Releasing Activity Disengages Cohesin’s Smc3/Scc1 Interface in a Process Blocked by Acetylation


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Releasing Activity Disengages Cohesin’s Smc3/Scc1 Interface in a Process Blocked by Acetylation

Graphical Abstract

Highlights
- Releasing activity disengages cohesin’s Smc3/Scc1 interface
- Releasing activity requires an interaction between the SMC ATPase domains
- Smc3 acetylation inhibits releasing activity

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In Brief
Cohesin entraps sister DNAs within a ring formed by its Scc1, Smc1, and Smc3 subunits. Beckouët et el. have shown that cohesin’s release from the DNA is mediated by disengagement of the Smc3/Scc1 interface. This process involves interaction between the SMC ATPase heads and is inhibited by Smc3 acetylation.

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Releasing Activity Disengages Cohesin’s Smc3/Scc1 Interface in a Process Blocked by Acetylation

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SUMMARY

Sister chromatid cohesion conferred by entrapment of sister DNAs within a tripartite ring formed between cohesin’s Scc1, Smc1, and Smc3 subunits is created during S and destroyed at anaphase through Scc1 cleavage by separase. Cohesin’s association with chromosomes is controlled by opposing activities: loading by Scc2/4 complex and release by a separase-independent releasing activity as well as by cleavage. Coentrapment of sister DNAs at replication is accompanied by acetylation of Smc3 by Eco1, which blocks releasing activity and ensures that sisters remain connected. Because fusion of Smc3 to Scc1 prevents release and bypasses the requirement for Eco1, we suggested that release is mediated by disengagement of the Smc3/Scc1 interface. We show that mutations capable of bypassing Eco1 in Smc1, Smc3, Scc1, Wapl, Pds5, and Scc3 subunits reduce dissociation of N-terminal cleavage fragments of Scc1 (NScc1) from Smc3. This process involves interaction between Smc ATPase heads and is inhibited by Smc3 acetylation.

INTRODUCTION

Sister chromatid cohesion essential for chromosome segregation is mediated by a multisubunit complex called cohesin (Guacci et al., 1997; Michaelis et al., 1997), which contains two SMC proteins, Smc1 and Smc3, and an α-kleisin subunit Scc1. Both Smc proteins form 50-nm-long intramolecular antiparallel coiled coils with a hinge/dimerization domain at one end and at the other an ATPase head domain formed from the protein’s N- and C-terminal sequences. They bind each other via their “hinges” to form V-shaped Smc1/Smc3 heterodimers (Nasmyth and Haering, 2005). Most remarkable is the manner by which the α-kleisin subunit binds to the ATPases at the vertices of this heterodimer. A pair of α-helices within Scc1’s N-terminal domain (NTD) forms a four-helical bundle with the coiled coil emerging from Smc3’s ATPase head (Gligoris et al., 2014), while a winged helix within its C-terminal domain (CTD) binds the base of Smc1’s ATPase, thereby creating a huge asymmetric tripartite ring. Sister chromatid cohesion is thought to be mediated by entrapment of sister DNAs within these rings (Haering et al., 2002), a concept known as the ring model. Bacterial Smc/kleisin complexes also form very similar tripartite rings (Bürmann et al., 2013) that entrap DNAs (Wilhelm et al., 2015), raising the possibility that all Smc/kleisin complexes operate as topological devices.

Coentrapment of sister DNAs within cohesin rings (Gligoris et al., 2014; Haering et al., 2008) takes place during replication and is accompanied by acetylation of a pair of conserved lysine residues within Smc3’s ATPase domain (K112 and K113) by an acetyltransferase called Eco1 (Ivanov et al., 2002; Nasmyth and Haering, 2009). Smc3 acetylation is essential for establishment of stable cohesion. It is maintained throughout G2 and M phases and only removed by a class I deacetylase called Hos1 in yeast and HDAC8 in mammalian cells (Beckouët et al., 2010; Rolef Ben-Shahar et al., 2008; Deardorff et al., 2012) upon cleavage of Scc1 by separase at anaphase onset, an event that opens the ring and destroys the connection between sister DNAs, triggering sister chromatid disjunction (Uhlmann et al., 1999).

Cohesin’s association with DNA, known as cohesin loading, depends on the ability of the ring to hydrolyse ATP bound to Smc1 and Smc3 (Arumugam et al., 2003, 2006), a process facilitated by the activity of a separate complex called Kollerin, which contains the Scc2 and Scc4 proteins (Ciolk et al., 2000). According to the ring model, loading involves passage of DNAs into the ring, which is proposed to take place via a gate created by transient dissociation of the Smc1/Smc3 hinge interface (Gruber et al., 2006). Cohesin rings can entrap in this manner either single DNA molecules or, following replication, a pair of sister DNAs (Gligoris et al., 2014).

Two mechanisms account for cohesin’s release from chromosomes. Best understood is cleavage of its kleisin subunit by separase (Uhlmann et al., 2000). The N- and C-terminal Scc1
fragments associated with Smc3 and Smc1 ATPase heads, respectively (Gruber et al., 2003), are subsequently degraded as daughter cells enter a new cell cycle. Degradation of the C-terminal fragment is mediated by the Ubr1 ubiquitin protein ligase (Rao et al., 2001), but the mechanism responsible for destroying the N-terminal fragment has yet to be elucidated. The second mechanism is independent of separase but requires a regulatory subunit associated with cohesin called Wapl (Gandhi et al., 2006; Kueng et al., 2006). It was initially called the prophase pathway because the process is greatly accelerated in animal cells as they enter mitosis and accounts for the release of most cohesin from chromosomes arms during this stage of the cell cycle.

It turns out that a releasing mechanism related to the prophase pathway operates throughout the cell cycle and is responsible for cohesin’s turnover on interphase chromatin, not only in animal cells (Gerlich et al., 2006) but also in yeast (Chan et al., 2012), where prophase-specific release does not occur and the entire pool of chromosomal cohesin is cleared by separase during anaphase. In addition to Wapl, releasing activity depends on K112 and K113 within Smc3 (in their unmodified state) and on two large regulatory subunits called Pds5 and Sccc3, which bind, respectively, to sequences within the N- and C-terminal halves of Sccc1 (Chan et al., 2013; Hara et al., 2014; Roig et al., 2014; Rowland et al., 2009).

Because it has the potential to destroy sister chromatid cohesion, releasing activity must somehow be neutralized after replication, at least for the chromosomal cohesin pool destined to hold sisters stably together during mitosis, and this is the function of acetylation of K112 and K113 by Eco1. In yeast, this modification appears sufficient to block releasing activity, but in animal cells it requires in addition recruitment of sorotonin (Nishiyama et al., 2010; Rankin et al., 2005). Key to the concept that the function of acetylation is to neutralize releasing activity was the finding that mutations within Wapl, Smc3, Pds5, and Sccc3 that bypass the lethality of eco1 mutants (Rowland et al., 2009; Tanaka et al., 2001) are all defective in releasing activity (Chan et al., 2012). Likewise, inactivation of Wapl bypasses the need for sorotonin in animal cells or Eco1 orthologs in animals (Nishiyama et al., 2010) and plants (De et al., 2014).

If loading is synonymous with entrapment of DNAs within cohesin rings, then release must involve their subsequent escape. The cohesin ring must have an exit as well as an entry gate for DNAs. A clue to the exit gate’s identity was the finding that co-translation fusion of Smc3’s C terminus to Sccc1’s N terminus creates a functional fusion protein that fails to turnover on chromosomes and, like releasing activity mutations, supresses eco1 lethality (Chan et al., 2012). If the exit gate opened by releasing activity were created by transient disengagement of Sccc1’s NTD from Smc3’s coiled coil, then fusion of the two proteins would create a topological barrier to DNA escape.

Though the properties of cohesin rings containing Smc3-Sccc1 fusion proteins are clearly consistent with releasing activity working by disengaging Sccc1 from Smc3, it does not address the key issue of whether or not releasing activity actually opens this interface in living cells. Under normal circumstances, this may be difficult to measure, as disengagement is presumably a fleeting process that is swiftly followed by re-engagement of Sccc1, whose CTD remains attached to Smc1. However, if releasing activity also operated on the N-terminal cleavage fragments (NScc1) created by separase, disengagement might lead to a permanent and thereby more readily measurable dissociation. We show here that this is indeed the case, namely that releasing activity is responsible for removing NScc1 from Smc3’s coiled coil, that this process likely involves engagement of Smc1 and Smc3 ATPases, and that the process is blocked by acetylation of Smc3’s K112 and K113 residues. Our findings constitute direct evidence that separase-independent release involves creation of a gate at the Smc3-Sccc1 interface from which previously entrapped DNAs can escape.

**RESULTS**

**NScc1 Is Stabilized by Releasing Activity Mutations**

We previously demonstrated that Smc3 is deacetylated by Hos1 in response to Sccc1 cleavage at the onset of anaphase (Beck-ouët et al., 2010). If releasing activity disengages Sccc1 from unacetylated Smc3, and if it also does so after Sccc1 has been cleaved, then the activity should detach irreversibly NScc1 from deacetylated Smc3, and this might stimulate proteolysis. To address this, wild-type and wpl1Δ cells arrested in metaphase by Cdc20 depletion were triggered to undergo anaphase and enter G1 (by reinduction of Cdc20). Western blot analysis revealed that wpl1Δ greatly delayed degradation of N- (Figure 1A) but not C-terminal fragments (see Figure S1A available online). This effect is not due to loss of Wapl per se, because it was also observed in mutants known to be defective in releasing activity, namely pds5S81R and sccc3E202K (Rowland et al., 2009) (Figures 1B and 1C). Note that Wapl is still recruited to chromosomal cohesin complexes containing Sccc3 E202K protein, and yet the mutation reduces NScc1 degradation as much as wpl1Δ.

**Releasing Activity Promotes Dissociation of NScc1 from Smc3**

Releasing activity might promote NScc1 degradation either directly, by interacting for example with the relevant proteolytic machinery, or indirectly, by promoting the fragment’s release from Smc3, which is a precondition for degradation. To measure the effect of releasing activity on disengagement of NScc1 from Smc3 per se, it is necessary to uncouple disengagement from proteolysis. This might be possible if proteolysis were a cell-cycle-dependent event and only occurred as cells undergo anaphase and enter G1. We therefore compared NScc1’s fate in wild-type and wpl1Δ cells when generated by induction of tobacco etch virus (TEV) protease in G2/M phase-arrested cells whose Sccc1 contained a TEV instead of a separase cleavage site at position 268 (Figure 1D). Interestingly, little or no degradation occurred in either wild-type or wpl1Δ mutant cells, at least during a 90 min window following cleavage (Figure 2A). NScc1 proteolysis is either an event associated only with separase cleavage or more likely one that only occurs from anaphase to G1.

Having discovered a condition in which NScc1 is stable, we were in a position to test whether Wapl influences dissociation. To do this, we used a version of Smc3 with a functional cysteine substitution within its coiled coil (S1043C) that can be efficiently crosslinked to a natural cysteine within Sccc1’s NTD (C56) (Figure 1D) in living cells using the homobifunctional sulfhydryl active reagent Bis-maleimidoethane (BMOE) (Gligoris et al., 2014). In cells arrested in G2/M by nocodazole, the presence or absence
In wpl1Δ mutant but not wild-type cells, 1–181 NScc1 (created by separase; Figure 1D) accumulates to high levels (Figure 2A). Moreover, it is crosslinked to Smc3 with an efficiency similar to FL Scc1 (Figure 2B), implying that NScc1 remains bound to Smc3 long after its creation at the previous anaphase. Induction of TEV protease in G2/M phase cells triggered cleavage of FL protein, creating similar amounts of 1–268 NScc1 (Scc1 TEV) in wild-type and mutant cells (Figure 2A). Despite this equality, treatment with BMOE induced more efficient crosslinking between Smc3 S1043C and C56 within 1–268 NScc1 in wpl1Δ than in wild-type (Figure 2B), implying that Wapl promotes dissociation. To exclude the possibility that this is due to a conformational change in the interaction between Scc1 and Smc3 rather than disengagement per se, we measured the amount of 1–268 NScc1 in immunoprecipitates of Smc3 from cells untreated with BMOE (Figure 2C). This confirmed that greater amounts of NScc1 remained associated with Smc3 in wpl1Δ than in wild-type cells (Figure 2C). Taken together, these data imply that releasing activity is required to disengage NScc1 from Smc3’s coiled coil. Note that because cleavage either by TEV or separase induces Smc3 de-acetylation, NScc1’s disengagement from Smc3 due to releasing activity must be from unacetylated Smc3 molecules.

Because most cohesin in G2/M is associated with chromosomes, NScc1 disengagement upon TEV cleavage could take place either shortly before or after cleaved complexes are released from chromatin. Indeed, if transient disengagement drives cohesin’s release from chromatin, then it must take place within complexes associated with chromatin. To address whether releasing activity also acts on cohesin whose cleavage had previously triggered dissociation from chromatin, we used BMOE-induced crosslinking to address the fate of separase-created 1–181 NScc1 in G2/M phase cells whose WPL1 gene is under control of the galactose-inducible GAL1-10 promoter (GAL-WPL1). In G2/M phase-arrested cells grown in the absence of galactose (i.e., without Wapl), 1–181 NScc1 accumulates and can be efficiently crosslinked to Smc3. Importantly, live imaging demonstrated that activation of separase at the metaphase to anaphase transition induces cohesin’s removal from chromatin even in the absence of Wapl (data not shown), so that the cohesin complexes containing NScc1 (but not Csc1, which will have been degraded by Ubr1) will be nucleoplasmic and not associated with chromatin.

Induction of Wapl by galactose had no effect on crosslinking between Smc3 S1043C and C56 within FL Scc1 but greatly reduced it within 1–181 NScc1 (Figure 2D). The amount of separase-generated 1–181 NScc1 immunoprecipitated with Smc3 in the absence BMOE also declined upon Wapl induction (Figure S1B). We conclude that Wapl promotes disengagement of NScc1 from Smc3 in soluble cohesin complexes that in all probability lack Scc1’s C-terminal cleavage fragment. A corollary is that although releasing activity requires Scc3, it does not require the latter’s association with its known Scc1 binding site, which will have been destroyed by Ubr1 (Roig et al., 2014).

**Imaging Wapl-Dependent NScc1 Dissociation**

To observe disengagement in living cells, we integrated 448 tetracycline operators (TetO) between the BMH1 and PDA1 genes in haploid or diploid WT cells that express both Scc3

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**Figure 1. Stability of Scc1 Cleavage Fragments in Releasing Activity Mutants**

(A–C) Wild-type (K17960), wpl1Δ (K20236), pds5-S81R (K20521), and Sc33-E202K (K20526) strains expressing CDC20 from the GAL promoter were grown to logarithmic phase at 25°C in YP medium containing galactose, transferred to galactose-free media to induce metaphase arrest (time 0), and anaphase triggered by galactose readdition. Separase cleavage of Scc1 was followed by western blotting, detecting N-terminal Myc tag on Scc1.

(D) Model of the ATPase domains of Smc1 and Smc3 in an engaged state driven by ATP binding. The separase cleavage site in Scc1 at position 181 is marked with a black asterisk; TEV sites at position 268 are marked with a red asterisk.

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of Wapl had little or no effect on BMOE-induced crosslinking between full-length (FL) Scc1 C56 and Smc3 S1043C proteins (Figure 2B). The lack of effect is unsurprising. In such cells, a large fraction of Smc3 is acetylated and should not, therefore, be subject to releasing activity. Moreover, it is doubtful that our crosslinking assay would detect transient disengagement of FL Scc1 even within unacetylated complexes.
and mCherry fused to the tetracycline repressor (Scc3-TetR and TetR-mCherry). Scc1 or Smc3 tagged with GFP at their C termini expressed in this strain colocalized with TetR-mCherry, indicating that the cohesin ring is specifically tethered at the TetO arrays by Scc3-TetR, which binds to the C-terminal half of Scc1. Interestingly, in both cases enrichment of GFP at TetO arrays was detectable not only in anaphase but also in telophase cells (Figures 3A and 3B), implying that separase cleavage does not provoke immediate removal/destruction of CScC1 bound to Smc1/Smc3 heterodimers. Unlike GFP attached to Scc1’s C terminus, GFP attached to its N terminus was never observed in telophase cells (Figure 3C). In contrast, it was invariably observed at the arrays in telophase wpl1Δ cells. We conclude that releasing activity removes most if not all NScc1 from Smc3 within 10 min of cleavage, a process more rapid than CScc1 degradation.

**Scc3’s Highly Conserved Surface Is Essential for Release, Not Loading**

If cohesin’s release from chromosomes is mediated by the transient disengagement of Scc1’s NTD from Smc3, and if dissociation of NScc1 is a valid measure of the latter, then all mutations known to inactivate releasing activity and capable of suppressing null alleles of eco1 should be defective in removing NScc1 from Smc3. We addressed first the role of cohesin’s regulatory subunit Pds5. Nonlethal mutations within the N-terminal domain of Pds5, for example pds5S81R, enable cells to proliferate in the absence of Eco1, greatly reduce cohesin’s turnover on chromosomes (Chan et al., 2012), and delay NScc1 degradation (Figure 1B). To address whether pds5S81R blocks its dissociation from Smc3, we analyzed the effect on BMOE-induced crosslinking between Smc3 S1043C and Scc1 C56. Figure 4C shows that the mutation greatly increases the crosslinking, implying that it does indeed delay dissociation.

We next addressed the role of Scc3. scc3E202K delays NScc1 degradation, suggesting that this mutation also delays NScc1’s
However, the pocket affected by this mutation is not particularly conserved, and we therefore turned our attention to the role of a highly conserved surface that sits underneath Scc3’s prominent nose (Figure 4A) (Hara et al., 2014; Roig et al., 2014). Given its extreme conservation, we expected that it participates in an essential process, such as cohesin loading. Surprisingly, substitution by alanine of seven highly conserved surface residues within the domain (Scc3 K364A, Y371A, K372A, T401A, K404A, W408A, R449A) is not lethal (Figures S2A and S2C). However, we observed that the 7A mutation affects releasing activity, as it permits proliferation of cells lacking Eco1 (Figure S2B). To evaluate the role of individual residues, we created a series of single mutations (K404E, K404A, Y405E, and Y405A) and used tetrad analysis to evaluate their ability to suppress lethality associated with eco1D. Of these, K404E suppressed lethality, and did so as efficiently as wap1D (Figures 4B and S2B). Gel filtration showed that K404E affects binding between purified FL Scc3 and Wapl proteins (Figure 4D), though it does not eliminate the interaction. Interestingly, scc3K404E...
(and the 7A mutant) does not affect accumulation of GFP tagged Wapl within pericentric chromatin in living cells (Figure S2C), implying that Wapl can still be recruited to chromosomal cohesin complexes. Crucially, scc3K404E caused an increase in NScc1-Smc3 crosslinking comparable to that caused by pds5S81R (Figure 4C). Remarkably, the very same mutation rescued a temperature sensitive allele of scc3K404E at 30°C, raising the possibility that Eco1 might have an essential target in addition to Smc3’s K112 and K113 residues.

An alternative explanation is that scc3M102K causes only a partial loss of releasing activity; that substitution of K112 and K113 by arginine also reduces releasing activity, though not below a level sufficient to restore viability; and that the combination of smc3K112R K113R and scc1M102K reduces releasing activity further, to a level compatible with proliferation in the absence of Eco1. Consistent with this scenario, scc1M102K suppressed lethality caused by eco1D in smc3K112R K113R cells (Figure S3B). Indeed, the presence or absence of ECO1 had little or no effect on proliferation of smc3K112R K113R scc1M102K double mutants, suggesting that K112 and K113 may after all be the only essential targets of Eco1.

Crucially, scc1M102K had little or no effect on the degree of crosslinking between NScc1 and an ectopically expressed epitope tagged Smc3 S1043C protein but caused a marked increase with Smc3 S1043C K112R K113R (RR) (Figures 5A and 5C). Note that smc3K112R K113R also had little effect in cells containing a wild-type SCC1 gene. These results identify mutations within the interface between Smc3 and Scc1’s NTD that affect both releasing activity and dissociation of NScc1 from Smc3’s coiled coil. Our findings that scc1M102K only causes a decrease in releasing activity sufficient to suppress eco1D
when combined with smc3K112R K113R and that both mutations are required to eliminate NScc1's dissociation from Smc3 further strengthens the correlation between these two phenomena and raises the possibility that NScc1 dissociation is actually responsible for cohesin's release from chromatin. How scc1M102K affects NScc1's dissociation from Smc3 at the hands of Wapl, Pds5, and Scc3 is presently unclear.

### Role of Residues within Smc3's ATPase Domain

To probe further the relationship between releasing activity responsible for eco1 mutant lethality and NScc1 dissociation, we measured the effect of mutations within Smc3's ATPase. Smc3 R107 is a highly conserved arginine within the upper of three parallel β sheets that underlie the loop containing K112 and K113 (Figure 5B). It faces inside the ATPase domain and mutation to either isoleucine or alanine enables robust proliferation of eco1J cells (Rowland et al., 2009). Figure 5B shows that, like scc3, wpl1, and pds5 suppressors, smc3R107I also blocks NScc1 dissociation as measured by crosslinking between Smc3 S1043C and Scc1 C56.

We next analyzed the effect of an smc3 mutation at the base of Smc3's ATPase domain (smc3D1189H) isolated not as an eco1J suppressor but by virtue of its ability to suppress the sensitivity to benomyl of wpl1J eco1J double mutants (Guacci et al., 2015). Like scc1M102K, smc3D1189H suppresses lethality associated with smc3K112R K113R, but not eco1J (Figure 5C). The finding that smc3D1189H causes only a modest increase in crosslinking between Smc3 S1043C and Scc1 C56 but that this effect is greatly increased when K112 and K113 are replaced by arginine. As in the case for scc1M102K, smc3K112R K113R enabled smc3D1189H to suppress lethality associated with eco1J (Figure S4A). The finding that smc3D1189H suppresses the lethality of smc3K112R K113R but not eco1J led to the suggestion that Eco1 must therefore have targets besides K112 and K113 (Guacci et al., 2015). Our data suggest an alternative and more plausible explanation, namely that cohesin complexes containing smc3K112R K113R have less releasing activity than unacetylated wild-type complexes. Thus, the properties of smc3D1189H do not imply that Eco1 has targets besides K112 and K113. The behavior of smc3D1189H reveals again an exquisite correlation between eco1J suppression and defective NScc1 dissociation. These findings have an interesting corollary. If one assumes that rescue by smc3D1189H of the benomyl sensitivity of wpl1J eco1J double mutants (Guacci et al., 2015) is due to a defect in releasing activity, then residual releasing activity must persist in wpl1J mutants.

### Smc3 Acetylation Blocks NScc1 Dissociation

To address whether Smc3 acetylation blocks disengagement of Scc1's NTD from Smc3, we asked whether the decline in crosslinking between NScc1 and Smc3 S1043C upon induction of Wapl with GAL-WPL1 depends on Smc3's deacetylation by Hos1. Figure 6A shows the decline is less pronounced in hos1J cells than in wild-type. In other words, NScc1's dissociation upon Wapl reactivation depends on the latter's prior deacetylation at the time of separase cleavage. The failure to remove NScc1 from Smc3 in hos1 mutant cells might contribute to the inability of Smc3 molecules that remain acetylated after anaphase to build cohesin during the subsequent cell cycle (Beckouët et al., 2010).

Another way of addressing whether acetylation blocks disengagement of NScc1 from Smc3 is to analyze the effect of mutations such as smc3K112Q K113Q that are thought to mimic acetylation. It has hitherto not been possible to address the effect of smc3K112Q K113Q on releasing activity directly, because the mutation causes a major decline in cohesin's loading throughout the genome (Hu et al., 2015). Without loading, it is not possible to measure release. Our finding that NScc1 dissociation appears to be a perfect surrogate for release solves this conundrum. All that is required is that Scc1 associated with Smc3 K112Q K113Q be cleaved by separase. Fortunately, this proves to be the case, and Figure 6B shows that smc3K112Q K113Q elevates crosslinking between NScc1 and Smc3S1043C to a degree comparable to wpl1J.

### Disengagement of NScc1 from Smc3 Involves Engagement of Smc ATPase Heads

To address the role of ATP hydrolysis, we used BMOE-induced crosslinking between Smc3 S1043C and Scc1 C56 to measure the effect of Smc3 E1155Q (Arumugam et al., 2003) on dissociation from Smc3 of NScc1 created in nocodazole-arrested cells by TEV protease. Figure 7A shows that crosslinking between Scc1 C56 and Smc3 S1043C is little affected by Smc3E1155Q, either when NScc1 is created by separase during the previous cell cycle or by TEV protease during nocodazole-induced G2/M phase arrest. This implies that the EQ mutation does not prevent dissociation. To confirm this, we compared the effect of inducing Wapl in GAL-WPL1 eco1J cells on BMOE-induced crosslinking between NScc1's C56 and Smc3 S1043C and Smc3 E1155Q S1043C. Interestingly, the smc3E1155Q mutation again had little or no effect (Figure S4B). Further confirmation that releasing activity dissociates NScc1 from Smc3 E1155Q ATPases is our finding that smc3K112Q K113Q causes a significant increase in NScc1 C56 crosslinking to Smc3 E1155Q S1043C (Figure 7B), suggesting that the dissociation reaction in Smc3 E1155Q complexes is affected by acetylation of K112 and K113.

We were unable to address the role of ATPase head engagement by analyzing mutations like smc3K39I, which affects binding of ATP to the Smc3 ATPase heads, or the signature motif mutation smc3S1128R, which compromises engagement with Scc1 ATPase heads (Arumugam et al., 2006), because neither Smc3 K39I nor Smc3 S1128R proteins accumulate within nuclei (data not shown) and may not even interact with Wapl, which is a nuclear protein. Nevertheless, in the course of our studies, a novel class of eco1J suppressor mutations has been discovered, namely smc3D1164E, which alters a key amino acid within the Smc1 ATPase's D loop, and smc1L1129V, which alters its signature motif (Figure S4C). These mutations cause a severe reduction in ATPase activity (without affecting ATP binding) but cause only a mild reduction in the amount of cohesin loaded throughout the genome (Elbatsh et al., 2016, in this issue of Molecular Cell). To test whether smc1D1164E or smc1L1129 affects dissociation of NScc1 from Smc3, we measured their effects on crosslinking between Smc3 S1043C and C56 from a version of Scc1 whose N terminus was tagged with multiple...
myc epitopes. We used this method instead of using an epitope tag on Scc1’s C terminus (as performed in all previous experiments), because tetrad analysis revealed that the latter was synthetic lethal with both smc1D1164E and smc1L1129V (data not shown). As shown in Figures 7C and S4C, both mutations increased crosslinking between NScc1 and Smc3. The equivalent residues within Smc3 are D1161 and L1126. smc3L1126V is viable despite reducing ATPase activity and cohesin’s association with chromosomes, but it fails to suppress the temperature sensitivity of an eco1ts strain (Elbatsh et al., 2016) and galactose-inducible WPL1 gene was grown at 25°C, arrested in G1 by pheromone, and permitted to go through S phase at 37°C in the presence of nocodazole. After addition of either galactose or glucose to induce Wapl expression (or not) samples were taken at times 0 and 60 min for in vivo crosslinking with 5 mM BMOE. Catenated monomers (CM), catenated dimers (CD).

Though our crosslinking assay is designed to measure NScc1 dissociation, it also reveals crosslinking between Smc3 S1043C and C56 from intact Scc1 molecules. We noticed that several mutations that affect ATP hydrolysis, in particular smc3E1155Q and possibly also smc3D1189H, appear to increase the efficiency of Scc1-Smc3 crosslinking. We do not understand the significance of this effect. Crucially, it does not correlate with smc3E1155Q on NScc1 dissociation implies that only part of a single ATP hydrolysis cycle is necessary. Though ATP hydrolysis per se is not required, an early event following the cooperative binding of ATP to Smc heads appears necessary. Abolition of releasing activity by smc1L1129V, but not by smc3L1126V, a pair of mutations that have equally severe effects on ATP hydrolysis and cohesin loading, suggests that the nature of ATPase head engagement required for release differs fundamentally from that required for loading. There may be more than one state of head engagement.

Though our crosslinking assay is designed to measure NScc1 dissociation, it also reveals crosslinking between Smc3 S1043C and C56 from intact Scc1 molecules. We noticed that several mutations that affect ATP hydrolysis, in particular smc3E1155Q and possibly also smc3D1189H, appear to increase the efficiency of Scc1-Smc3 crosslinking. We do not understand the significance of this effect. Crucially, it does not correlate with
defects in NScc1 dissociation, and we therefore presume that it is immaterial to our measurements of NScc1 cleavage fragment crosslinking.

The Mechanism by which Acetylation Blocks Release

The finding that NScc1 release depends on the precise form of interaction between Smc1 and Smc3 ATPase heads raises the possibility that acetylation blocks release by regulating this process. For the lack of an assay to measure directly the state of interaction between Smc1 and Smc3 ATPase heads in vivo, we investigated whether the chemical status of K112 and K113 affects the behavior of Smc3 E1155Q cohesin complexes. The inability of these complexes to hydrolyze ATP stabilizes head engagement and prevents cohesin’s translocation into pericentric sequences from its loading sites at core centromeres (Hu et al., 2011, 2015). These complexes are nevertheless capable of releasing NScc1 from Smc3. If acetylation blocked a step prior to that blocked by E1155Q, for example a certain type of head engagement, then K112Q K113Q should block the ability of acetylation to dissociate the Smc3 head engagement necessary for association of Smc3E1155Q with its centromeric loading sites, raising the possibility that it blocks release via a related mechanism.

Releasing Activity Induces Exit of DNAs from Cohesin Rings

To prove that releasing activity does actually induce DNAs to escape entrapment by cohesin rings, we created an eco1-1(G211H) GAL-WPL1 yeast strain with cysteine pairs at all three interfaces (hence 6C) making up the cohesin ring (Gligoris et al., 2014). Cells (containing a circular 2.5 kb minichromosome) growing at 25°C were arrested in G1 by pheromone and then permitted to undergo S phase at 37°C (the restrictive temperature for eco1-1(G211H)) in the presence of nocodazole. The G2/M-arrested cells were treated with BMOE and DNAs associated with 6C crosslinked cohesin immune-precipitated, heated in the presence of SDS at 65°C before gel electrophoresis, and minichromosome DNA detected by Southern blotting. The

Figure 7. Disengagement of NScc1 from Smc3 Requires a Single Round of ATP Hydrolysis

(A) Strains K23070 (MATα SMC3 URA3::SMC3 S1043C-HA6 MYC3-SCC1(TEV268) YEp-PGAL1-TEV), K23067 (MATα SMC3 URA3::smc3 S1043C E1155Q-HA6::URA3 MYC3-SCC1(TEV268) YEp-PGAL1-TEV), and K23068 (MATα SMC3 SMC3 S1043C-HA6 MYC3-SCC1(TEV268) wpl1Δ YEp-PGAL1 TEV) were treated and analyzed as described in Figure 2B.

(B) Strains K24217 (MATα SMC3 URA3::SMC3 S1043C-PK6::URA3 MYC3-SCC1(TEV268)), K24218 (MATα SMC3 URA3::smc3K112 K113Q S1043C-PK6 MYC3-SCC1(TEV268)), and K24219 (MATα SMC3 URA3::smc3E1155Q S1043C-PK6 MYC3-SCC1(TEV268)) were grown and analyzed as described in Figure 6B.

(C) Exponentially growing strains K23070, K23068, K24911 (SMC3smc3L1126VS1043C::LEU2MYC3-SCC1(TEV268)) and K24523 (SMC3 S1043C::ADE2 MYC3-SCC1(TEV268) smc1 L1129V) were treated as described in Figure 6B and analyzed by western blotting using anti-MYC antibodies.

(D) Shown is a model for how releasing activity dissociates Scc1-NTD from Smc3’s coiled coil leading to escape of entrapped DNAs in a process involving ATP-dependent engagement of SMC ATPase heads. Acetylation of Smc3 residues K112 and K113 is suggested to inhibit ATP-dependent head engagement.
absence of Wapl permits such cells to build cohesion, and this is apparent in the formation of sister DNAs catenated by cohesin rings containing BMOE-induced crosslinks at all three interfaces (catenated dimers or CDs). Monomeric DNAs catenated by covalently closed cohesin rings (CMs) are also apparent. Crucially, addition of galactose to the G2/M phase cells, which restores Wapl to cells lacking Smc3 acetylation, causes a rapid decline of both CDs and CMs as well as the amount of monomeric DNAs merely coprecipitated with cohesin (Figure 6C). A modest decline in CDs, but not CMs, was also observed in the absence of galactose, possibly due to the extended incubation at 37°C. We conclude that releasing activity does indeed catalyze escape of DNAs from inside cohesin’s ring.

DISCUSSION

The kinetics of cohesin’s association with chromosomes is determined by loading mediated by the Scc2/4 kollerin complex and release mediated by cohesin’s Wapl, Pds5, and Scc3 subunits. If loading involves entrapment of DNAs within cohesin’s ring, then release must involve their subsequent escape through an exit gate. We describe here definitive evidence that DNAs are indeed induced to escape in this manner in living cells (Figure 7D). The observation that cohesin containing an Smc3-Scc1 fusion protein loads but cannot dissociate suggests that the exit gate is situated at the interface between Scc1’s NTD and the coiled coil emerging from Smc3’s ATPase. However, this is not proof. Fusion of Smc3 to Scc1 could conceivably block release not by creating a topological barrier to DNA exit but instead by altering the ATPase’s conformation, which could have an indirect effect on release via an exit gate situated elsewhere. It was vital, therefore, to address whether or not cohesin’s releasing activity does indeed disengage Scc1’s NTD from Smc3.

The technology we adopted is chemical crosslinking induced by the bifunctional thiol-specific reagent BMOE, which induces within minutes efficient crosslinks between Smc3’s S1043C within Smc3’s coiled coil and a natural cysteine (C56) in Scc1’s NTD. We suspect that our failure to discern any effect of releasing activity on the efficiency of crosslinking within intact cohesin complexes can be attributed to release being a transient process. When Scc1 NTDs reassociate with Smc3, they will be again subject to crosslinking, which will subsequently trap them in the associated state. Our discovery that degradation of Scc1’s N-terminal separase cleavage fragment is largely abolished by mutations that inactivate releasing activity provided a solution to this conundrum. It led to the realization that dissociation could be made irreversible and hence observable by measuring crosslinking between Smc3’s coiled coil and N-terminal Scc1 cleavage fragments (NScc1), be they created naturally during anaphase by separase or artificially by inducing cleavage with TEV protease.

Using this assay, we have analyzed a wide variety of mutations known to have severe defects in releasing activity (because they permit proliferation in the absence of Eco1). Some of these mutations have been described previously while others are described in this or the accompanying paper (Elbatsh et al., 2016). In every single case where mutations suppress eco1Δ, they also reduce NScc1 dissociation. Striking examples are scc1M1022K and smc3D1189H, which only suppress eco1Δ when the residues within Smc3’s ATPase domain (K112 and K113) that must normally be acetylated by Eco1 are replaced by arginine. Smc3K112R K113R not only prevents acetylation but also compromises releasing activity, not to an extent that avoids lethality, but to an extent that exacerbates defects caused by scc1M1022K and smc3D1189H.

The strength of this correlation means that it is hard to avoid the conclusion that releasing activity does indeed induce dissociation of the interface between Scc1’s NTD from Smc3’s coiled coil. The previous finding that fusion of Smc3’s C terminus to Scc1’s N terminus also abrogates releasing activity and suppresses eco1Δ lethality implies that dissociation of Scc1 from Smc3 is not only an intrinsic aspect of releasing activity but is also required for this phenomenon. Both sets of observations are consistent with the notion that cohesin loading involves entrapment of DNA inside cohesin rings and that release from chromosomes is mediated by their escape through a gate created by dissociation of Scc1’s NTD from Smc3’s coiled coil.

Our conclusion that cohesin’s release from chromosomes is mediated by dissociation of NSc1 from Smc3’s coiled coil has an important ramification. Hitherto, it has not been possible to measure directly the effect of K112 and K113 acetylation by Eco1 on release, as modification of these residues interferes with loading as well as release. Our finding that releasing activity-dependent NScc1 dissociation can also be observed within cohesin complexes that cannot load onto chromosomes means that it has been possible to measure the effect of smc3K112Q K113Q, which is presumed to mimic the acetylated state. Crucially, this double mutant appears to abrogate dissociation of NScc1 from Smc3 as severely as all known releasing activity mutations, implying that K112 and K113 acetylation does indeed neutralize releasing activity, as has long been suspected. Our finding that the Hos1 deacetylase is important for NScc1’s dissociation from Smc3 during anaphase confirms that this conclusion applies equally well to physiological acetylation.

The mechanism by which Scc1’s N-terminal domain is dissociated from Smc3’s coiled coil is clearly a complex process, as it depends on Wapl, Pds5, Scc3, residues throughout Smc3’s ATPase head; as well as the D loop that is the target of Eco1; and on residues within the D loop and signature motif of Smc1’s ATPase head. The latter is a crucial discovery, as it implies that ATP-driven engagement of Smc1 and Smc3 ATPase heads is involved. In this regard, the process shares properties with the cohesin loading reaction. Indeed, it is tempting to speculate that acetylation blocks both loading and release, because it blocks ATPase head engagement, a proposal consistent with our finding that smc3K112Q K113Q greatly reduces association of Smc3E1155Q with centromeric loading sites. We have previously suggested that the acetylation state of the KKD loop controls the switch between Smc3’s ATPase binding pocket via R61, which sits on top of a short z-helix connecting these structures. Smc3 R61Q is lethal and compromises cohesin loading, and it will be interesting to establish whether the mutation also abolishes releasing activity. We acknowledge that the lack of any effect of smc3K105Q K106Q mutations on the ATPase activity of purified human trimeric Smc1/Smc3/Scc1 complexes (Ladurner...
et al., 2014) appears contrary to our proposal that acetylation affects the state of head engagement. We note, however, that these assays were performed in the absence of Scc3, Pds5, or Wapl, which might alter the properties of cohesin’s ATPase heads. Moreover, the role of acetylation of vertebrate cohesin is presently unclear. While acetylation is a prerequisite for Sororin association, it is not sufficient to counteract releasing activity (which requires Sororin association to acetylated cohesin). It is therefore conceivable that any effect on the ATP hydrolysis cycle might manifest only after Sororin association.

It is nevertheless important to point out that loading is not greatly affected by wpl1, pds5, scc3, and smc3 mutations that greatly reduce release. Likewise, fusion of Smc3 to Scc1, which abrogates release, still permits cohesin loading not only in yeast (Gruber et al., 2006; Haering et al., 2008) but also in Drosophila (Eichinger et al., 2013). Thus, while opening of the Scc1-Smc3 interface is an obligate aspect of release, it is not an obligate aspect of loading, a fact difficult if not impossible to reconcile with the suggestion that this same interface is also cohesin’s main DNA entry gate (Murayama and Uhlmann, 2015). Indeed, there is evidence that cohesin’s DNA entry gate is situated instead at the Smc1/Smc3 hinge interface (Gruber et al., 2006).

We conclude that though release and loading share certain properties, such as involvement of an intermediate involving head engagement, they are nevertheless distinct processes with very different outcomes for the state of cohesin’s association with chromatin. We speculate that release involves an intermediate in which DNA previously trapped inside heterotrimeric rings is ejected from the lumen created by engagement of Smc1 and Smc3 ATPase heads, permitting its escape when this subsequently triggers disengagement of the Scc1-Smc3 interface (Figure 7D).

There is now incontrovertible evidence that the sister DNAs of circular minichromosomes can be entrapped by heterotrimeric cohesin rings, a phenomenon that is currently the only plausible explanation for how cohesin holds sister chromatids together. How DNAs enter cohesin rings remains very unclear. This paper provides a plausible pathway for its subsequent exit, namely through a gate created by disengagement of the Scc1-Smc3 interface. The existence of this pathway constitutes therefore an important endorsement of the notion that cohesin and its relatives do indeed act as topological devices, as originally proposed by the ring model. Lastly, our finding that releasing activity opens the interface between Scc1 and Smc3, with the implication that this creates a gate for DNA to exit cohesin rings, suggests that topological entrapment is a universal mechanism for cohesin’s stable or semistable association with chromatin fibers and not merely one that applies to minichromosomes, which hitherto is the only instance where this has been directly demonstrated.

**EXPERIMENTAL PROCEDURES**

**Yeast Cell Culture**
All strains are derivatives of W303 (K699). Strain numbers and relevant genotypes of the strains used are listed in the figure legends. Details of strain construction are in Supplemental Experimental Procedures. Cells were cultured at 25°C in YEP medium with 2% glucose unless stated otherwise.

**Sequence Alignment**
For the multiple alignment conservation, the following sequences were included: Zygosaccharomyces rouxii (C5DW3M), Saccharomyces cerevisiae (P40541), Ashbya gossypii (MMYD6), Homo sapiens (Q6P275), Xenopus laevis (Q9DG1N), Danio rerio (B0V0X2), Drosophila melanogaster (Q9VM62), Daphnia pulex (E9FY68), Brugia malayi (A8QED2), Vitis vinifera (D7TP60), Candida albicans (C4YFQ5), Schizosaccharomyces pombe (O13816), and Sordaria macrospora (F7W0E2).

**In Vivo Chemical Crosslinking and Minichromosome IP**
In vivo chemical crosslinking and minichromosome IP were performed as described by Gilgors et al. (2014) and as detailed in the Supplemental Experimental Procedures.

**Live-Cell Imaging**
Exponentially growing cells were placed on 2.5% agarose pads made of synthetic complete medium plus glucose. Live-cell imaging was performed under a spinning disk confocal system (PerkinElmer UltraVIEW) with an EMCCD camera (Hamamatsu) mounted on an Olympus IX81 microscope with Olympus 100N.A. objectives. Image acquisition was done at 25°C. Seventeen to twenty-one z-stacking images were acquired, and image deconvolution was done by using Velocity software with seven iterations and 95% confidence. Fresh samples were prepared every 10 min.

**Calibrated ChIP-Sequencing**
Calibrated ChIP-seq was performed as detailed in (Hu et al., 2015).

**ACCESSION NUMBERS**
The sequencing data have been deposited in GEO with the accession number GSE76890.

**SUPPLEMENTAL INFORMATION**
Supplemental Information includes four figures and Supplemental Experimental Procedures and can be found with this article at http://dx.doi.org/10.1016/molcel.2016.01.026.

**AUTHOR CONTRIBUTIONS**
F.B., M.S., M.B.R., and K.-L.C. designed and conducted the experiments. J.G.S., P.B., B.H., N.P., T.G., A.G.S., and L.S. conducted the experiments. K.N. designed the experiments and wrote the paper.

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