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The cohesin ring concatenates sister DNAs

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Summary

Sister chromatid cohesion, which is essential for mitosis, is mediated by a multi-subunit protein complex called cohesin whose Scc1, Smc1, and Smc3 subunits form a tripartite ring structure. It has been proposed that cohesin holds sister DNAs together by trapping them inside its ring. To test this, we used site-specific cross-linking to create chemical connections at the three interfaces between the ring's three constituent polypeptides, thereby creating covalently closed cohesin rings. As predicted by the ring entrapment model, this procedure produces dimeric DNA/cohesin structures that are resistant to protein denaturation. We conclude that cohesin rings concatenate individual sister minichromosome DNAs.

Introduction

Sister chromatid cohesion is mediated by a multi-subunit complex called cohesin that contains four core subunits: Smc1 and Smc3, members of the SMC (Structural Maintenance of Chromosomes) protein family, and two non-SMC subunits, Scc1/Mcd1, a member of the kleisin family, and Scc3/SA ^{1,2}. Sister chromatid disjunction occurs when all chromosomes have been bi-oriented and is triggered by site-specific cleavage of cohesin's Scc1 subunit by separase ³. Cohesin's Smc1 and Smc3 both form rod shaped molecules that heterodimerize via "hinge" domains situated at the ends of 30 nm long intra-molecular antiparallel coiled coils ^{4,5}. ATPase "heads" at the other ends are connected by cohesin's Scc1 kleisin subunit, thereby forming a tripartite ring with a 35 nm diameter ^{4,6}. It has been proposed that cohesin holds sister chromatids together by trapping sister DNAs inside its ring. By severing Scc1, separase is envisioned to open the ring and thereby release sister DNAs from their topological embrace.

To investigate the physical nature of sister chromatid cohesion, we have recently used sucrose gradient sedimentation and gel electrophoresis to purify from yeast cohesed sister chromatids of small circular minichromosomes ⁷. The minichromosome dimers are composed of individual DNAs packaged into nucleosomes that are converted to monomers by cleaving Scc1 or by linearizing their DNA. Importantly, their formation depends on centromeres as well as cohesin (see Supplementary Information and Fig. S1). These data are consistent with (but do not prove) the notion that cohesin attaches to chromatin using a topological mechanism. However, they do not exclude the possibility that cohesion requires the non-topological association of cohesin rings bound to different chromatin fibres ⁸. If cohesin holds dimeric minichromosomes together by trapping them inside its ring, then introducing covalent connections between the Smc1/Smc3 hinge, Smc1 head/Scc1-C and Smc3 head/Scc1-N interfaces should create a chemically circularized cohesin ring within which sister DNAs would be trapped even after protein denaturation (Fig. 1a). Here we describe experiments that test this crucial prediction.

Covalent connection of the three cohesin ring subunits

To connect cohesin subunit interfaces covalently, we used the homobifunctional thiol-reactive chemicals dibromobimane (bBBr) and bis-maleimidoethane (BMOE) that bridge thiol groups up to 5 Å or 8 Å respectively ^{9,10} (Supplementary Fig. S2). We created a homology model ¹¹ of the yeast Smc1/Smc3 hinge heterodimer based on the homodimeric *Thermotoga maritima* crystal structure ⁴ and identified two juxtaposed side-chains that we mutated to cysteines. Incubation of the engineered Smc1/Smc3 hinge dimer with either bBBr or BMOE caused efficient cross-linking within a few minutes (Fig. 1b). Crucially, cross-linking was dependent on both cysteine substitutions. We used the same approach to connect the loop between the two β-strands of Scc1's winged-helix to a β-strand in Smc1's

ATPase head ¹² (Fig. 1c). Because no structural information is available for the interface between Smc3's ATPase and Scc1's N-terminal domain, we expressed Smc3 and Scc1 as a fusion protein, using a long flexible linker containing triple target sequences for the TEV protease to connect Smc3's C-terminus with Scc1's N-terminus (Smc3-TEV-Scc1) ¹³. To create cohesin rings that could be chemically circularized by BMOE or bBBr, the cysteine substitutions were introduced into Smc3-TEV-Scc1 and Smc1.

Cohesin circularization produces SDS resistant minichromosome dimers

A 2.3 kbp circular minichromosome ⁷ was introduced into yeast strains whose Smc1 and Smc3-TEV-Scc1 polypeptides contained either all four cysteine substitutions or only a subset of these. After nocodazole arrest and cell lysis, extracts were centrifuged through sucrose gradients and fractions containing monomeric and dimeric minichromosomes detected by native agarose gel electrophoresis and Southern blotting. Importantly, dimeric minichromosomes could still be isolated from yeast cells whose cohesin ring subunits had been engineered to permit cohesin circularization. The cysteine substitutions had little adverse effect, but the fusion of Scc1 to Smc3 roughly halved the fraction of dimeric minichromosomes (Fig. 2a). This was not surprising as the fusion causes partial cohesion defects in vivo 13. DTT, sucrose and other low molecular weight contaminants were removed from the gradient fractions by dialysis and cohesin subunits treated with bBBr, BMOE, or merely DMSO solvent. After quenching the reaction by re-addition of DTT, SDS was added to a final concentration of 1% and the samples were heated to 65°C for four minutes. The denatured samples were finally electrophoresed in agarose gels containing ethidium bromide and minichromosome DNA was detected by Southern blotting.

Dimer fractions from control cells expressing unmodified cohesin contained four species of DNA (Fig. 2b, panel F). The fastest migrating and most predominant are supercoiled monomers (①). Due to SDS in the loading buffer, the next two species (② and ③) were poorly resolved from each other. These DNAs co-migrated with monomeric nicked circles produced by nicking enzyme after removal of nucleosomes with 2 M KCl (Fig. 2d) and with (infrequently) intertwined (i.e. concatenated) supercoiled DNAs isolated from a topoisomerase II mutant (Fig. 2e) and therefore include both of these species of DNA. The nicking enzyme treatment revealed that about 10% of DNAs from dimer (but not monomer) fractions are DNA-DNA concatemers (Fig. 2d and data not shown). The least abundant species (④) migrated more slowly than two intertwined supercoiled circles (③) but more rapidly than two intertwined nicked circles generated by treatment with nicking enzyme (⑤). We conclude that these DNAs correspond to one supercoiled circle intertwined with one nicked circle.

Treatment of dimer fractions with bBBr or BMOE had no effect on the electrophoresis profile of minichromosome DNAs (Fig. 2b, F). Moreover, the very same pattern was observed when dimeric minichromosomes were isolated from strains expressing the Smc3-TEV-Scc1 fusion (Fig. 2b, E) and cross-linkable cysteine pairs at either the Smc1/Scc1 or the Smc1/Smc3 interface (Fig. 2b, C and D). In contrast, bBBr and BMOE but not DMSO alone caused the appearance of two additional species of DNA when dimers were isolated from a strain containing the Smc3-TEV-Scc1 fusion and cysteine pairs at both interfaces (Fig. 2b, B). The more abundant (⑦) migrated slightly more slowly than intertwined supercoiled circles whereas the less abundant (⑧) migrated slightly more slowly than supercoiled circles intertwined with nicked circles. Their electrophoretic mobilities and the fact that neither was detected when identical cross-linking reactions were conducted with monomer fractions (Fig. 2b, A) suggest that they represent novel dimeric forms. Their formation occurs at the expense of supercoiled and nicked monomeric circles.

Our data suggest that the faster form $(\bar{\mathcal{D}})$ is a dimer of supercoiled monomeric circles associated with cohesin while the slower form (®) is a dimer between supercoiled and nicked circles associated with cohesin. Consistent with this, both were converted to a form (9) that co-migrates with intertwined nicked circles (5) when treated with nicking enzyme (Fig. 2d). The extra mass of cohesin probably has little effect on the electrophoretic mobility of this slow running species. Importantly, neither novel dimer was produced when just one of the four cysteine substitutions was lacking (Supplementary Fig. S3a), implying that they arise due to the simultaneous cross-linking of both cysteine pairs at the Smc1/Smc3 hinge and Smc1/Scc1 interfaces. Cross-linked dimers were also produced when minichromosomes were isolated from cycling cultures (Supplementary Fig. S3a), suggesting that their formation is not an artefact caused by arresting cells with nocodazole. Finally, no slower migrating species could be observed when DNA was linearized with a restriction enzyme after cross-linking (Supplementary Fig. S3b). In conclusion, covalent closure of the cohesin ring converts dimeric but not monomeric minichromosomes to a dimeric form that is resistant both to SDS and to 2 M KCl (native dimers are converted to monomers at 0.5-1 M KCl; Supplementary Fig. S4).

Circularized cohesin holds individual DNAs together

To test whether the SDS-resistant dimers produced by cohesin circularization are indeed monomeric DNAs held together by cohesin, we employed two-dimensional (2D) gel electrophoresis. Denatured cross-linked samples were resolved on an agarose gel as before (the first dimension) and then electrophoresed perpendicularly through a thin zone of agarose or agarose containing proteinase K into a second agarose gel (the second dimension). Proteinase K should digest any proteins before DNAs enter the second gel and DNAs that ran as dimers in the first dimension should run as monomers in the second

dimension if they were initially held together by a proteinaceous (i.e. cohesin) connection. In the absence of proteinase K, all DNA species migrate identically in first and second dimensions and therefore lie on a diagonal line (Fig. 3a). Several species also ran on the diagonal in the presence of proteinase K, namely monomeric supercoils (①), monomeric nicked circles (②), intertwined supercoils (③), and nicked circles intertwined with supercoils (④) (Fig. 3a). In contrast, DNAs of presumptive dimers of two supercoiled minichromosomes held together by cohesin (⑦) migrated as monomeric supercoils in the second dimension (⑦→①), while presumptive supercoiled-nicked circle dimers held together by cohesin (⑧) split into monomeric supercoils (⑧→①) and nicked circles (⑥→②). We conclude that chemical circularization of cohesin associated with native dimeric minichromosomes is accompanied by the cross-linking of monomeric DNAs to create SDS-resistant but protease sensitive dimers.

The protease containing 2D gel revealed two new types of low abundance DNAs. The first (⑥) migrated considerably slower than monomeric supercoils in the first dimension but ran as monomeric supercoils in the second dimension (⑥→①). These DNAs were only detected in monomeric or dimeric minichromosome preparations in which cohesin rings had been covalently closed (data not shown). They presumably correspond to rare supercoiled monomers whose migration is retarded by their association with (entrapment by) a chemically circularized cohesin ring. The second species (⑩) co-migrated with cohesin-mediated supercoiled dimers in the first dimension but with intertwined supercoils (and nicked circles) in the second dimension. These DNAs could correspond either to monomeric nicked circles associated with cohesin or, more likely, to intertwined supercoils that are also associated with cohesin.

If cohesin circularization by bBBr and BMOE cross-linking per se is responsible for the formation of SDS-resistant minichromosome dimers, then cleavage of the cohesin ring should be sufficient to release the monomeric DNAs. To test this, we incubated cross-linked dimeric minichromosome preparations with or without TEV protease to cleave the linker connecting Smc3 and Scc1. The presence of TEV greatly reduced both types of DNA dimers induced by cohesin's circularization (⑦ and ⑧), which was accompanied by a corresponding increase in monomeric DNAs (① and ②) (Fig. 3b). This effect was clearly caused by cleavage of the TEV sites in the Smc3-Scc1 linker because DNA dimers produced by circularization of cohesin with a TEV-resistant Smc3-Scc1 linker were unaffected by TEV protease (Fig. 3b). We conclude that the SDS-resistant association of sister DNAs induced by cross-linking cohesin's three subunits does not merely accompany the circularization of cohesin but actually depends on it.

Are minichromosomes held together by single or double cohesin rings?

The simplest explanation for the cross-linking results is that sister DNAs are topologically trapped within single (monomeric) cohesin rings (Supplementary Fig. S5a). An alternative albeit more complicated possibility envisions entrapment of sister DNAs by rings that are themselves topologically intertwined (Supplementary Fig. S5b) or by dimeric cohesin rings. Only two cysteine cross-links are needed for entrapment by single rings, while four are required by double ring models. If we knew the efficiency with which bBBr and BMOE cross-link the Smc1/Smc3 and Smc1/Scc1 interfaces and the number of cohesin bridges, then we could calculate the fraction of DNAs that should be trapped as dimers according to the two models and compare these to what is actually observed.

To estimate the protein cross-linking efficiency, we spiked cross-linkable minichromosome dimer preparations with purified Smc1/Smc3 hinge or Smc1 head/Scc1-C before crosslinking with bBBr or BMOE and denaturation. One half of the reactions was run on SDS-PAGE and the fraction of proteins cross-linked measured after SYPRO ruby staining (Fig. 4a). The other half was run on an agarose gel and the fraction of DNAs dimerized measured by Southern blotting (Fig. 4b). The fraction of rings expected to be cross-linked at both interfaces, which is given by multiplying individual cross-linking efficiencies, was 30% for both bBBr and BMOE. We would therefore expect 30% of DNAs to be dimerized if held together by a single cohesin ring but only 9% by a double ring. Estimating the actual number of bridges is harder because the gradient fractions contain much cohesin not associated with minichromosomes (data not shown). However, if we assume that a single bridge is sufficient to hold sister DNAs together and that cross-bridges form in vivo and survive fractionation in vitro with a defined probability (λ), then the fraction of chromosomes f(x) with 0, 1, 2, ... n bridges should fit a Poisson distribution. f(0) can be measured directly, namely by measuring the fraction of monomeric minichromosomes (Supplementary Fig. S6a) and DNA-DNA concatemers (Fig. 2d), which permits calculation of λ (see Supplementary Information). Because f(0) is large, most native dimeric minichromosomes are predicted to have a single bridge. Taking this into account, the single and double ring models predict 32% and 10% dimerization, respectively. The observed value with both reagents was 30% (Fig. 4b), which is inconsistent with the double ring model and close to that predicted by the single ring model.

Single and double ring models also make different predictions for heterozygous diploids that express equal amounts of TEV-cleavable and TEV-resistant Smc3-Scc1 fusion proteins (Supplementary Fig. S6b). In the case of one cross-bridge, the single ring model predicts that 50% of cross-linked dimers should survive cleavage of half the cohesin rings. In contrast the double ring model predicts only 25% because cleavage of just one ring is sufficient to destroy dimers held together by intertwined rings. We isolated

minichromosome dimers from cleavable and non-cleavable haploids, a 1:1 mixture of the two, and from heterozygous diploids. These were cross-linked, treated either with TEV protease or a non-catalytic TEV mutant, denatured with SDS and run on an agarose gel. The fraction of DNAs dimerized by cross-linking was measured by scanning Southern blots. This revealed that about 50% of cohesed minichromosomes survived TEV treatment when isolated from heterozygous diploids as well as the 1:1 mixture of haploids (Fig. 4c). These data fit the single but not the double ring model. We note that the latter also predicts that a sizeable fraction of cross-linked dimers (⑦ and ⑧) from heterozygous diploids should be converted by TEV cleavage to supercoiled monomers associated with cohesin (⑥), which is not observed (Fig. 4c).

Discussion

Our cross-linking experiments suggest that sister minichromosome DNAs are entrapped by a single monomeric ring. Importantly, they exclude the possibility that the connection between sister DNAs is mediated by non-topological interactions between cohesin complexes associated with each sister ⁸. Given the specificity of the cross-linking by bBBr and BMOE, there is no reason to suppose that putative interactions between cohesin rings will have been cross-linked in our experiments. Double ring models envisioning topological cohesin-cohesin interactions or a gigantic ring formed by two cohesin complexes are difficult to reconcile with the findings that the fraction of DNAs dimerized is almost identical to the fraction of cohesin rings circularized and not to the square of this fraction and that cleavage of half the cohesin rings reduces dimers to 50% and not 25%.

Our conclusion that sister chromatin fibres of dimeric minichromosomes are threaded through cohesin rings provides a simple and potentially adequate mechanism to explain cohesin's ability to hold sister chromatids together. Cohesin could therefore be considered a "concatenase". It will be important to address whether it uses the same mechanism at loci farther away from core centromeres, whether it sometimes traps individual chromatin fibres and if so whether it is capable of forming chromatin loops. We detected rare instances where the individual DNA trapping occurred on our minichromosomes, namely monomeric DNAs that upon cohesin circularization are retarded in their electrophoretic mobility (form ®) in a manner that is destroyed by ring cleavage. Cohesin is known to associate with chromatin prior to DNA replication, when it cannot be involved in holding sisters together. Moreover, it can associate with replicated chromosomes in a manner that does not lead to cohesion between sisters ^{12,14}. We suggest that cohesin frequently does trap individual chromatin fibres and that its activity in postmitotic cells in metazoan might involve this type of action ^{15,16}. If we are correct in concluding that cohesin is a novel type of concatenase, then it is not implausible to imagine that other SMC-kleisin complexes such as condensin and its bacterial equivalent have related activities. Indeed, the deep

evolutionary roots of these types of complexes suggest that the ability to concatenate DNA may have been an activity without which DNA genomes could not have evolved.

Online Methods

Yeast strains

All strains are derived from W303. Genotypes are listed in Supplementary Table 1.

Model of the S. cerevisiae Smc1/Smc3 hinge structure

A structure model of the yeast Smc1/Smc3 hinge structure was created with the Modeller program ¹¹ using an alignment of *S. cerevisiae* Smc1 aa residues 488-690 and Smc3 aa residues 496-699 with aa residues 475-679 of the *T. maritima* SMC protein and the coordinates of pdb file 1GXL.

Expression, purification and cross-linking of cohesin subunit domains

Sequences encoding aa 494-705 of the *S. cerevisiae* Smc3 hinge domain followed by an internal ribosome binding site and sequences encoding aa 486-696 of the *S. cerevisiae* Smc1 hinge domain fused to a C-terminal His₆ tag were cloned by PCR into the pET28 expression vector. Cysteine mutations were introduced by overlap extension PCR. The Smc1/Smc3 hinge domains were co-expressed in *E. coli* strain BL21(DE3)-RIPL (Stratagene) at 20°C for 5 h after induction with 0.25 mM IPTG. Cells were lysed in 50 mM NaP_i pH8.0, 300 mM NaCl containing Complete EDTA free protease inhibitor mix (Roche) and the complex was purified via Ni²⁺-chelating affinity chromatography followed by gelfiltration on a Superdex 200pg 26/60 column (GE Healthcare) in TEN buffer (20 mM TRIS-HCl pH8.0, 1 mM EDTA, 1 mM NaN₃) +100 mM NaCl +2 mM DTT. The Smc1 head domain bound to Scc1-C was expressed in insect cells using the baculovirus system and purified as described ¹².

Purified proteins were re-buffered into reaction buffer (25 mM NaP_i pH7.4, 50 mM NaCl, 10 mM MgSO₄, 0.25% Triton X-100) via a Superdex G-25 column, adjusted to 0.5 mg/ml and mixed quickly into 1/25th volume of DMSO, 5 mM bBBr (Sigma), or 25 mM BMOE (Pierce). Both cross-linkers were dissolved in DMSO just before use. After 10 min incubation at 4°C, sample loading buffer containing β-mercaptoethanol was added, the samples were heated for 3 min at 90°C and run on an SDS-PAGE followed by Coomassie blue staining. Cross-linking reached a maximum after a few minutes at 4°C.

Minichromosome preparation and cross-linking

Yeast strains containing the 2.3 kbp minichromosome were grown, arrested in nocodazole and lysed by spheroplasting as described ⁷, with the exception that sodium citrate and sodium sulfite in the lysis buffer were replaced by 300 mM NaCl to increase minichromosome yield. Extracts were loaded onto an SW41 10-30% sucrose gradient in 25 mM HEPES-KOH pH8.0, 50 mM KCl, 10 mM MgSO₄, 0.25% Triton X-100, 1 mM DTT,

1 mM PMSF. Gradients were run for 15 h at 18,000 rpm and fractionated. Fraction aliquots were separated on a 1% agarose gel containing 0.5 μ g/ml ethidium bromide as described ⁷. Gels were transferred under alkaline conditions by capillary blotting onto Immobilion-NY+ membrane (Millipore). The blots were hybridized with a ³²P-labelled probe for the 2.3 kbp minichromosome sequence, exposed to imaging plates, scanned on an FLA-7000 image analyzer (Fujifilm) and quantified using ImageQuant.

Minichromosome monomer or dimer peak fractions (~300 μl) were dialysed for 4 h against 500 ml reaction buffer at 4°C in a Float-a-lyzer (SpectraPor) with 100 kDa MW cut-off. The dialysis buffer was replaced three times. 24 μl dialysed fraction were mixed quickly into 1 μl DMSO, 5 mM bBBr, or 25 mM BMOE (both freshly dissolved in DMSO) and incubated at 4°C for 10 min. Final concentrations of 200 μM bBBr or 1 mM BMOE were optimal for cross-linking (Supplementary Fig. S7). Bis-malemide based cross-linkers with longer spacers than BMOE, like BMB or BMH (Pierce), could be used with similar efficiency (data not shown). The reaction was quenched by the addition of 1.25 μl 210 mM DTT. For TEV cleavage, 24 μl of the quenched cross-linking reaction was mixed with 1μl 5 mg/ml wild-type or C151A mutant TEV protease in TEV buffer (TEN buffer +50 mM NaCl +2 mM DTT) or TEV buffer only and incubated at 30°C for 1 h. Protein was denatured for 4 min at 65°C after the addition of 2.8 μl 10% SDS. The denatured samples were mixed with 3 μl 80% sucrose containing 0.02% bromophenol blue and 20-25 μl of the mixture were loaded onto a 0.8% agarose gel containing 0.5 μg/ml ethidium bromide. Gels were run at 4°C for 14 h at 1.4 V/cm and blotted and hybridized as before.

For 2D gels, lanes from the first dimension agarose gels were cut out and placed at the top of a second 0.8% agarose gel (20×20 cm) containing 0.5 μ g/ml ethidium bromide, leaving an approximately 5 mm wide slot between the lane and the gel. The slot was filled with 60°C warm 0.8% agarose in TAE. Proteinase K was dissolved in TAE and mixed with the pre-warmed agarose solution to a final concentration of 0.2 mg/ml just before casting. Second dimension gels were run at 4°C for 8 h at 2 V/cm, blotted and hybridized as before.

For nicking minichromosome DNA, 200 μ l cross-linked samples were first dialysed against 500 ml reaction buffer +2 M KCl +1 mM DTT at 4°C for 4 hours to remove nucleosomes followed by 2 h dialysis against reaction buffer + 1mM DTT to remove salt and 2 h dialysis against reaction buffer +20% sucrose +1 mM DTT to re-concentrate the samples. Dialysis buffers were replaced every hour. 1 μ l *Nb.BsrDI* (10 U/ μ l, NEB) was added to 27 μ l sample followed by 10 min incubation at 50°C, addition of SDS to 1% and denaturation as above.

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Figure legends

Figure 1. Making covalently closed cohesin rings. a Experimental strategy: fusion of the Smc3 C-terminus with the Scc1 N-terminus and chemical cross-linking of engineered cysteine residues at the Smc1/Smc3 and Smc1/Scc1 interfaces creates a covalently closed cohesin ring. Circular sister DNAs should remain cohesed even after protein denaturation if and only if they are trapped inside cohesin's ring. **b** A homology model of the Smc1/Smc3 hinge interface to the homodimeric bacterial structure (PDB 1GXL) identifies two juxtaposed Cβ atoms in Smc1's helix H11 and Smc3's helix H6 at a distance compatible with cross-linking when mutated to cysteine. Wild-type and cysteine mutant yeast Smc1/Smc3 hinge domain dimers purified after recombinant expression in E. coli were incubated with DMSO, or final concentrations of 200 µM bBBr, or 1 mM BMOE, denatured and run on SDS-PAGE followed by Coomassie Blue staining. c The structure of the yeast Scc1 C-terminus bound to Smc1's head domain (PDB 1W1W) identifies two juxtaposed side-chains that should allow cross-linking when mutated to cysteine. Purified complexes of wild-type and cysteine mutant yeast Smc1 head domain bound to Scc1's Cterminal domain after recombinant expression in insect cells were incubated with DMSO, 200 µM bBBr, or 1 mM BMOE, denatured and run on SDS-PAGE followed by Coomassie Blue staining. The low level crosslinking observed for the Smc1/Scc1(A547C) combination likely results from a reaction of BMOE with the engineered cysteine in Scc1 and the nearby ε-amino group of Smc1's lysine residue K20.

Figure 2. Covalent cohesin circularization creates SDS-resistant minichromosome **DNA dimers. a** Extracts from yeast strains harbouring the minichromosome and expressing wild-type cohesin (top) or Smc3-TEV-Scc1 and Smc1 containing the engineered cysteine pairs (bottom) were separated by gradient centrifugation followed by native gel electrophoresis and Southern blotting. Minichromosome dimers sediment faster but electrophorese slower than monomers. The fraction of dimers is reduced by the Smc3-TEV-Scc1 fusion. **b** Monomer or dimer gradient fractions from yeast strains in which all three (K14856; panels A and B), only two (K14857, K14859; C and D), one (K14858; E), or no (K14860; F) ring subunit interface(s) can be covalently linked were dialysed against reaction buffer, treated with DMSO, bBBr, or BMOE, denatured with 1% SDS at 65°C, run on an agarose gel, and minichromosome DNA was detected by Southern blotting. Monomer and dimer fractions contain supercoiled (1) and nicked (2) monomeric minichromosomes. Dimer fractions also contain supercoiled-supercoiled (3) and supercoiled-nicked (4) concatenated minichromosome DNAs. Only samples in which all three cohesin ring subunit interfaces have been covalently linked contain additional slower migrating bands, presumably corresponding to supercoiled-supercoiled (⑦) and supercoiled-nicked (®) cohesed minichromosomes. c Input fractions for cross-linking

reactions A-F were run without denaturation on an agarose gel and Southern blotted. **d** Treatment of non-cross-linked or cross-linked minichromosomes with nicking enzyme after removal of nucleosomes by high salt converts supercoiled (①), supercoiled-supercoiled (③, ⑦) or supercoiled-nicked (④, ⑧) minichromosomes to nicked (②) or nicked-nicked (⑤, ⑨) forms. **e** Gradient dimer fractions from a topoisomerase II mutant strain (K15029) grown at the restrictive temperature were denatured with SDS. Concatenated minichromosomes (short exposure) co-migrate with bands ② and ③ of K14856 (long exposure) on the same agarose gel. **f** Schemata of minichromosome conformations.

Figure 3. Cohesin's covalent circularization holds individual DNAs together. a Crosslinked and denatured samples were run on an agarose gel, lanes cut out, and run on a second agarose gel in perpendicular direction. A thin slot between the first dimension lane and the second gel was filled with either agarose or agarose containing 0.2 mg/ml proteinase K. In the absence of protease, all bands run on a diagonal. In the presence of protease, supercoiled-supercoiled cohesed dimers (⑦) run as supercoiled monomers (⑦→①) in the second dimension, while supercoiled-nicked cohesed dimers (⑧) split up into supercoiled ($\$ \rightarrow \textcircled{1}$) and nicked ($\$ \rightarrow \textcircled{2}$) monomers in the second dimension, more clearly visible in a longer exposure (panel below). A faint band presumably corresponding to cohesin bound supercoiled monomers (6) is converted into naked supercoiled monomers (⑥→①) upon protease cleavage. An additional band (⑩) running off the diagonal presumably corresponds to supercoiled-supercoiled concatenated minichromosomes that had bound cohesin. **b** Cross-linked minichromosome samples were treated with or without TEV protease before SDS protein denaturation, agarose gel electrophoresis and Southern blotting. The slower migrating bands (②, ⑧) corresponding to cohesed minichromosomes disappear upon TEV incubation if the Smc3-Scc1 linker contains triple target sequences for the protease (TEV cleavable), but not if the -1 sites of the triple target sequence have been mutated to lysine (TEV resistant).

Figure 4. Minichromosomes are cohesed by single cohesin rings. a Protein-protein cross-linking efficiencies by bBBr and BMOE were measured by spiking dimer fractions (K14856) with purified double cysteine mutants of Smc1/Smc3 hinge or Smc1/Scc1-C complex preparations at a final concentration of 3 μM (saturated association of the nanomolar affinity interactions between the two domains). After cross-linking, half of the reactions were run on an SDS-PAGE and band intensities were measured after SYPRO Ruby staining. **b** The other half of the reactions were run on an agarose gel and the fractions of cohesed minichromosomes (⑦+®/total) were quantified after Southern blotting. **c** Minichromosome dimer fractions from haploid strains expressing Smc3-Scc1 fusions containing TEV target sequences (K14856) or resistant to TEV cleavage (K15089),

from a 1:1 mixture of the two haploid strains, or from a diploid strain containing one TEV cleavable and one TEV resistant SMC3-SCC1 allele (K15267) were cross-linked with bBBr, treated with wild-type or a non-catalytic (nc; C151A) mutant of TEV protease, denatured, run on an agarose gel and Southern blotted. Lane profiles were recorded, the peaks corresponding to cohesed supercoiled minichromosome dimers (\bigcirc) integrated and the peak area after incubation with active versus non-catalytic TEV protease compared. In case the number of cohesin cross-bridges follows a Poisson distribution, we would expect 53% or 27% of cross-linked dimers surviving TEV treatment for single or double ring models, respectively. The observed value is $54\% \pm 6\%$ (mean of three independent experiments \pm standard deviation).

Supplementary Information

Minichromosome cohesion depends on centromeres

To confirm that dimeric minichromosomes detected in sucrose gradients ⁷ reflect minichromosome cohesion detected *in vivo* ¹⁷, we tested whether their generation depends on centromeres. To do this, we created a yeast strain containing a minichromosome whose *CEN4* sequences were flanked by target sites for the site-specific *Zygosaccharomyces rouxii* recombinase (Supplementary Fig. S1a), which was expressed from the galactose-inducible *GAL1* promoter. An exponential culture growing in non-inducing raffinose medium was split and galactose added to one half and glucose to the other. Both sets of cells were arrested in G1 by α factor and then transferred to pheromone-free medium containing nocodazole, which caused them to undergo DNA replication and arrest in a mitotic state. Due to leaky expression of the recombinase, only half of the minichromosomes in glucose grown cells contained *CEN4* prior to DNA replication (Supplementary Fig. S1b). Due to efficient deletion, few if any contained *CEN4* in galactose grown cells.

Minichromosomes containing *CEN4* sequences, either because they contained no recombinase sites or because they had not undergone recombination between them, sediment as dimers and monomers (Supplementary Fig. S1c, top and middle panels). In contrast, minichromosomes whose *CEN4* sequences had been deleted prior to DNA replication sediment almost exclusively as monomers (Supplementary Fig. S1c, lower panel), suggesting that *CEN4* is required for the formation and/or maintenance of minichromosome cohesion. If so, all dimeric minichromosomes from glucose grown cells, which contained equal numbers of intact and deleted minichromosomes, should possess *CEN4* sequences. To test this, monomer and dimer fractions were run on an agarose gel after removal of associated proteins by SDS heat denaturation. This showed that most DNAs migrated as supercoiled circular monomers. Strikingly, almost all DNAs from the dimer fraction had not undergone *CEN4* deletion while most DNAs from the monomer fraction had done so (Supplementary Fig. S1d). We conclude that minichromosomes containing *CEN4* can produce (and maintain) cohesin-mediated dimers during (and after) DNA replication but those lacking *CEN4* cannot.

Calculation of the number of cross-bridges following a Poisson distribution

Following the Poisson distribution, the distribution of *x* of cross-bridges between minichromosomes will be

(1)
$$f(x;\lambda) = \frac{\lambda^x e^{-\lambda}}{x!}$$

f(0) can be determined from the fraction of minichromosomes without cross-bridges (x=0), e.g. the fraction of monomers plus the fraction of dimers that are held together by concatenation (but not by cross-bridges). The percentage of monomers can be directly quantified from gradient blots and is approximately 78% (Supplementary Fig. 4). The fraction of concatemers in the dimer fractions can be directly measured after relaxation of non-crosslinked DNAs with nicking enzyme (Fig. 3d). Alternatively, the ratio of nicked to supercoiled DNA can be measured from monomer fractions (Fig. 4a, panel A, 2/(1+2)). This number can then be used to calculate the proportion of nicked (2) and supercoiled-supercoiled concatemers (3) in the dimer fractions (Fig. 4a, panel B), yielding the percentage of concatemers (3) in the dimer fractions (Fig. 4a, panel B), yielding the dimers are concatemers. We estimate that approximately $78\% + 10\% \cdot 22\% = 80\%$ of minichromosomes are not cohesed, therefore $2 = -\ln 0.8$. The distribution then is

fraction of total circles fraction of cohesed circles
$$x=0 f(0;-ln0.8)=80.0\%$$

$$x=1 f(1;-ln0.8)=17.8\% f^*(1)=f(1;-ln0.8)/(1-f(0;-ln0.8))=89.3\%$$

$$x=2 f(2;-ln0.8)=2.0\% f^*(2)=f(2;-ln0.8)/(1-f(0;-ln0.8))=10.0\%$$

$$x=3 f(3;-ln0.8)=0.1\% f^*(3)=f(3;-ln0.8)/(1-f(0;-ln0.8))=0.7\%$$

$$\vdots \vdots \vdots$$

We then calculated the fraction of dimers p that are held together by at least one SDS-resistant cross-bridge using the measured cross-linking efficiency e=30% at both Smc1-Smc3 and Smc1-Scc1 interfaces (Fig. 5c):

single ring model double ring model
$$x=1$$
 $p(1) = e = 0.30$ $p'(1) = e^2 = 0.09$ $x=2$ $p(2) = 1-(1-e)^2 = 0.51$ $p'(2) = 2e^2 - e^4 = 0.17$ $x=3$ $p(3) = 1-(1-e)^3 = 0.66$ $p'(3) = 3e^2-3e^4+e^6 = 0.25$ \vdots $\Sigma p(x)\cdot f^*(x) = \textbf{0.32}$ $\Sigma p'(x)\cdot f^*(x) = \textbf{0.10}$

In a diploid strain expressing one TEV cleavable and one TEV non-cleavable allele (fraction of non-cleavable molecules n=50%) of the Smc3-Scc1 linker, the fraction of dimers with at least one non-cleaved cross-bridge will be

single ring model double ring model
$$x=1$$
 $p''(1) = n = 0.50$ $p'''(1) = n^2 = 0.25$ $x=2$ $p''(2) = 1-(1-n)^2 = 0.75$ $p'''(2) = 2n^2 - n^4 = 0.44$ $x=3$ $p'''(3) = 1-(1-n)^3 = 0.88$ $p'''(3) = 3n^2-3n^4+n^6 = 0.58$ \vdots $\Sigma p'''(x)\cdot f^*(x) = \mathbf{0.53}$ $\Sigma p'''(x)\cdot f^*(x) = \mathbf{0.27}$

Supplementary Methods

Centromere loop-out

Cells were inoculated into YEP + 2% raffinose (YEPR) from an overnight culture in –TRP + 2% raffinose and grown at 30°C to OD₆₀₀=0.6. Galactose (gal) or glucose (glc) was added to a final concentration of 2%. After 30 min cultures were diluted to OD₆₀₀=0.15 and α factor added to a final concentration of 2 µg/ml. Additional α -factor was added to 1.5 µg/ml each after two 40 min intervals. Cells were collected by centrifugation, washed with four culture volumes YEPD, resuspended in YEPD containing 10 µg/ml nocodazole and grown for 1.5 h at 30°C. Genomic DNA was prepared as described in the Supplemental Materials and digested with *EcoRI*. Cleavage fragments were resolved on a 0.8% agarose gel, Southern blotted and probed for the *TRP1* gene.

Genomic DNA preparation

Yeast cells from 25-30ml asynchronous culture were resuspended in 200µl SCE buffer (1 M sorbitol, 0.1 M sodium citrate pH 7.0, 60 mM EDTA) +0.1 M β -ME + 1 mg/ml zymolyase T-100) and incubated for 1 h at 37°C with occasional shaking. Spheroplasts were lysed by addition of 200 µl SDS lysis buffer (2% SDS, 0.1M TRIS-HCl pH 9.0, 50 mM EDTA) and incubation for 5 min at 65°C followed by addition of 200 µl 5 M potassium acetate and centrifugation at 16,000 × g after 20 min incubation on ice. 450 µl supernatant was mixed with 1 ml isopropanol and 200 µl 5 M ammonium acetate and precipitated genomic DNA was pelleted by 30 sec centrifugation at 2300 × g. All excess liquid was removed and the pellet was dissolved by 1 h incubation in 90 µl TE at 37°C. 10 µl 5 M ammonium acetate and 200 µl isopropanol were added and genomic DNA was pelleted as above. The DNA pellet was washed with 1 ml 70% ethanol and dissolved in 50 µl TE containing 0.2 mg/ml RNaseA. For restriction digest, 5 µl were used in a 20 µl reaction.

Preparation of concatenated minichromosomes from a top2 strain

Strain K15029 was grown at 23°C to log phase in –TRP + 2% raffinose media. Cultures were diluted to OD_{600} =0.15 and α -factor was added in three steps to final concentrations of

2, 1.5 and 1.5 μ g/ml, respectively, in 40 min intervals at 23°C. Cells were collected by centrifugation and α -factor was removed by washing with four culture volume YEPR at 4°C. Cells were resuspended in 35.5°C warm YEPR containing 10 μ g/ml nocodazole and grown for 1.5 h at 35.5°C. Cells were harvested and genomic DNA was prepared by spheroplasting as described above, with the exception that all steps including lysis were performed at 37°C using buffers pre-warmed to 35.5 °C. Cleared lysates were loaded on pre-chilled 10-30% sucrose gradients and processed as described above.

Purification of catalytically active and inactive TEV protease

Wild-type and catalytically inactive mutant (C151A) TEV protease ¹⁸ fused to an N-terminal His₆ tag were expressed from the pET9d vector in *E. coli* strain BL21(DE3)-RIPL (Stratagene) after induction with 0.25 mM IPTG for 6 h at 22°C. Cells were lysed in 50 mM TRIS-HCl pH8.0, 1 mM β-mercaptoethanol, 1 mM PMSF. TEV protease was purified from the clarified extract via Ni²⁺-chelating affinity chromatography followed by gelfiltration on a Superdex 200pg 26/60 column (GE Healthcare) in TEN buffer (20mM TRIS-HCl pH 8.0, 1 mM EDTA, 1 mM NaN₃) +50 mM NaCl +2 mM DTT. TEV protease eluted as a single band at the expected delay volume and was concentrated to 5 mg/ml by ultrafiltration.

Testing cleavage of the Smc3-Scc1 linker

Protein extracts from 50 ml asynchronous cultures were prepared by glass bead lysis and the Smc3-Scc1 fusion protein was immunoprecipitated via its C-terminal HA₆ epitope tag with 16B12 antibody as described previously 6 . Immunoprecipitation beads were split an incubated in TEV buffer containing 0.1 mg/ml TEV protease or TEV buffer only for 2 h at 16°C before addition of loading buffer, SDS-PAGE and Western blotting. The blot was probed with 3F10 antibody (Roche) against the HA₆ epitope and exposed to film.

Supplementary Table

Table 1. Yeast strains.

- K14856 MATalpha, smc3::HIS3, smc1::kanMX, scc1::kanMX, leu2::SMC1(G22C, K639C)-myc9::LEU2, ura3::pSCC1-SMC3(E570C)-TEV3-SCC1(A547C)-HA6::URA3, [2.3 kbp TRP1-ARS1-CEN4]
- K14857 *MATa*, *smc3::HIS3*, *smc1::kanMX*, *scc1::kanMX*, *leu2::SMC1(G22C)-myc₉::LEU2*, *ura3::pSCC1-SMC3-TEV3-SCC1(A547C)-HA₆::URA3*, [2.3 kbp TRP1-ARS1-CEN4]
- K14858 MATa, smc3::HIS3, smc1::kanMX, scc1::kanMX, leu2::SMC1-myc9::LEU2, ura3::pSCC1-SMC3-TEV3-SCC1-HA6::URA3, [2.3 kbp TRP1-ARS1-CEN4]
- K14859 *MATalpha, smc3::HIS3, smc1::kanMX, scc1::kanMX, leu2::SMC1(K639C)-myc9::LEU2, ura3::pSCC1-SMC3(E570C)-TEV3-SCC1-HA6::URA3, [2.3 kbp TRP1-ARS1-CEN4]*
- K14860 *MATa*, scc1::SCC1(TEV3)-HA₆::HIS3, [2.3kbp TRP1-ARS1-CEN4]
- K15029 MATa, top2-4, [2.3 kbp TRP1-ARS1-CEN4]
- K15089 MATalpha, smc3::HIS3, smc1::kanMX, scc1::kanMX, leu2::SMC1(G22C, K639C)-myc9::LEU2, ura3::pSCC1-SMC3(E570C)-noTEV-SCC1(A547C)-HA6::URA3, [2.3 kbp TRP1-ARS1-CEN4]
- K15249 *MATalpha*, *smc3*::*HIS3*, *scc1*::*kanMX*, *smc1*::*kanMX4*, *leu2*:*SMC1*-*myc*₉::*LEU2*, *ura3*::*pSCC1*-*SMC3*(*E570C*)-*TEV3*-*SCC1*(*A547C*)-*HA*₆::*URA3*, [2.3 kbp TRP1-ARS1-CEN4]
- K15250 MATalpha, smc3::HIS3, scc1::kanMX, smc1::kanMX4, leu2:SMC1(K639C)-myc9::LEU2, ura3::pSCC1-SMC3(E570C)-TEV3-SCC1(A547C)-HA6::URA3, [2.3 kbp TRP1-ARS1-CEN4]
- K15251 MATalpha, smc3::HIS3, smc1::kanMX, scc1::kanMX, leu2::SMC1(G22C, K639C)-myc₉::LEU2, ura3::pSCC1-SMC3(E570C)-TEV3-SCC1(A547C)-HA₆::URA3, [2.3 kbp TRP1-ARS1-CEN4]
- K15252 *MATa*, *smc3*::*HIS3*, *scc1*::*kanMX*, *smc1*::*kanMX4*, *leu2*::*SMC1*(*G22C*)-*myc9*::*LEU2*, *ura3*::*pSCC1*-*SMC3*(*E570C*)-*TEV3*-*SCC1*(*A547C*)-*HA*₆::*URA3*, [2.3 kbp TRP1-ARS1-CEN4]
- K15301 MATa, mhra::RS-HMR-EI-RS, leu2::pGAL1-RecR::LEU2 ×2, [2.6 kbp TRP1-ARS1-RS-CEN4-RS]
- K15351 MATa, leu2::pGAL1-Rec::LEU2, [2.3kbp TRP1-ARS1-CEN4]
- K15267 MATa/MATalpha, smc3::HIS3/smc3::HIS3, smc1::kanMX4/smc1::kanMX4, scc1::kanMX/scc1::kanMX, leu2::SMC1(G22C, K639C)-myc9::LEU2/leu2::SMC1(G22C, K639C)-myc9::LEU2, ura3::pSCC1-SMC3(E570C)-TEV3-SCC1(A547C)-HA6::URA3/ura3::pSCC1-SMC3(E570C)-noTEV-SCC1(A547C)-HA6::URA3, [2.3 kbp TRP1-ARS1-CEN4]

All strains are *ade2-1, trp1-1, can1-100, leu2-3,112, his3-11,15, ura3, GAL, psi*+ unless noted otherwise.

Supplementary Figure Legends

Figure S1. Minichromosome cohesion requires centromeres. a Induction of Z. rouxii recombinase leads to loop-out of the minichromosome centromere region flanked by recombination sites (RS). b Recombinase expression was induced by galactose (+gal) or repressed by glucose (+glc) addition to synchronous cultures of yeast strains harbouring minichromosomes with centromeres flanked by RS sites (K15301) or without RS sites (K15351) before arresting cells in G1 by α-factor addition. Cells were released from the pheromone arrest into nocodazole containing medium to re-arrest them in a mitotic state. To estimate the efficiency of *CEN4* loop-out, genomic DNA was prepared from asynchronous cultures, 30 min after glucose or galactose addition, or after cells had arrested in α-factor or nocodazole. DNA was linearized with *EcoRI* and probed against the TRP1 sequence after Southern blotting. Filled circles indicate minichromosomes that still contain CEN4, open circles indicates minichromosomes that have looped out CEN4. A star marks a restriction fragment of the genomic TRP1 locus. The fraction of minichromosomes that have excised CEN4 was quantified and blotted. c Extracts from the nocodazole arrested strains were fractionated by sucrose gradient centrifugation. Gradient fractions were electrophoresed through native agarose gels and minichromosome DNA detected by Southern blotting. Open triangles indicate monomeric and closed triangles indicate dimeric minichromosomes. **d** DNA from monomer (open bars) and dimer (filled bars) minichromosome gradient fractions were deproteinized by heating for 5 min in 1% SDS, separated by agarose gel electrophoresis, and Southern blotted to measure the fraction of minichromosomes that have looped-out CEN4.

Figure S2. Covalent connection of adjacent cysteine residues. Reaction schemes for cross-linking juxtaposed thiol groups with **a** dibromobimane (bBBr) and **b** bismaleimidoethane (BMOE).

Figure S3. SDS-resistant dimer formation after cross-linking requires all four cysteine substitutions at the Smc1/Smc3 and Scc1/Smc1 interfaces and is lost upon DNA linearization. a Extracts were prepared from strains expressing the Smc3(E570C)-TEV3-Scc1(A547C) fusion protein containing cysteines in the Smc3 hinge domain the Scc1 C-terminus but lacking both (K15249), either one (K15252, K15250), or none (K15251) of the cysteines in Smc1's hinge and head domains from nocodazole arrested or asynchronous cells and run on sucrose gradients. Dimer fractions were isolated, treated with DMSO, bBBr, or BMOE and analysed by Southern blotting after protein denaturation. Cohesed minichromosome bands (②) were only detectable when all four cysteines are present. **b** Monomer or dimer fractions from a strain (K14856) whose cohesin rings could be covalently circularized were treated with DMSO, bBBr, or BMOE,

denatured and run on an agarose gel after incubation with *BgIII* restriction enzyme or buffer only. No slower migrating cross-linked dimer bands can be observed after DNA linearization, consistent with the idea that a topological association between circularized cohesin rings and minichromosomes requires intact rings and DNA circles.

Figure S4. Salt sensitivity of native minichromosome dimers. Minichromosome monomer and dimer gradient fractions from a yeast strain expressing Scc1 with a triple tandem repeat of TEV target sites (K14860) were dialysed for 12 h against reaction buffer containing the indicated concentrations of KCl before removal of high salt by dialysis against reaction buffer without KCl and incubation with TEV protease or buffer only. About 50% of dimers are converted to monomers after incubation with 0.5 M KCl and 100% with 1 M KCl or more.

Figure S5. Alternative models for the topological connection of minichromosomes by cohesin rings. a The single ring model envisions that minichromosomes are held together by their topological entrapment within individual cohesin rings. b The double ring model proposes the entrapment of each minichromosomes within separate cohesin rings that are connected by intertwinning.

Figure S6. Quantification of minichromosome monomer fraction and Smc3-Scc1 TEV cleavage efficiency. a Gradient fractions from a yeast strain expressing the Smc3-TEV-Scc1 fusion and cysteine pairs at both Smc1/Smc3 and Smc1/Scc1 interfaces (K14856) were run on an agarose gel and Southern blotted. Signal intensities were quantified using a phosphorimager and plotted as percentage of total signal. b Extracts were prepared from haploid strains expressing a single copy of an Smc3-Scc1 fusion protein connected by a TEV-cleavable (K14856) or TEV-resistant (K15089) linker, or from a diploid strain expressing both alleles as single copies (K15267). The fusion proteins were immunoprecipitated via their C-terminal HA₆ tag an the immunoprecipitation beads were incubated with TEV protease or buffer only. The beads were boiled in SDS loading buffer, proteins resolved by SDS-PAGE and the gel was immunoblotted against the HA epitope.

Figure S7. Optimization of cross-linker concentrations. Dimer fractions from a yeast strain expressing the Smc3-TEV-Scc1 fusion and cysteine pairs at both Smc1/Smc3 and Smc1/Scc1 interfaces (K14856) were incubated with increasing concentrations of BMOE (top) or bBBr (bottom) after DTT had been removed by dialysis. The reaction was quenched after 10 min by re-addition of DTT and samples were denatured with 1% SDS, resolved on a 0.8% agarose gel and Southern blotted. Cross-linking efficiencies (⑦/total) are plotted against cross-linker concentration (log-scale). Optimal cross-linking occurred at 1 mM BMOE or 200 μM bBBr.













