Sister Chromatid Cohesion

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During S phase, not only does DNA have to be replicated, but also newly synthesized DNA molecules have to be connected with each other. This sister chromatid cohesion is essential for the biorientation of chromosomes on the mitotic or meiotic spindle, and is thus an essential prerequisite for chromosome segregation. Cohesion is mediated by cohesin complexes that are thought to embrace sister chromatids as large rings. Cohesin binds to DNA dynamically before DNA replication and is converted into a stably DNA-bound form during replication. This conversion requires acetylation of cohesin, which in vertebrates leads to recruitment of sororin. Sororin antagonizes Wapl, a protein that is able to release cohesin from DNA, presumably by opening the cohesin ring. Inhibition of Wapl by sororin therefore “locks” cohesin rings on DNA and allows them to maintain cohesion for long periods of time in mammalian oocytes, possibly for months or even years.

DNA replication during the synthesis (S) phase generates identical DNA molecules, which, in their chromatinized form, are called sister chromatids. The pairs of sister chromatids remain united as part of one chromosome during the subsequent gap (G2) phase and during early mitosis, in prophase, prometaphase, and metaphase. During these stages of mitosis chromosomes condense, in most eukaryotes the nuclear envelope breaks down, and in all species chromosomes are ultimately attached to both poles of the mitotic spindle. Only once this biorientation has been achieved for all chromosomes, the sister chromatids are separated from each other in anaphase and transported toward opposite spindle poles of the mother cell, enabling its subsequent division into two genetically identical daughter cells.

This series of events critically depends on the fact that sister chromatids remain physically connected with each other from S phase until metaphase. This physical connection, called sister chromatid cohesion, opposes the pulling forces that are generated by microtubules that attach to kinetochores and thereby enables the biorientation of chromosomes on the mitotic spindle (Tanaka et al. 2000b). Without cohesion, sister chromatids could therefore not be segregated symmetrically between the forming daughter cells, resulting in aneuploidy. For the same reasons, cohesion is essential for chromosome segregation in meiosis I and meiosis II. Cohesion defects in human oocytes can lead to aneuploidy, which is thought to be the major cause of spontaneous abortion, because only a few types of aneuploidy are compatible with
viability, such as trisomy 21 (Down syndrome), trisomy 18 (Edwards syndrome), and trisomy 13 (Patau syndrome) (Hunt and Hassold 2010). Studying the mechanisms of cohesion is therefore essential for understanding how the genome is passed properly from one cell generation to the next.

In addition, sister chromatid cohesion facilitates the repair of DNA double-strand breaks in cells that have replicated their DNA, where such breaks can be repaired by a homologous recombination mechanism that uses the undamaged sister chromatid as a template (for review, see Watrin et al. 2006). Furthermore, mutations in the proteins that are required for sister chromatid cohesion can cause defects in chromatin structure and gene regulation, and can in rare cases lead to congenital developmental disorders, called Cornelia de Lange syndrome, Roberts/SC Phocomelia syndrome, and Warsaw Breakage syndrome (for review, see Mannini et al. 2010).

**SISTER CHROMATID COHESION IS MEDIATED BY COHESIN**

Sister chromatid cohesion depends on a multi-subunit protein complex called cohesin (Fig. 1). Three of cohesin’s subunits—Smc1, Smc3, and Scc1/Rad21/Mcd1—form an unusual ring-shaped structure (Anderson et al. 2002; Haering et al. 2002). Smc1 and Smc3 are members of the “structural maintenance of chromosomes” (SMC) protein family. Members of this family are also found in other DNA-binding protein complexes, such as condensins, the Smc5/6 complex, and in bacterial SMC complexes. Like other SMC proteins, Smc1 and Smc3 are highly elongated molecules almost 50 nm in length that contain a central rod-shaped domain that is composed of anti-parallel coiled coils. Smc1 and Smc3 bind directly to each other at one of their ends via a “hinge” domain. At the other end of their coiled-coil domains, Smc1 and Smc3 form a nucleotide-binding domain (NBD). These NBDs form two ATP-binding sites, where one molecule of Mg^{2+}-ATP is simultaneously bound to Walker A and Walker B motifs in the NBD of one SMC subunit and to an ABC signature motif in the other SMC subunit (Haering et al. 2004). The NBDs of Smc1 and Smc3 are connected by the kleisin (“bridging”) subunit Scc1, with Scc1’s amino terminus being bound to Smc3 and its carboxyl-terminal Winged helix domain being bound to Smc1 (Haering et al. 2002). These interactions between Smc1, Smc3, and Scc1 lead to the formation of a tripartite ring with an inner diameter of 30–40 nm.

The integrity of this ring structure is essential for its ability to mediate sister chromatid cohesion (Gruber et al. 2003), and experimental observations using yeast mini-chromosomes indicate that cohesin mediates cohesion by topologically embracing the two sister chromatids (Ivanov and Nasmyth 2005; Haering et al. 2008). It has therefore been proposed that cohesin mediates cohesion by embracing both sister chromatids as a single ring (for review, see Nasmyth 2011). Other models propose that only a single DNA molecule can be embraced by a cohesin ring, and that two cohesin rings have to interact to connect two sister chromatids (Huang et al. 2005; Zhang et al. 2008b).

A fourth cohesin subunit, called Scc3 in yeast and stromal antigen (SA1/2) in higher eukaryotes, binds directly to Scc1 (Haering et al. 2002). Scc3 is also essential for sister chromatid cohesion (Michaelis et al. 1997; Nishiyama et al. 2010). In vertebrate somatic cells, cohesin...
complexes contain either one of two different SA/STAG proteins, called SA1 and SA2 (Losada et al. 2000; Sumara et al. 2000). In most organisms, Scc1 and Scc3/SA interact with several other proteins called Wapl/Rad61, Pds5, and Sororin (Table 1). These proteins are believed to determine how stably the cohesin ring interacts with DNA, and if these interactions are able to mediate cohesion (see below).

In many organisms, different kleisin subunits and isoforms of Smc1, Scc3/SA, and Pds5 exist, some of which are differentially expressed between somatic and meiotic cells. For example, in mammalian oocytes and spermatocytes, many cohesin complexes contain Smc1β instead of Smc1 (the latter is therefore also called Smc1α), and the kleisins Rec8 or Rad21L instead of Scc1. Table 1 provides an overview regarding the different isoforms of these and other cohesin subunits and their regulators that have been identified in different species.

LOADING OF COHESIN ONTO DNA DEPENDS ON ADHERIN, COHESIN’S ATPASE ACTIVITY, AND POSSIBLY OPENING OF THE COHESIN RING

Cohesin is loaded onto DNA before DNA replication, in vertebrates already at the end of mitosis in telophase (Fig. 2). This process depends on cohesin’s ATPase activity (Arunugam et al. 2003; Weitzer et al. 2003) and on a separate complex, called adherin (Furuya et al. 1998) or kollerin (Nasmyth 2011). Because cohesin is thought to embrace DNA as a ring, it has furthermore been proposed that cohesin rings have to be opened transiently to allow entry of the DNA into the ring (Gruber et al. 2003).

Adherin is a chromatin-bound protein complex composed of two subunits, a large protein (with a molecular mass of 316 kDa in humans), called Scc2 in budding yeast and NIPBL in mammals, and a smaller subunit, called Scc4 in yeast and MAU2 in mammals (Table 1). In Xenopus eggs, adherin is recruited to chromatin by binding to pre-replicative complexes (pre-RCs) (Gillespie and Hirano 2004; Takahashi et al. 2004), whereas in yeast, adherin is particularly enriched at centromeres but is also found on chromo-
Table 1. Subunits and regulators of cohesin in yeasts, fruit flies, and humans

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<thead>
<tr>
<th></th>
<th>Saccharomyces cerevisiae</th>
<th>Schizosaccharomyces pombe</th>
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<td>Hos1</td>
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*Expressed specifically in meiosis and largely/partially replace mitotic subunit.

b Also expressed in B cells.

c Paralog of CTCF specifically expressed in testis.
the two NBDs (Hopfner et al. 2000), and in the case of ABC transporters, this "engagement" step facilitates transport of substrate molecules through the transmembrane channel (for review, see Rees et al. 2009). ATP hydrolysis reverts these conformational changes by leading to disengagement of the NBDs. Similarly to how one cycle of ATP binding and hydrolysis leads to

Figure 2. The cohesin cycle in vertebrate cells. A schematic illustration showing cohesin loading, cohesion establishment, and cohesin removal from chromosomes in telophase/G1 phase, S phase, and mitosis, respectively. (Small circles) Cohesin complexes; (light gray circles) cohesin complexes that are dynamically bound to DNA; (dark gray circles) "cohesive" complexes that are stably bound to DNA (Gerlich et al. 2006).

Figure 3. Dynamic and stable binding modes of cohesin to DNA. A schematic drawing illustrating how adherin-mediated separation of the hinge domains of Smc1 and Smc3 might create an entry gate for DNA into the cohesin ring, whereas a transient Wapl-mediated dissociation of Scc1 from the SMC subunits may release cohesin from DNA by generating an exit gate (Nasmyth 2011; Chan et al. 2012). Wapl's ability to open this exit gate might be inhibited by sororin, which binds to cohesin in vertebrates following Smc3 acetylation and possibly other unidentified events during DNA replication (Nishiyama et al. 2010).
transient opening of the transmembrane channel in ABC transporters, it is therefore possible that ATP binding and hydrolysis would lead to transient opening of an entry gate for DNA in the cohesin ring.

But how could binding of ATP to the NBDs of Smc1 and Smc3 lead to separation of the hinge domains of Smc1 and Smc3, which are almost 50 nm away from the NBDs? Atomic force microscopy and crystallographic studies have revealed that ATP binding to the signature motif of Rad50 is not only accompanied by NBD engagement but also induces a dramatic conformational change that alters the angle at which the coiled-coil domain of Rad50 emanates from the NBD (Moreno-Herrero et al. 2005; Lammens and Hopfner 2010; Lim et al. 2011; Williams et al. 2011). This conformational change is mediated by helices in Rad50, called the signature-coupling helices, which move relative to each other and thereby function as a lever that exerts force on the coiled-coil domain. It is therefore conceivable that ATP binding and hydrolysis also induce changes in the coiled-coil domains of cohesin, and thereby could affect interactions between Smc1’s and Smc3’s dimerization domains over relatively long distances.

Further support for the hypothesis that cohesin loading onto DNA is controlled by ATP hydrolysis comes from the recent identification of a cohesin mutant in the Walker B motif of Smc1 and Smc3 that might be locked in a transition state of the loading reaction (Hu et al. 2011). This mutant binds to DNA in an Sccl-dependent manner, in particular at centromeres, but is unable to associate with DNA stably and to mediate cohesion. Based on analogy with ABC transporters, it has been speculated that this mutant might be locked in a state where the NBDs are engaged but the hinge domains are physically separated, which would explain why these mutants can interact with DNA only transiently.

COHESIN IS POSITIONED AT SPECIFIC SITES IN THE GENOME

Chromatin immunoprecipitation (ChIP) experiments performed in several different organisms have revealed that cohesin can predominantly be detected at specific sites in the genome, indicating that cohesin is not distributed randomly along the length of chromosomes (for review, see Peters et al. 2008). For example, in budding yeast and fission yeast, large amounts of cohesin are found at centromeres and pericentric regions, and smaller amounts are found in discrete cohesin attachment regions (CARs) along chromosome arms, which on average are 10 kb apart from each other. Because adherin is essential for the ability of cohesin to associate with DNA, it is generally assumed that cohesin loading occurs at adherin-binding sites. However, ChIP experiments in budding yeast and fission yeast indicate that most adherin complexes are bound to different regions of the genome from cohesin, with the exception of centromeres where both complexes are abundant (Lengronne et al. 2004; Schmidt et al. 2009; Hu et al. 2011; for a conflicting view, see Kogut et al. 2009). It has therefore been proposed that cohesin is first loaded onto DNA at adherin sites and that it is subsequently translocated from these sites to pericentric regions or CARs (Lengronne et al. 2004). Consistent with this notion, the Walker B mutant of cohesin, which is thought to be locked in a transition state of the loading reaction, accumulates at yeast centromeres but to a much lesser degree than wild-type cohesin at pericentric regions and CARs, as if cohesin were normally loaded predominantly at centromeres and subsequently moved from there to chromosomal arm regions (Hu et al. 2011).

How cohesin is moved from one site to the other remains unknown, but many CARs in budding and fission yeast are found in regions of convergent transcription, and it has been shown that the induction of regulatable genes leads to removal of cohesin from an intragenic position to the end of the transcription unit (Lengronne et al. 2004). This finding and biochemical experiments that suggest that cohesin rings are able to “slide off” short linearized yeast mini-chromosomes in vitro (Ivanov and Nasmyth 2005) have led to the proposal that cohesin is able to move laterally along DNA while encircling it. According to this idea, RNA polymerases or the native transcripts and RNPs produced by them would simply push cohesin out...
of actively transcribed regions. If this is true, the pattern of cohesin distribution that is seen along yeast chromosome arms could largely be an indirect consequence of gene activity, rather than a pattern that is specified because cohesin could only fulfill its function in particular chromosomal locations. An exception to this rule could be the centromere, where a high number of cohesin complexes are thought to resist the pulling forces that are generated by the mitotic spindle.

Because cohesin mediates sister chromatid cohesion from yeast to men, the principle mechanisms of how cohesin associates with DNA might have evolved early during the evolution of eukaryotes, or possibly even before, because DNA-binding SMC complexes related to cohesin exist in all kingdoms of life. Although this may well be the case for the initial loading process in which the cohesin rings entrap DNA, it is less clear if the subsequent positioning of cohesin in the genome occurs according to the same principles in different organisms. In contrast to the situation in yeasts, where cohesin and adherin can be found in different chromosomal locations, the two complexes largely colocalize in the Drosophila genome, where in many cases they occupy the entire length of actively transcribed genes (Misulovin et al. 2008). Yet a different situation is found in mammalian cells, where NIPBL is frequently located at the promoters of active genes, often together with mediator, a large transcriptional coactivator complex, and cohesin (Kagey et al. 2010). However, although some cohesin can be detected at the majority of NIPBL sites (Kagey et al. 2010), the opposite is not true. There are many more cohesin-binding sites detectable in ChIP experiments (estimates range up to 55,000 sites for the nonrepetitive part of mammalian genomes) (Schmidt et al. 2010) than NIPBL sites, implying that also in mammals, cohesin loading may occur at sites that are distinct from the sites where cohesin finally accumulates. But unlike in yeasts, there is presently no evidence that cohesin is moved to these positions by the transcriptional machinery.

Instead, there are numerous observations that indicate that the pattern of cohesin distribution in mammalian genomes is specified by DNA sequence elements that are recognized by CCCTC binding factor (CTCF), a protein that binds to DNA via one or several of its 11 zinc finger motifs. CTCF is found at the majority, but possibly not at all cohesin-binding sites (for review, see Merkenschlager 2010). Interestingly, partial depletion of CTCF or mutation of CTCF-binding sites reduces cohesin binding at these sites, without detectably reducing the total amount of cohesin on DNA (Parelho et al. 2008; Stedman et al. 2008; Wendt et al. 2008). The implication is that CTCF is dispensable for loading cohesin onto DNA but required for a subsequent positioning step. This situation is particularly evident at the H19-Igf2 locus. In this chromosomal location, the maternal and paternal alleles are differentially methylated, leading to different gene expression patterns on the two alleles within one and the same cell. This “imprinted” gene expression pattern depends on CTCF, which can bind to an imprinting control region (ICR) on the maternal allele but not on the paternal allele, where cytosine methylation prevents CTCF binding to the ICR. Interestingly, cohesin binds only to the ICR on the maternal allele, where CTCF is present, but not on the paternal allele, where CTCF is not bound, further supporting the hypothesis that CTCF specifies where cohesin accumulates once it has been loaded onto DNA (Rubio et al. 2008; Stedman et al. 2008; Wendt et al. 2008; Nativio et al. 2009).

Recent experiments have shown that the carboxy-terminal region of CTCF can directly bind to the SA2 subunit of mammalian cohesin (Xiao et al. 2011), suggesting that cohesin may be recruited to CTCF-binding sites through direct protein–protein interactions. But how cohesin moves from its presumed loading sites where NIPBL is present to CTCF sites remains unknown. Either cohesin can somehow be translocated along DNA, or loading would normally have to occur directly at CTCF sites. In the latter case, one would have to postulate that NIPBL is present, at least transiently, at many more CTCF sites than could be detected in ChIP experiments so far, or NIPBL might be able to promote cohesin loading in cis. In this case, an NIPBL site would have to be able to interact
with multiple CTCF sites, possibly in a dynamic fashion that depends on chromatin folding.

**RECRUITMENT OF COHESIN TO CTCF SITES MAY DETERMINE CHROMATIN STRUCTURE**

There is presently no evidence that cohesin has to accumulate at CTCF sites to be able to mediate sister chromatid cohesion, because no cohesion defects have been observed after depletion of CTCF, despite the fact that less cohesin is detected at CTCF sites under these conditions (Wendt et al. 2008). Furthermore, cohesin does not colocalize with CTCF on *Drosophila* chromosomes, and there is no evidence that a CTCF-related protein colocalizes with cohesin in the genome of budding yeast or fission yeast, further indicating that cohesin’s role in cohesion is independent of CTCF. However, cohesin depletion does cause defects in gene expression (for review, see Dorsett and Ström 2012), and in mammalian cells, some of these defects resemble the defects that are seen after depletion of CTCF. For example, cohesin depletion changes the levels of H19 and Igf2 transcripts in a similar way as depletion of CTCF, implying that both proteins are required for imprinted gene expression at the H19-Igf2 locus (Wendt et al. 2008; Nativio et al. 2009). Because CTCF is required for recruitment of cohesin to the ICR, whereas the opposite is not true, CTCF appears not to be sufficient for imprinted gene expression. Instead, the main function of CTCF in this process might be to recruit cohesin, and it could therefore be cohesin that affects transcription more directly than CTCF. If this were true, then cohesin positioning in mammalian cells might not be an indirect consequence of gene expression, as has been proposed to be the case in yeast, but may on the contrary control patterns of gene expression.

It is believed that cohesin controls gene expression by mediating long-range chromatin interactions, for example, by forming physical contacts between different CTCF sites that could lead to the formation of chromatin loops. This hypothesis is based on the notion that activity of gene expression depends on promoter–enhancer interactions, and that the proximity between promoters and enhancers might be controlled topologically through the formation of chromatin loops or more complex higher-order chromatin structures. For example, CTCF is thought to function as a transcriptional insulator on the maternal allele of the H19-Igf2 locus by inducing the formation of a chromatin loop, which topologically separates the H19 enhancer from the promoters of the Igf2 gene. Given that cohesin is able to mediate sister chromatid cohesion, that is, it can physically connect two DNA molecules in *trans*, it is conceivable that cohesin also tethers different sequence elements (CTCF sites) on the same chromosome in *cis*, leading to the formation of intrachromatid loops. Support for this hypothesis has come from “chromatin conformation capture” (3C) experiments (for review, see Seitan et al. 2012), but how generally important cohesin and CTCF are for regulating chromatin structure remains to be determined.

**SISTER CHROMATID COHESION IS ESTABLISHED DURING DNA REPLICATION AND COINCIDES WITH STABILIZATION OF COHESIN ON DNA**

Cohesin can bind to DNA both before and after DNA replication, but it can establish sister chromatid cohesion normally only during S phase (Uhlmann and Nasmyth 1998). This and other observations indicate that cohesion establishment is coupled to DNA replication. It is attractive to think that cohesion establishment must occur at or in the direct vicinity of replication forks, because such a scenario could explain how cohesion is only established between sister chromatids, and not between non-sister DNA molecules. Consistent with this possibility, numerous replication fork components are required for cohesion (Table 2), and Eco1/Ctfl, an acetyltransferase required for establishment of sister chromatid cohesion (Skibbens et al. 1999; Toth et al. 1999), is thought to travel along DNA with replication forks (Lengronne et al. 2004). Even though cohesion may normally only be established at replication forks, recent experiments in budding yeast have revealed that cohesion can also be established de novo after
DNA replication if DNA has been damaged by double-strand breaks (see below). In addition, in this case, cohesion establishment depends on Eco1. However, in the absence of Eco1 and either DNA replication or DNA damage, the association of cohesin with DNA is not sufficient to establish sister chromatid cohesion. The implication is that it is not the binding of cohesin to DNA per se but an Eco1-dependent process and possibly other DNA replication or DNA damage-dependent events that make cohesin “cohesive.” An interesting hint as to what may determine the “cohesiveness” of cohesin has come from fluorescence recovery after photobleaching (FRAP) experiments in mammalian cells. These experiments revealed that cohesin binds to DNA dynamically in G1 phase with a residence time of several minutes, whereas many cohesin complexes bind to DNA much more stably after DNA replication, with a residence time of at least many hours (Gerlich et al. 2006). Thus, the ability of cohesin to mediate cohesion coincides with stabilization of cohesin on DNA, implying that stable binding of cohesin to DNA may be an inherent property of cohesive cohesin, and that this property may be required to maintain cohesion from S phase until the subsequent metaphase. Cohesin’s ability to bind to DNA for long periods of time may be particularly important in mammalian oocytes. In these cells cohesion is established during pre-meiotic S phase, which occurs before birth, and then has to be maintained until completion of meiosis, which only occurs during ovulation many months or years later (Hunt and Hassold 2010).

**Table 2. DNA replication proteins required for sister chromatid cohesion**

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<th><em>S. pombe</em></th>
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<td>And1</td>
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Cohesion Acetylation is Essential for Sister Chromatid Cohesion

Eco1’s essential role in sister chromatid cohesion is the acetylation of cohesin’s Smc3 subunit on two lysine residues, K112 and K113 in budding yeast, and K105 and K106 in human Smc3.
Both residues are well conserved in many eukaryotes, as are orthologs of Eco1, of which there are two different ones in vertebrates, called Esco1 and Esco2 (Table 1) (Hou and Zou 2005). Eco1 orthologs are also required for sister chromatid cohesion in fission yeast (Tanaka et al. 2000a), Drosophila (Williams et al. 2003), and human cells (Hou and Zou 2005), and in the latter case, both enzymes contribute to Smc3 acetylation (Zhang et al. 2008a; Nishiyama et al. 2010). The acetylation of Smc3 can be reverted by specific enzymes. In human cells, the major enzyme responsible for Smc3 deacetylation is HDAC8 (Dear dorff et al. 2012). In budding yeast, the deacetylase Hos1 is required for this process (Beckouet et al. 2010; Borges et al. 2010; Xiong et al. 2010). Hos1 deletion does not cause cohesion defects; however, it becomes crucial for cohesion either when Eco1 is inactivated (Beckouet et al. 2010) or when de novo Smc3 synthesis is suppressed (Borges et al. 2010). These observations raise the interesting possibility that cohesin may have to be acetylated during the process of cohesion establishment, whereas it may be impossible to convert previously acetylated cohesin into a cohesive state.

Although the importance of Smc3 acetylation for sister chromatid cohesion is well established, it is less clear how this modification enables cohesin to become cohesive. The acetylated lysine residues in Smc3 are predicted to be located close to the ATP-binding pocket in the NBD domain. Interestingly, mutations that are thought to mimic the acetylated state of Smc3 abolish a dominant-negative effect that Walker B mutations in Smc3 have on wild-type cohesin (Heidinger-Pauli et al. 2010). Because Smc3 mutations that prevent ATP binding also suppress this dominant effect, it has been proposed that Smc3 acetylation regulates the ATP-binding and hydrolysis cycle of cohesin, either by reducing ATP binding to cohesin or by preventing ATP-dependent interactions between cohesin and its loading machinery. Furthermore, cohesin is normally only acetylated when bound to chromatin. Together with the genetic interactions between acetyl-mimicking and Walker B mutations, this observation raises the possibility that Smc3 acetylation on chromatin prevents rebinding of ATP to cohesin that has entrapped DNA. According to this hypothesis, Smc3 acetylation could stabilize cohesin on DNA by blocking cycles of ATP binding and hydrolysis.

An alternative explanation for how acetylation may contribute to cohesion establishment has come from the observation that the essential function of Eco1 becomes dispensable in fission yeast and budding yeast if the cohesin-associated proteins Wapl and Pds5 are inactivated, or if specific residues in the cohesin core subunits Smc3 and Scc3 are mutated (Tanaka et al. 2001; Ben-Shahar et al. 2008; Unal et al. 2008; Zhang et al. 2008a; Rowland et al. 2009; Sutani et al. 2009). Because Wapl promotes the release of cohesin from DNA in vertebrates (see below), it has been proposed that Smc3 acetylation by Eco1 might stabilize cohesin on DNA by inactivating Wapl (Ben-Shahar et al. 2008; Unal et al. 2008). Alternatively, it has been argued that Wapl, Pds5, Smc3, and Scc3 are required for an activity that prevents cohesion establishment, and that Smc3 acetylation does not prevent the Wapl-mediated release of cohesin from DNA but instead overcomes this “anti-establishment” activity (Rowland et al. 2009). In vertebrates, the existing evidence supports the idea that Smc3 acetylation mediates cohesion by preventing dissociation of cohesin from DNA because in these species, cohesin acetylation leads to the recruitment of sororin, a protein that prevents release of cohesin from DNA (see below).

**SORORIN STABILIZES COHESIN–DNA INTERACTIONS AND MEDIATES SISTER CHROMATID COHESION BY ANTAGONIZING WAPL**

Sororin is a protein that has only been identified in metazoans so far, where it is essential for sister chromatid cohesion (Rankin et al. 2005) and for stabilizing cohesin on chromatin during DNA replication (Schmitz et al. 2007). Sororin only binds to cohesin once Smc3 has been acetylated. However, this modification is not...
sufficient for sororin recruitment because DNA replication is also necessary for this event (Lafont et al. 2010; Nishiyama et al. 2010). Remarkably, sororin’s essential function in sister chromatid cohesion becomes dispensable when Wapl is depleted from cells or *Xenopus* egg extracts, indicating that sororin might mediate cohesion by inhibiting Wapl (Nishiyama et al. 2010). This hypothesis could explain how sororin stabilizes cohesin on DNA (Schmitz et al. 2007), because the release of cohesin from DNA depends on Wapl (see below).

In vitro, sororin can displace Wapl from its binding partner Pds5, but there is no evidence that sororin can fully dissociate Wapl from cohesin (Nishiyama et al. 2010). It has therefore been proposed that sororin changes the topology of how Wapl and Pds5 interact with cohesin, and that these conformational changes inhibit Wapl’s ability to release cohesin from DNA. As a result, cohesin may become stably “locked” on DNA and may thus be able to mediate sister chromatid cohesion for prolonged periods of time (Fig. 3).

It remains unknown if cohesion depends on sororin in all species because related proteins have been identified in some but not all eukaryotes. Sororin exists in vertebrates and *Drosophila*, where it is called Dalmatian, and related proteins containing a conserved carboxy-terminal “sororin domain” have been identified in many other metazoans, including cephalochordates, echinoderms, cnidaria, placozoa, and in plants, but not yet in yeasts and worms (Nishiyama et al. 2010). It therefore remains to be understood if Smc3 acetylation regulates cohesin also in yeast and worms by recruiting a sororin-related protein or by a different mechanism.

**DNA DOUBLE-STRAND BREAKS ALSO INDUCE COHESION ESTABLISHMENT**

Although cohesion is normally established during DNA replication, DNA double-strand breaks also trigger de novo cohesion establishment even in postreplicative budding yeast cells in an Eco1-dependent manner (Strom et al. 2007; Unal et al. 2007). In this case, it is thought that acetylation of Scc1 by Eco1, but not of Smc3, is required for cohesion establishment (Heidinger-Pauli et al. 2009). This cohesion can be established independently of DNA replication and is induced by double-strand breaks not only on the damaged but also on undamaged chromosomes that are present in the same cells. These observations imply that a diffusible cohesion establishment factor, which is inactivated after DNA replication, is somehow reactivated in response to DNA damage. This factor is presumably Eco1 itself because Eco1 is normally degraded after S phase, but stabilized following DNA damage, and because overexpression of Eco1 is sufficient to induce cohesion establishment after DNA replication (Lyons and Morgan 2011).

**COHESIN IS REMOVED FROM MITOTIC CHROMOSOMES IN TWO STEPS**

Because cohesin physically connects sister chromatids, sister chromatids can only be separated from each other and transported toward opposite spindle poles in anaphase if cohesin is removed from chromosomes and cohesion is dissolved. This can be achieved by either of two mechanisms. In vertebrates, and presumably in most other metazoans, most cohesin complexes are removed from chromosome arms already in prophase and prometaphase, whereas cohesin at centromeres is protected from this “prophase pathway” and maintains cohesion until anaphase onset (Wäizenegger et al. 2000). It is this differential removal of cohesin from chromosome arms and centromeres, respectively, that causes the characteristic “X-shape” (or “V-shape” in species where centromeres are close to one of the telomeres) of chromosomes in prometaphase and metaphase (Gimenez-Abian et al. 2004).

**WAPL RELEASES COHESIN FROM DNA**

The dissociation of cohesin from chromosomes in prophase and prometaphase depends on Wapl (Gandhi et al. 2006; Kueng et al. 2006) and on phosphorylation of cohesin’s SA2 subunit (Hauf et al. 2005) by Polo-like kinase.
(Plk1) and possibly other mitotic kinases, such as cyclin-dependent kinase 1 (Cdk1) and Aurora B (Losada et al. 2002; Sumara et al. 2002; Hegemann et al. 2011). This pathway of removing cohesin from mitotic chromosomes might use the same mechanism by which cohesin is released from DNA in interphase because deletion of Wapl also increases the residence time and the steady-state levels of cohesin on DNA in interphase (Kueng et al. 2006). In mitosis, this release mechanism might become activated by mitotic kinases, either by phosphorylation of SA2, which is located next to Wapl in the cohesin complex (Gandhi et al. 2006; Kueng et al. 2006), or by phosphorylation of Wapl and Pds5, which are also phosphorylated in mitosis in a Plk1-dependent manner (Hegemann et al. 2011). Because sororin antagonizes Wapl, sororin also has to be inactivated in mitosis to enable Wapl to release cohesin from DNA. In addition, sororin inactivation is mediated by phosphorylation (Nishiyama et al. 2010), which has been reported to depend on Cdk1 (Dreier et al. 2011) or Plk1 (Zhang et al. 2011). At the end of mitosis, sororin is further degraded following ubiquitylation by the ubiquitin ligase APC/C<sup>Cdh1</sup>, possibly to prevent precocious formation of cohesin complexes that become too stably bound to DNA (Rankin et al. 2005). Activation of the prophase pathway of cohesin dissociation may therefore depend on mitotic phosphorylation of several proteins: sororin, SA2 (and presumably also SA1), and possibly Wapl and Pds5. In addition, adherin is also removed from chromosomes in prophase (Gillespie and Hirano 2004; Watrin et al. 2006), implying that cohesin cannot be loaded onto DNA any longer in early mitosis. Together, these events may shift the equilibrium between soluble and DNA-bound cohesin complexes so that during prophase and prometaphase most cohesin is removed from chromosome arms.

Although the majority of cohesin complexes are removed from mitotic chromosomes in vertebrate cells by Wapl, some cohesin complexes can only be removed by a different mechanism that depends on the protease separase, which cleaves the Scc1 subunit and thereby opens the cohesin ring (see below). Unlike separase, Wapl releases cohesin from DNA presumably by enabling opening of the cohesin ring without cleaving one of its subunits (Waizenegger et al. 2000; Kueng et al. 2006; Peters et al. 2008). It has been proposed that the Wapl-mediated opening of the cohesin ring occurs via an “exit gate” for DNA that is different from the proposed “entry gate” (Fig. 3) (Nasmyth 2011; Chan et al. 2012). Whereas the latter is thought to be created by separation of the hinge domains of Smc1 and Smc3 (see above), the former has been proposed to require dissociation of Scc1 from the SMC subunits (Nasmyth 2011). There is some precedence that such a conformational change could occur from structural studies of MukB, an SMC-related protein required for chromosome segregation in bacteria (Woo et al. 2009). MukB forms a homodimer whose NBDs interact with two carboxy-terminal Winged helix domains of the MukF protein, similarly to how Scc1’s carboxy-terminal Winged helix binds to Smc1 (Haering et al. 2002). Interestingly, ATP-mediated engagement of the MukB’s NBDs results in detachment of one of MukF’s Winged helix domains from MukB (Woo et al. 2009), implying that conformational changes could also lead to transient dissociation of Scc1 from cohesin’s SMC subunits.

POSSIBLE COEVOLUTION OF COHESIN ACETYLATION AND WAPL-MEDIATED COHESIN RELEASE FROM DNA

In budding yeast, it has been reported that a Plk1-dependent mechanism removes cohesin from meiotic chromosomes (Yu and Koshland 2005), but it is unknown if this pathway depends on Wapl, and in mitotic yeast cells, most if not all cohesin is cleaved by separase (Uhlmann et al. 1999). It is therefore unclear if there is a prophase pathway of cohesin dissociation in budding yeast that is related to the one in vertebrates. However, because Wapl antagonizes the function of Eco1 (see above), Wapl may also be required for releasing cohesin from DNA in budding yeast. Direct support for this notion has come from studies in fission yeast, where, as in vertebrates, Wapl is required for dissociation of cohesin from DNA (Feytout et al. 2011).
Nasmyth and Schleiffer (2004) pointed out that the genome of the simple eukaryote *Encephalitozoon cuniculi* lacks a gene that encodes an Eco1-related enzyme. We found 19 other species in whose genome we could not identify an Eco1-related gene (Table 3). Interestingly, in seven of these 20 species without a detectable Eco1 gene, at least one of the two lysine residues that are acetylated by Eco1 in yeast is replaced by another amino acid residue or missing. In contrast, we found only a single species (*Trypanosoma brucei*) with an Eco1 gene in which the KK motif in Smc3 was not conserved. More interestingly, of the seven species in whose genomes we could not identify an Eco1 gene but could find a mutated or missing KK motif in Smc3, six species have genes encoding either Pds5 or Wapl orthologs. The presence of Pds5 and Wapl therefore coincides in many cases either with the presence of Eco1, or with mutations in the lysine residues that are acetylated by Eco1 in other species. These observations raise the interesting possibility that the ability of Pds5 and/or Wapl to release cohesin from DNA may have coevolved with Smc3 acetylation by Eco1.

### Table 3. Coevolution of Smc3 acetylation by Eco1 and cohesin release by Wapl

<table>
<thead>
<tr>
<th>Organism</th>
<th>Smc3</th>
<th>Pds5</th>
<th>Wapl</th>
<th>Eco1</th>
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<tr>
<td><strong>Species with an Eco1 ortholog</strong></td>
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<tr>
<td><em>Homo sapiens</em></td>
<td>KK</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td><em>Ciona intestinalis</em></td>
<td>KK</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><em>Xenopus laevis</em></td>
<td>KK</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><em>Arabidopsis thaliana</em></td>
<td>KK</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><em>Saccharomyces cerevisiae</em></td>
<td>KK</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><em>Schizosaccharomyces pombe</em></td>
<td>KK</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td><em>Drosophila melanogaster</em></td>
<td>KK</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td><em>Trypanosoma brucei</em> TREU927</td>
<td>KQ</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
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<td>KQ</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td><em>Dictyostelium fasciculatum</em></td>
<td>NK</td>
<td>+</td>
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<td>KK</td>
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<td>HK</td>
<td>+</td>
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<tr>
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<td>KK</td>
<td>+</td>
<td>+</td>
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<td>KK</td>
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<tr>
<td><em>Volvoc carteri f. nagariensis</em></td>
<td>—</td>
<td>+</td>
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<tr>
<td><em>Cryptosporidium parvum</em></td>
<td>QK</td>
<td></td>
<td></td>
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<tr>
<td><em>Aureococcus anophagefferens</em></td>
<td>KK</td>
<td></td>
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<tr>
<td><em>Encephalitozoon cuniculi</em> GB-M1</td>
<td>KK</td>
<td></td>
<td></td>
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<tr>
<td><em>Schistosoma japonicum</em></td>
<td>KK</td>
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(KK) Lysine residues that correspond to the ones that have been shown to be acetylated by Eco1 in *S. cerevisiae* Smc3. In the sequence of *Volvoc carteri f. nagariensis* Smc3, the region containing these lysine residues is missing. Gene or protein sequences are available upon request.

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chromosomes arms but leaves cohesin behind at centromeres, where cohesion is maintained until the onset of anaphase (Waizenegger et al. 2000). These centromeric cohesin complexes are protected from the prophase pathway by a protein called shugoshin, or Sgo1, which recruits the protein phosphatase 2A complex (PP2A) to centromeres (for review, see Gutiérrez-Caballero et al. 2012). PP2A has been proposed to protect cohesin from the prophase pathway by reverting the phosphorylation of SA2 at centromeres (Kitajima et al. 2006), but it is conceivable that PP2A also has other critical substrates whose dephosphorylation is required to maintain centromeric cohesion, such as sororin, Wapl, or Pds5. A second protein that has been shown to protect cohesin at centromeres is a protein kinase called Haspin. Haspin’s only known substrate is the core histone H3, which can be phosphorylated on threonine 5, but how Haspin contributes to protection of cohesin at centromeres remains to be understood (for review, see Higgins 2010).

Those cohesin complexes that are protected from the prophase pathway by shugoshin-PP2A can only be removed from chromosomes by separase, a protease that cleaves the Scc1 subunit of cohesin at two distinct sites and thereby opens the cohesin ring (Uhlmann et al. 1999, 2000; Waizenegger et al. 2000). This process is essential for sister chromatid separation because either expression of non-cleavable Scc1 or inactivation of separase prevents proper sister chromatid separation (for review, see Peters et al. 2008).

In budding yeast, Scc1 cleavage by separase is also a prerequisite for Smc3 deacetylation by Hos1 (Beckouet et al. 2010; Borges et al. 2010). If cohesin contains non-cleavable Scc1, Smc3 is not deacetylated at the time of anaphase onset even though cyclin B is degraded. Vice versa, Scc1 cleavage triggers Smc3 deacetylation in metaphase arrested cells in the absence of cyclin B degradation. Although Scc1 cleavage is a prerequisite for cohesin dissociation from chromatin, cleavage per se is not necessary for deacetylation in vitro because release of cohesin from DNA by nuclease digestion is sufficient for Smc3 deacetylation even if Scc1 remains intact (Borges et al. 2010). Separase may therefore enable cohesin deacetylation by simply releasing cohesin from chromatin, with which Hos1 is not associated (Borges et al. 2010).

**CONCLUDING REMARKS**

It is one of the most fundamental aspects of life that new cells are generated by division of older cells and that the newly formed daughter cells share most if not all properties of their mothers because the daughters inherit an identical copy of the mother cell’s genome. The two key processes that constitute this chromosome cycle are the precise duplication of the genome by DNA replication and the subsequent segregation of the two copies of the genome by chromosome segregation. Although the basic logic of this cycle was understood already during the early days of cell biology in the 19th century, it has only become obvious in the more recent past that understanding the mechanisms of sister chromatid cohesion will also be essential for understanding the eukaryotic cell division cycle. It is now well established that sister chromatid cohesion is mediated by cohesin, but exactly how cohesin performs this and possibly other functions and how these are controlled during the cell cycle and during cell differentiation remain incompletely understood. Numerous important questions will have to be answered to build a comprehensive molecular model of how cohesin mediates these functions: How and where does adherin load cohesin onto DNA? Is cohesin translocated from loading sites to its final binding sites? If so, how, and by which mechanism does CTCF recruit cohesin to these sites in vertebrates? How does Wapl release cohesin from DNA, and how is this process prevented by sororin or possibly by other mechanisms that depend on cohesin acetylation? Is cohesin converted into a cohesive form at replication forks? If so, how are cohesin acetyltransferases and sororin recruited to these forks, and exactly how do the two sister DNA strands become connected by cohesin at or in the vicinity of replication forks? What happens to cohesin when DNA or RNA polymerases or other enzymes move along DNA? Can these enzymes move through...
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cohesin rings, or bypass them, or push them along DNA, or do they have to evict cohesin transiently from DNA? Is cohesin really able to mediate cohesion for months or years, as has been speculated to be the case in mammalian oocytes? How does cohesin's ability to mediate cohesion relate to its role in gene regulation? Does cohesin mediate the latter function through the formation of intrachromatid loops? If yes, how is the ability of cohesin to connect DNA strands targeted to sister chromatids when cohesion is established, but to different DNA sequences in cis during the process of gene regulation? Answering these and further questions will be an exciting task for the future, and an essential prerequisite for understanding how chromosomes are organized and passed from one cell generation to the next.

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# Sister Chromatid Cohesion

Jan-Michael Peters and Tomoko Nishiyama

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