin Smc3 by Hos1 is important for efficient removal of cohesins from chromosomes, which ensures timely chromosome segregation in mitosis.

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Smc3 Deacetylation by Hos1 Facilitates Efficient Dissolution of Sister Chromatid Cohesion during Early Anaphase

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SUMMARY

Cohesins establish sister chromatid cohesion during S phase and are removed when cohesin Scc1 is cleaved by separase at anaphase onset. During this process, cohesin Smc3 undergoes a cycle of acetylation: Smc3 acetylation by Eco1 in S phase stabilizes cohesin association with chromosomes, and its deacetylation by Hos1 in anaphase allows re-use of Smc3 in the next cell cycle. Here we find that Smc3 deacetylation by Hos1 has a more immediate effect in the early anaphase of budding yeast. Hos1 depletion significantly delayed sister chromatid separation and segregation. Smc3 deacetylation facilitated removal of cohesins from chromosomes without changing Scc1 cleavage efficiency, promoting dissolution of cohesion. This action is probably due to disengagement of Smc1-Smc3 heads prompted by de-repression of their ATPase activity. We suggest Scc1 cleavage per se is insufficient for efficient dissolution of cohesion in early anaphase; subsequent Smc3 deacetylation, triggered by Scc1 cleavage, is also required.

INTRODUCTION

Cohesion between sister chromatids is established during DNA replication and removed when cells enter anaphase. Timely regulation of cohesion is crucial; if cohesion is lost precociously, or its removal is delayed, chromosome mis-segregation could result. Sister chromatid cohesion relies on the tetrameric cohesin complex, which is composed of Scc1 (also called Mcd1 or Rad21), Scc3, Smc1, and Smc3, which forms a ring structure embracing sister chromatids (Nasmyth and Haering, 2009).

Cohesins are loaded onto chromosomes in telophase or G1 phase, by the cohesin loader. During the subsequent S phase, Smc3 is acetylated by Eco1 acetyl-transferase (also called Ctf7). Smc3 acetylation prevents ATPase activity of the Smc1-Smc3 heads, which in turn inhibits opening of the Smc3-Scc1 interface (Chan et al., 2012; Beckouët et al., 2016; Çamdere et al., 2015; Eibatsh et al., 2016; Huber et al., 2016; Murayama and Uhlmann, 2015) and counteracts the activity of Wpl1 (also called Wapl and Rad61) (Rolef Ben-Shahar et al., 2008; Rowland et al., 2009; Unal et al., 2008; Zhang et al., 2008). Smc3 acetylation leads to replicated sister chromatids being stably trapped inside the cohesin ring complex. Sister chromatid cohesion is thus maintained until the onset of anaphase, when the cysteine protease separase becomes active and cleaves Scc1 (Nasmyth and Haering, 2009; Uhlmann, 2001). This leads to opening of the cohesin ring complex and removal of cohesion, allowing chromosome segregation to opposite spindle poles. During this process, Scc1 cleavage also allows Hos1 deacetylase to remove acetyl groups from Smc3 (Beckouët et al., 2010; Borges et al., 2010; Chan et al., 2013; Deardorff et al., 2012; Xiong et al., 2010). It is suggested that this deacetylation of Smc3 allows it to be reused for cohesion in the next cell cycle, although this recycling is not essential for cell growth as new (and therefore non-acetylated) Smc3 is also expressed. These processes are essentially conserved from yeast to humans (Uhlmann, 2010).

It is unclear, however, whether Smc3 deacetylation has any immediate effect in regulating cohesins at anaphase onset when the deacetylation actually occurs. Moreover, although Scc1 cleavage leads to removal of cohesins from chromosomes (Uhlmann et al., 2000), it is unknown whether Scc1 cleavage itself is sufficient for this process or whether any events downstream of Scc1 cleavage are also involved. Here we address these questions using budding yeast as a model organism.

RESULTS

Cohesin Deacetylase Hos1 Promotes Efficient and High-Fidelity Sister Chromatid Segregation during Anaphase

To address the role of Smc3 deacetylation in anaphase, we aimed to abrogate the function of Hos1 deacetylase. Deletion of the hos1 gene weakens sister chromatid cohesion, presumably due to a lack of Smc3 recycling for the next cell cycle (Beckouët et al., 2010; Borges et al., 2010). Therefore, to investigate immediate effects of abrogated Hos1 function in anaphase, we tagged the hos1 gene with an auxin-inducible degron and induced degradation by adding auxin (Nishimura et al., 2009). Hos1 was depleted within 20–40 min after adding auxin (Figure 1A). Smc3 deacetylation, which normally occurs in anaphase, was impaired after Hos1 depletion (Figure 1B), as reported in hos1-deleted cells (Beckouët et al., 2010; Borges et al., 2010; Xiong et al., 2010). After depletion of Hos1 following...
Figure 1. Depletion of Hos1 Leads to a Delay in Chromosome Segregation

(A) Hos1 was rapidly depleted using the auxin-induced degron system. Cells with (T11218) and without (T11219) an auxin-induced degron tag (aid) for HOS1 were incubated with auxin (NAA 0.5 mM), and analyzed after 20, 40, and 60 min by western blotting with an anti-AID tag antibody.

(B) HOS1 wild-type with NAA and hos1-aid with NAA were analyzed for release from G1 arrest using western blotting with an anti-AID tag antibody and antibodies to Smc3-Ac and Cdc28.

(C) Histone SPB Myo1

(D) Chromosome XV

(E) % of cells

(legend continued on next page)
G1 phase, sister chromatid cohesion was still maintained in metaphase as robustly as it was in Hos1 wild-type cells (Figure S1A).

After Hos1 depletion following G1 phase, we investigated chromosome segregation by visualizing chromosomes using histones fused with cyan fluorescent protein (CFP). In the same cells, we also visualized spindle poles with mCherry red fluorescent protein to evaluate spindle length and a myosin ring at the bud neck to monitor cytokinesis. In most of the wild-type cells, chromosomes completed segregation to opposite poles (i.e., histone signals split into two) within 30 min after anaphase onset, whereas in the majority of Hos1-depleted cells, it took substantially longer for chromosomes to complete segregation (Figure 1C). We also depleted Hos1 during metaphase arrest (rather than following G1), subsequently released cells to anaphase, and found similar outcomes (Figure S1B). We next analyzed segregation of a chosen chromosome (chromosome XV) by marking three loci (CEN, HIS, and TEL) with tet operators that could bind TetR-GFP (green fluorescent protein) fusion proteins, thus visualizing them as small GFP dots. In wild-type cells, CEN, HIS, and TEL dots—in that order—showed segregation to opposite poles (Renshaw et al., 2010). After Hos1 depletion following G1 phase, it took longer for these dots to complete their segregation (Figure 1D). Thus, chromosome segregation is substantially delayed during anaphase after Hos1 depletion.

In wild-type cells, anaphase spindle elongation takes place in two phases: an initial, rapid elongation and a subsequent, slower elongation (Straight et al., 1997). In Hos1-depleted cells, the initial, rapid spindle elongation occurred normally but the subsequent slower elongation was retarded and the completion of cytokinesis was significantly delayed (Figures S1C and S1D). We next addressed whether the spindle assembly checkpoint (SAC) is involved in the delay of chromosome segregation in Hos1-depleted cells. This was not the case, since Hos1-depleted cells showed a similar delay in chromosome segregation in the absence of the SAC (mad2 deletion) (Figure S1E).

We then analyzed the fidelity of chromosome segregation with Hos1 depletion. We visualized TEL (right telomere) on chromosome XV as a GFP dot. Following anaphase and cytokinesis with Hos1 depletion, the proportion of cells with no TEL dot or two TEL dots increased from 4.6% to 13.3% (Figure 1E). By contrast, such an increase was not found in Hos1 wild-type cells. Thus Hos1 depletion leads to an increased rate of chromosome mis-segregation.

Hp3 acetylation counteracts the Wpl1 function of opening the Smc3-Scc1 interface (see references in Introduction), so Hos1 depletion might delay chromosome segregation by counteracting the Wpl1 function. If so, we expected that Wpl1 depletion would show a similar outcome to that of Hos1 depletion. To test this, we depleted Wpl1 using an auxin-inducible degron (Figure S1F). After Wpl1 depletion, there was no substantial delay in chromosome segregation, in contrast to Hos1 depletion (Figure 1C). Collectively, Hos1 depletion delays chromosome segregation and increases chromosome mis-segregation. The delay in chromosome segregation is not dependent on the SAC or due to suppression of Wpl1 function.

Hos1 Facilitates Removal of Sister Chromatid Cohesion and Chromosome-Bound Cohesins during Early Anaphase without Affecting Scc1 Cleavage

Is the delay in chromosome segregation found with Hos1 depletion due to impaired regulation of sister chromatid cohesion? To address this, we visualized CEN and HIS3 loci as GFP dots on chromosome XV (Figure 2A, diagram); CEN and HIS3 dots were distinguishable, as the former segregated earlier and showed a higher intensity than the latter (see Figure 2B, cell images). We evaluated separation timing of sister HIS3 loci after anaphase onset (Figure 2A, graph). In Hos1-depleted cells, sister

(B) Smc3 acetylation remained in anaphase after depletion of Hos1. Cells of T11219 and T11218 (see A) were released from G1 arrest to YPAD medium (at 0 min) containing NAA to deplete Hos1 aid. Mating pheromone was re-added to the culture after bud formation to re-arrest cells in the following G1 phase. The Smc3 acetylation (Smc3-Ac) was checked every 20 min by western blotting with anti-acetyl Smc3 (Borges et al., 2010) (top) and anti-Cdc28 (bottom, loading control) antibody.

(C) Segregation of chromosomes was delayed significantly after Hos1 depletion but only marginally after Wpl1 depletion. hos1-aid (T11218) and wpl1-aid (T11432) cells with HTB2-CFP, SPC42-mCherry, and MYO11-mCherry—where Htb2 is a histone h2B, Spc42 is a spindle-pole-body (SPB) component, and Myo1 is a component of the acto-myosin ring at the bud neck—were released from G1 arrest to YPAD medium containing 0.5 mM NAA. Control wild-type cells (T11219) were treated in the same way. From 80 min after release from G1, images were acquired every 2 min for 2 hr. Representative time-lapse images show chromosome segregation during anaphase (left; scale bar, 1 μm). Anaphase onset (time 0 in the image) is defined as the time when the distance between two SPBs reached 2.5 μm. The time of completion of chromosome segregation—i.e., disappearance of histone signals at the bud neck following their segregation—was plotted in individual cells (right). The bars in the graph show mean and SEM. Orange squares in middle (Hos1 depletion) show the time when time-lapse observation finished before chromosome segregation was completed. p values were obtained by t test. In the examples (left), chromosome segregation completed at 22, 44, and 22 min (from top to bottom). See Movies S1, S2, and S3.

(D) Segregation of HIS3 and TEL loci on chromosome XV was delayed in Hos1-depleted cells. HOS1 wild-type (T10830) and hos1-aid (T10829) cells with Tet repressors fused with GFP (TetR-GFP) and tet operators (tetO) inserted at three loci on chromosome XV (CEN, HIS3, TEL; top left diagram) were treated as in (C). From 80 min after release from G1, images were acquired every 4 s for 45 min. Representative time-lapse images (top right) show segregation of the three GFP-labeled loci during anaphase (scale bar, 1 μm). The white, red, and yellow arrows indicate sister CEN, HIS3, and TEL loci, respectively, which were identified as they segregate in this order (Renshaw et al., 2010). Anaphase onset (time 0 in the image) was defined as the time when the distance between two CENs reached 3 μm. Time required for HIS3 and TEL locus segregation (defined as the time when CEN-HIS3 or CEN-TEL distance becomes <1.5 μm in the bud) is plotted in the graph (bottom) where bars indicate mean and SEM. p values were obtained by t test. See Movies S4 and S5.

(E) Hos1 depletion leads to an increase in chromosome mis-segregation. HOS1 wild-type (T11113) and hos1-aid (T11112) cells with SPC42-mCherry, TetR-GFP, and TEL-tetO (on chromosome XV) were treated as in (B). Images were acquired during arrest in G1 (before NAA addition; 1st G1) and at 180 min after release from G1 arrest (after NAA treatment and cytokinesis; 2nd G1). Cells with zero and two TEL dots were counted (graph on left) and representative images are shown (right). The cell, highlighted in color, carried two TEL dots. Sample numbers (from left to right): n = 424, 657, 480, and 450. p values were obtained by Fisher’s exact test (one versus zero/two dots). n.s., not significant. See also Figure S1.
HIS3 dots took longer to separate, on average, than they did in wild-type cells. Sister HIS3 separation may occur either before or after spindle pulling forces are applied to this locus through CEN and chromosome arms. We next evaluated the frequency of sister HIS3 separation after anaphase onset but before any spindle-pulling force was applied on this locus. To monitor this, the ADE2 locus was visualized as a CFP dot on chromosome XV (Figure 2B, diagram; Renshaw et al., 2010). We focused on the period when sister CEN dots were pulled toward opposite spindle poles but sister ADE2 dots were not, and we evaluated separation of sister HIS3 dots (Figure 2B). After Hos1 depletion, more cells showed associated (non-separated) sister HIS3 dots during this period than was the case in wild-type cells (Figure 2B, graph). Thus, in Hos1-depleted cells, sister chromatid cohesion...
persists longer in early anaphase before any spindle force is applied.

We next compared cellular defects caused by Hos1 depletion and hos1 gene deletion (hos1-D). Hos1 depletion freshly removed Hos1 in the current cell cycle, whereas in hos1-D cells Hos1 was not functional in both previous and current cell cycles. In hos1-D cells, cohesion was weakened in metaphase (Figure S2A, left), presumably due to failure in re-cycling Smc3 for the next cell cycle (Beckouët et al., 2010; Borges et al., 2010). Intriguingly, hos1-D cells showed only a marginal delay in completing chromosome segregation, in contrast to cells with “fresh” Hos1 depletion (Figure S2A, right); probably a weaker cohesion in metaphase offset a delay in cohesion removal in anaphase, with hos1-D.

Is the persistent cohesion found during early anaphase in Hos1-depleted cells due to a delay in the removal of cohesins? To address this question, we let cells synchronously enter anaphase (by Cdc20 depletion and subsequent re-expression) but prevented them from entering the next cell cycle (Figure S2B). To address this question, we let cells synchronously enter anaphase (by Cdc20 depletion and subsequent re-expression) but prevented them from entering the next cell cycle (Figure S2B). In this situation, Hos1 wild-type and Hos1-depleted cells showed similar spindle elongation during anaphase (Figures S2C and S2D). We then quantified the amount of chromosome-bound cohesin Smc1 by chromatin immunoprecipitation and quantitative PCR (ChIP–qPCR) at two chromosome loci (Figure 2C). From 10 to 25 min after Cdc20 re-expression, peaks of Smc1 were still found above the background all along chromosomes in Hos1-depleted cells, whereas most of these were abolished in Hos1 wild-type cells. Thus, Hos1 depletion delays cohesin removal from chromosomes in early anaphase.

Smc1 cleavage by separase leads to removal of cohesins at the onset of anaphase (Uhlmann et al., 2000). We next addressed whether the efficiency of Smc1 cleavage was affected by Hos1 depletion. For this, we quantified the C-terminal Scc1 cleavage product (Scc1-C) upon entry into anaphase (Figure 2D, left). Scc1-C was detected at 10 and 20 min, and its amount was similar in wild-type and Hos1-depleted cells. Scc1-C degradation relies on Ubr1, which is the E3 ubiquitin ligase of the N-end rule pathway (Rao et al., 2001), and so we also quantified Scc1-C after Ubr1 depletion (Figure 2D, right). In this condition, the amount of Scc1-C increased at 20 min and subsequently remained high; the amount was similar in wild-type and Hos1-depleted cells. Thus, Hos1 depletion does not change the efficiency of Scc1 cleavage by separase. Hos1 facilitates removal of cohesins from chromosomes in early anaphase without changing Scc1 cleavage efficiency.

**Smc3 Deacetylation by Hos1 at K112 and K113 Leads to Efficient Cohesin Removal in Anaphase and Timely Chromosome Segregation**

We next addressed whether deacetylation of Smc3 by Hos1 is crucial for removal of cohesins from chromosomes and timely sister chromatid segregation. Smc3 is acetylated at lysines (K) 112 and 113 by Eco1 during S phase (Rolef Ben-Shahar et al., 2008; Unal et al., 2008) and deacetylated by Hos1 in early anaphase (Beckouët et al., 2010; Borges et al., 2010; Xiong et al., 2010). If Smc3 deacetylation by Hos1 at K112 and K113 is important for removal of cohesins in early anaphase, we can make the following two predictions: first, acetyl-Smc3 would be bound to anaphase chromosomes longer in Hos1-depleted cells; second, non-acetyl smc3 mutants—e.g., replacement of lysines with arginines at these sites (K112R K113R)—would rescue timely chromosome segregation in Hos1-depleted cells.

We tested the first prediction by quantifying acetyl-Smc3 (at K112 and K113) that was bound to chromosomes. Acetyl-Smc3 was detected on fixed and immobilized chromosomes (Figure S3A). In metaphase, acetyl-Smc3 was detected at a similar level in Hos1 wild-type and Hos1-depleted cells (Figure 3A). However, in anaphase, more acetyl-Smc3 was detected on chromosomes in Hos1-depleted cells. For comparison, we quantified HA-tagged Smc3 on chromosomes, and this showed a similar behavior to that of acetyl-Smc3 (Figure S3B). Thus, acetyl-Smc3 indeed remains longer on anaphase chromosomes in Hos1-depleted cells.

It is not straightforward to test the second prediction, since non-acetyl smc3 mutants (K112R K113R) are lethal (Rolef Ben-Shahar et al., 2008; Unal et al., 2008). This lethality is due to the destabilized Smc3-Scc1 interface, which hampers establishment of cohesion (Chan et al., 2012). We attempted to keep non-acetyl smc3 mutant cells alive by fusing the C terminus of Smc3 to the N terminus of Scc1 (Smc3-Scc1). Cells with an Smc3-Scc1 fusion protein remain viable even if their original SCC1 and SM3C are deleted (Gruber et al., 2006). We introduced non-acetyl K112R K113R mutations to the Smc3-Scc1 fusion and found that this non-acetyl Smc3-Scc1 fusion can maintain cell viability, as can the “wild-type” Smc3-Scc1 fusion (Figure S3C). The non-acetyl and “wild-type” Smc3-Scc1 fusions could establish and maintain cohesion similarly in metaphase, though cohesion in these cells is slightly weaker than in Smc3+ Scc1+ (non-fusion) control cells (Figure S3D).

We then measured the period from anaphase onset to completion of chromosome segregation in these cells (Figure 3B). First, we found that in the presence of the Smc3-Scc1 fusion, Hos1 depletion caused a delay in completion of chromosome segregation (Figure 3B; left versus middle in graph). The delay by Hos1 depletion was smaller with the Smc3-Scc1 fusion than it was with non-fusion Smc3+ Scc1+ (compare Figures 3B and 1C), presumably due to slightly weaker cohesion in metaphase (Figure S3D). Second, in Hos1-depleted cells, non-acetyl Smc3-Scc1 fusion alleviated a delay in chromosome segregation observed with “wild-type” Smc3-Scc1 fusion (Figure 3B; middle versus right in graph). Therefore, at least in the context of Smc3-Scc1 fusion, Smc3 non-acetyl mutants rescue timely chromosome segregation in Hos1-depleted cells. We also found that this rescue is independent of the SAC (Figure S3E). In summary, Smc3 K112 and K113 are Hos1 substrates whose deacetylation is crucial for efficient chromosome segregation.

**Engineered Smc3 Cleavage by Separase Rescues Efficient Chromosome Segregation in Hos1-Depleted Cells**

How does Smc3 deacetylation by Hos1 facilitate cohesin removal in early anaphase? It was suggested that Smc3
acetylation inhibits ATPase activity of Smc1-Smc3 heads and that, while ATP hydrolysis is prohibited, Smc1-Smc3 heads remain engaged (Beckouët et al., 2016; Çamdere et al., 2015; El-batsh et al., 2016; Huber et al., 2016; Murayama and Uhlmann, 2015) (Figure 4A). Smc3 deacetylation could disengage Smc1-Smc3 heads and thus facilitate the exit of chromosomes from the ring composed of Smc1 and Smc3 (and from the entire cohesin ring, since Scc1 is cleaved at anaphase onset). This model would give the following prediction: if there were a way to open up the Smc1-Smc3 ring without relying on opening Smc1-Smc2-R). These insertion sites were selected so as not to perturb the Smc3 coiled-coil structure (Gruber et al., 2003). As a control, mutated separase cleavage sites were also inserted in the same way (Smc3-2D).

When cells expressed Smc3-2R or -2D as the sole Smc3 protein, these Smc3 mutants supported robust cohesion as did Smc3 wild-type (Figure S4A). As expected, we could detect an Smc3 cleavage product with Smc3-2R, but not with Smc3-2D (Figure 4B, lanes 2 and 3). To confirm that the engineered Smc3 cleavage is able to open the Smc1-Smc3 ring, we asked

Figure 3. Smc3 Deacetylation at K112 and K113 by Hos1 Leads to Efficient Cohesin Removal in Anaphase and Timely Chromosome Segregation

(A) A larger amount of acetyl-Smc3 remains on anaphase chromosomes in Hos1-depleted cells. HOST1 wild-type (T13179) and hos1-aid (T13180) cells with PGAL-CDC20 Spc42-mCherry were arrested in metaphase and subsequently released to anaphase, as in Figure 2C. NAA was added to deplete Hos1-aid, as in Figure 2C. Cells in metaphase arrest and in anaphase (10–15 min after CDC20 re-expression) were taken, and chromosomes were fixed and immobilized on a slide glass immediately after cell lysis. Representative cells are shown on the left. The distance between two SPBs (right, top) and acetyl-Smc3 (ac-Smc3) signals on chromosomes (right, bottom) were quantified. DNA signals showed elongation, and the distance between two ac-Smc3 signals was enlarged in this assay when cells proceeded from metaphase to anaphase (Renshaw et al., 2010). Cells in metaphase, early anaphase, and late anaphase were defined by SPB-SPB distance (right, bottom). Scale bars, 1 μm; a.u., arbitrary unit; n.s., no significant difference. p value was obtained by t test. Bars and error bars show means and SEMs.

(B) Smc3 non-acetyl mutants rescue timely chromosome segregation in Hos1-depleted cells. HOST1 wild-type SMC3-SCC1 fusion (T12684), hos1-aid SMC3-SCC1 fusion (T12665), and hos1-aid SMC3-K112R K113R–SCC1 fusion (T12666) cells with SPCC42-mCherry, MYO1-mCherry, and HTB2-CPF were treated, and images were acquired as in Figure 1C. In these cells, the original SMC3 and SCC1 genes were deleted. Representative time-lapse images show chromosomes segregation during anaphase (scale bar, 1 μm). Anaphase onset (0 min) is defined, and time required for completion of chromosome segregation was analyzed (graph) as in Figure 1C. p values were obtained by t test. In the examples shown, chromosome segregation completed at 24, 42, and 24 min (from top to bottom).

See also Figure S3.
(A) Diagram shows that Smc3 acetylation inhibits ATPase activity of Smc1-Smc3 heads, and Smc1-Smc3 heads remain engaged until ATP is hydrolyzed (top) (Beckouet et al., 2016; Çamdere et al., 2015; Ebahs et al., 2016; Huber et al., 2016; Murayama and Uhlmann, 2015). In the absence of Smc3 acetylation, ATP is hydrolyzed and Smc1-Smc3 heads are disengaged (middle). Subsequent ATP binding leads to opening of the Smc3-Scc1 interface and destabilizes cohesin association with chromosomes (bottom).

Figure 4. Engineered Smc3 Cleavage by Separase Rescues Efficient Chromosome Segregation in Hos1-Depleted Cells

(B) WT: wild-type
OE: overexpression

(C)

(D)

(E)

(legend continued on next page)
if Smc3-2R could rescue chromosome segregation in the presence of an Scc1 whose separase cleavage sites were mutated (Scc1-2D; Uhlmann et al., 1999). Indeed, in the presence of Scc1-2D, we observed a partial rescue of chromosome segregation by Smc3-2R, but not by Smc3-2D (Figure 4C, orange and blue bars). Since Smc3 is not a natural target of separase, Smc3-2R cleavage by separase may not be efficient, but it may be made more so if separase is overexpressed. With separase overexpression (Figure S4B), cohesin regulation was largely normal (Figures S4C and S4D), but Smc3-2R showed a higher rate of chromosome segregation in the presence of Scc1-2D (Figure 4C, magenta bars). Thus, the engineered cleavage of Smc3 by separase can open the cohesin ring when non- cleavable Scc1 is present.

We then addressed whether the Smc3 cleavage (with separase overexpression) rescues timely chromosome segregation in Hos1-depleted cells (Figure 4D). Hos1 depletion delayed chromosome segregation in the majority of cells with Smc3-2D. Importantly, this delay was significantly alleviated with Smc3-2R in Hos1-depleted cells. By contrast, in Hos1 wild-type cells, neither Smc3-2D nor Smc3-2R changed the timing of chromosome segregation (Figure S4E). Thus, the engineered Smc3 cleavage can rescue timely chromosome segregation in Hos1-depleted cells. The result is consistent with Smc3 deacetylation by Hos1 facilitating disengagement of the Smc1-Smc3 heads to promote removal of cohesins from chromosomes at anaphase onset.

DISCUSSION

Our study suggests that Smc3 deacetylation by Hos1 facilitates cohesin removal and efficient dissolution of sister chromatid cohesion in early anaphase (Figure 4E). This action is presumably due to de-repression of the ATPase activity of Smc1-Smc3 heads prompted by Smc3 deacetylation, which leads to Smc1-Smc3 head disengagement. It has been shown that Smc3 deacetylation by Hos1 is triggered by Scc1 cleavage by separase (Beckouet et al., 2010; Chan et al., 2013). Thus Scc1 cleavage and subsequent Smc1-Smc3 head disengagement together open up the cohesin ring complex, facilitating sister chromatin separation and segregation. Both steps are required to fully open up the cohesin ring complex. In this context, Scc1 cleavage per se is not sufficient for efficient removal of cohesin; Smc3 deacetylation of Hos1 (which is triggered by Scc1 cleavage) is also required. It has been thought that Scc1 cleavage by separase is the final regulatory step to release cohesin and to dissolve cohesion at anaphase onset. Our results provide evidence that there is another regulatory step required for efficient cohesin release and cohesion dissolution on chromosomes. Nevertheless, while Scc1 cleavage is essential for cohesin removal (Uhlmann et al., 1999), Smc3 deacetylation by Hos1 is important only for efficient cohesin removal (Figure 2C). Without Hos1, Smc1-Smc3 head disengagement would be delayed but would eventually occur (Figure 4E). We assume that Smc1-Smc3 head disengagement takes place spontaneously, but with a delay, in Hos1-depleted cells.

Our results suggest that Smc3 K112 and K113 are important targets of Hos1, whose deacetylation promotes chromosome segregation in early anaphase (Figure 3). However, we cannot rule out that deacetylation of additional targets by Hos1 also contributes to this process. For example, it has been recently reported that coiled-coil regions of Smc1 and Smc3 are acetylated (in addition to Smc3 K112 and K113), facilitating association of Smc1 and Smc3 in these regions (Kulemzina et al., 2016). It is currently unknown whether Hos1 deacetylates these sites. If this is the case, it is possible that deacetylation of these sites by Hos1 facilitates dissociation of Smc1 and Smc3 in their coiled-coil regions and promotes removal of cohesins after Scc1 cleavage.

The cohesion structure and the Smc3 acetylation cycle are well conserved in evolution from yeast to humans. It is therefore
likely that the mechanism found in yeast in this study is conserved in higher eukaryotes. Indeed, HDAC8 has been identified as the Smc3 deacetylase in human cells, and its inactivation leads to longer retention of cohesins, including cleaved Scc1, on chromosomes during anaphase (Deardorff et al., 2012). Further study may find a delay in chromosome segregation and cytokinesis with inactive HDAC8 in human cells, as we have found here in budding yeast. Mutations in HDAC8 are associated with Cornelia de Lange syndrome, a dominantly inherited congenital disorder (Deardorff et al., 2012). It is possible that delays in chromosome segregation are associated with development of the disease.

STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures, one table, and five movies and can be found with this article online at https://doi.org/10.1016/j.molcel.2017.10.009.

AUTHOR CONTRIBUTIONS

Conceptualization, Methodology, Visualization, and Writing (Original Draft) – S.L. and T.U.T.; Investigation, Formal Analysis, and Validation – S.L. and Z.Y.; Resources and Data Curation – S.L.; Funding Acquisition, Supervision, Project Administration, and Writing (Review & Editing) – T.U.T.

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REFERENCES


### STAR Methods

**KEY RESOURCES TABLE**

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EXPERIMENTAL MODEL AND SUBJECT DETAILS

All yeast *Saccharomyces cerevisiae* strains, used in this study, had a W303 background (derived from K699 and K700 from the Nasmyth laboratory). The genotypes of the yeast strains are shown in Table S1.

METHOD DETAILS

**Yeast strains and cell culture**

Methods for yeast culture have been described previously (Amberg et al., 2005; Renshaw et al., 2010). To synchronize cells in the cell cycle, yeast cells were arrested in G1 phase by treatment with yeast mating hormone (α or a factor) and subsequently released to fresh media (Amberg et al., 2005; O’Reilly et al., 2012). Cells were cultured at 25°C in YPA medium (1% yeast extract, 2% peptone, 0.01% adenine hydrochloride) containing 2% glucose (YPAD), unless otherwise stated. To activate MET3 promoter, cells were incubated in methionine drop-out medium. The MET3 promoter was suppressed by adding 2 mM methionine to the relevant medium. To activate GAL1-10 promoter, cells were incubated in YPA medium containing 2% raffinose for at least 2.5 h, and subsequently 2% galactose was added. SPC42, MYO1 and NIC96 genes were tagged with mCherry or with 4x3mCherry at their C terminus at their original gene loci using a one-step PCR method with plasmids pKS391 (mCherry-natNT2; a gift from Ken Sawin lab) and pT909 (4x3mCherry-natNT2) as PCR templates. HTB2 was tagged with CFP at its C terminus at the original gene locus using a one-step PCR method with plasmid pKT101 (CFP-spHIS5; EUROSCARF) as a PCR template. SCC1 and SMG3 were tagged with six tandem copies of HA at their C terminus at the original gene locus using a one-step PCR method with plasmid pYM15 (EUROSCARF) as a template. HOS1 and WPL1 (RAD61) were tagged with three tandem copies of mini aid tag at their C terminus by one-step PCR method, using plasmid pMK151 (3xmini-aid-kanMX; Kubota et al., 2013) as a template. Their protein degradation was facilitated within cells carrying rice *Oryza sativa* TIR1 (osTIR1, expressed from ADH1 promoter) in the presence of 0.5 mM auxin NAA (1-naphaleneacetic acid) (Nishimura et al., 2009). Strains smc3D, scc1D and mad2D were generated with a PCR-based gene deletion method (Amberg et al., 2005), using the plasmids pFA6-αphN1, pFA6-natNT2 and pFA-kanMX4 (EUROSCARF), respectively. TetR-GFP, TetR-3xCFP, 3xCFP-LacI, P<sub>GAL</sub>-CDC20 and P<sub>MET3</sub>-CDC20 were used in our previous study (Renshaw et al., 2010), in which the original papers about these constructs are cited. Construction of cdc15-as1(L99G) (D’Aquino et al., 2005),
SPB–SPB distance was < 2.5 m when cells proceeded from metaphase to anaphase. Cells in metaphase, in early anaphase and in late anaphase were defined as elongated and the distance between two SPBs was enlarged, as demonstrated previously (Renshaw et al., 2010), in chromosome spreads, chromosomes showed elongation and the distance between two SPBs was enlarged. Chromosome spreads (chromosome fixation and immobilization) (Renshaw et al., 2010; Tanaka et al., 1997). Subsequently, they were processed for immunostaining using mouse monoclonal anti-acetyl-Smc3 (a gift from Shirahige lab; Borges et al., 2010) or anti-HA (16B12) antibody. The primary antibody was visualized by incubation with the secondary antibody goat anti-mouse IgG (Alexa Fluor 647). Chromosome DNA was stained with Hoechst. Spindle Pole Bodies (SPBs) were visualized by expression of Spc42 fused with mCherry.

**Construction of Smc3 cleavable by separase**

The Nasmyth group previously constructed Smc3 whose coiled-coil region could be cleaved by TEV protease (Gruber et al., 2003). For this, they inserted a TEV recognition sequence at amino acid positions 250 and 968, where coiled-coil probability is low and the insertion is therefore less likely to disrupt the coiled-coil structure. To construct Smc3 whose coiled-coil region could be cleaved by separase (Smc3-2R), we inserted a separase recognition sequence (TSLEVGR; Sullivan et al., 2004), instead of a TEV recognition sequence, at the same 250 and 968 amino acid positions (between D250 and G251 and between D968 and F969). As a control, we also inserted a mutated separase recognition sequence (TSLEVGD) at the same sites. The Smc3 coding sequence with the mutations, plus 350 bp upstream and 282 bp downstream sequences, were cloned into an integrative yeast vector pRS405. At the C terminus of the Smc3 open reading frame, six tandem copies of myc epitope tags were inserted. The plasmid constructs were inserted at leu2 locus in the diploid T10954 (MATa/α SMC3/smc3J) strain. Transformed diploid cells were sporulated, tetrad were dissected, and SMc3-2R smc3J and SMc3-2D smc3J haploid cells were obtained. These haploid cells showed a normal growth rate in YPAD media.

**Live-cell imaging and image analysis**

For live-cell imaging yeast cells were immobilized on a glass-bottom dish coated with concanavalin A, and time-lapse images were acquired and analyzed, as described previously (Kalantzaki et al., 2015; Tanaka et al., 2010). Briefly, we used DeltaVision Core and Elite microscopes (GE Healthcare Life Sciences), an UPlanSapo 100 x objective lens (Olympus; numerical aperture 1.40), SoftWoRx software (GE Healthcare Life Sciences), and CoolSnap HQ2 and Cascade II 512B CCD cameras (Photometrics) for image acquisition. At 25°C, we acquired seven to nine (0.7 μm apart) z sections, which were subsequently deconvolved, projected to two-dimensional images and analyzed with Velocity software (Improvision). For Figures, if cells drifted during live-cell imaging, the positions of frames of microscope images were adjusted so that the same cells are in middle of the frames at different time points. The time of completion of chromosome segregation was judged by disappearance of histone signals at the bud neck following their segregation; the disappearance of histone signals at the bud neck was confirmed after enhancing histone fluorescence signals on image analysis software. The completion of cytokinesis was monitored by detection of Myo1 at the bud neck (Wloka and Bi, 2012).

**Western blotting**

To detect myc-, HA-, AID-tagged proteins, acetyl-Smc3 and Cdc28, monoclonal anti-myc (SE11), monoclonal anti-HA (16B12, Covance), polyclonal anti-AID (a gift from Masato Kanemaki lab), monoclonal anti-acetyl-Smc3 (a gift from Katsu Shirahige lab; Borges et al., 2010) and polyclonal anti-Cdc28 (Santa Cruz) antibodies were used, respectively. To detect primary antibodies, HRP-conjugated secondary antibodies (Abcam, GE Healthcare) and IRDye secondary antibodies (LI-COR) were used. The amount of proteins was quantified using software ImageLab 4.1 (Bio-Rad) after blots were scanned using the ChemiDoc imaging system (Bio-Rad).

**Chromatin immunoprecipitation**

Chromatin immunoprecipitation (ChIP), followed by quantitative PCR (ChiP-qPCR) and high throughput DNA sequencing (ChiP-seq), was carried out as described previously (Natsume et al., 2013) with some modification: Cells were incubated with 1% formaldehyde for overnight at 4°C for crosslink. To stop crosslink, glycine was added. Subsequently, cells were washed and lysed, followed by immunoprecipitation using anti-HA antibody (16B12) and magnetic beads (Dynabeads Protein A). For quantitative PCR following ChiP, we used Rotor-Gene 6000 (Corbett) and SYBR Green PCR kit (QIAGEN). Control cells lacking HA tag for SMC1 (T11877) were also analyzed in Figure 2C and showed a low background (IP/input < 0.0003). High throughput DNA sequencing, following ChiP, was carried out at Edinburgh Genomics. The sequence results of ChiP-seq were converted to graphs (Figure S2E) using Integrative Genomic Viewer version 2.4 (y axis is set to 0–1500). To identify peaks in ChiP-seq, MACS version 2.1.0 (Feng et al., 2012) was used to call peaks directly from BAM files, with options “–gsize 12e6–mfold 3 100.” For each detected peak, its position, enrichment with respect to background and statistical significance were calculated.

**Chromosome spreads (chromosome fixation and immobilization)**

Chromosomes were fixed and immobilized on a slide glass, immediately after cell lysis, as described previously (chromosome spreads) (Renshaw et al., 2010; Tanaka et al., 1997). Subsequently, they were processed for immunostaining using mouse monoclonal anti-acetyl-Smc3 (a gift from Shirahige lab; Borges et al., 2010) or anti-HA (16B12) antibody. The primary antibody was visualized by incubation with the secondary antibody goat anti-mouse IgG (Alexa Fluor 647). Chromosome DNA was stained with Hoechst. Spindle Pole Bodies (SPBs) were visualized by expression of Spc42 fused with mCherry. As demonstrated previously (Renshaw et al., 2010), in chromosome spreads, chromosomes showed elongation and the distance between two SPBs was enlarged, when cells proceeded from metaphase to anaphase. Cells in metaphase, in early anaphase and in late anaphase were defined as those in which SPB–SPB distance was < 2.5 μm, 2.5–5.5 μm and > 5.5 μm, respectively. The signals of Smc3-HA or acetyl-Smc3 on chromosomes were quantified using Imaris software (Bitplane).
QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical analyses were carried out using Prism software (Graphpad). Methods of statistic tests are stated in each relevant figure legend. The null hypotheses in these tests were that the samples were collected randomly and independently from the same population. All p-values were two-tailed, and the null hypotheses were reasonably discarded when p-values were < 0.05.

DATA AND SOFTWARE AVAILABILITY

ChIP-seq data have been deposited in the GEO database under the accession number GEO: GSE96841. Original images used in figures have been deposited to Mendeley Data and are available at https://doi.org/10.17632/m64mz2tbvh.1.