Clamping of DNA shuts the condensin neck gate

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Condensin is a structural maintenance of chromosomes (SMC) complex needed for the compaction of DNA into chromatids during mitosis. Lengthwise DNA compaction by condensin is facilitated by ATP-driven loop extrusion, a process that is believed to be the fundamental activity of most, if not all, SMC complexes. In order to obtain molecular insights, we obtained electron cryomicroscopy structures of yeast condensin in the presence of a slowly hydrolyzable ATP analog and linear as well as circular DNAs. The DNAs were shown to be “clamped” between the engaged heterodimeric SMC ATPase heads and the Ycs4 subunit, in a manner similar to previously reported DNA-bound SMC complex structures. Ycg1, the other non-SMC subunit, was only flexibly bound to the complex, while also binding DNA tightly and often remaining at a distance from the head module. In the clamped state, the DNA is encircled by the kleisin Brn1 and the two engaged head domains of Smc2 and Smc4. The Brn1/Smc2/Smc4 tripartite ring is closed at all interfaces, including at the neck of Smc2. We show that the neck gate opens upon head engagement in the absence of DNA, but it remains shut when DNA is present. Our work demonstrates that condensin and other SMC complexes go through similar conformations of the head modules during their ATPase cycle. In contrast, the behavior of the Ycg1 subunit in the condensin complex might indicate differences in the implementation of the extrusion reactions, and our findings will constrain further mechanistic models of loop extrusion by SMC complexes.

Significance

DNA needs to be compacted to fit into nuclei and during cell division, when dense chromatids are formed for their mechanical segregation, a process that depends on the protein complex condensin. It forms and enlarges loops in DNA through loop extrusion. Our work resolves the atomic structure of a DNA-bound state of condensin in which ATP has not been hydrolyzed. The DNA is clamped within a compartment that has been reported previously in other structural maintenance of chromosomes (SMC) complexes, including Rad50, cohesin, and MukBEF. With the caveat of important differences, it means that all SMC complexes cycle through at least some similar states and undergo similar conformational changes in their head modules, while hydrolyzing ATP and translocating DNA.

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dissociation from DNA requires opening of the Smc3–kleisin (Sccl) neck interface, which depends on an accessory protein called Wapl and also ATP binding (9, 10).

In vivo cysteine cross-linking analyses of cohesin (11, 12), SMC-ScpAB (13, 14), and MukBEF (15) have been used to reveal topological compartments in SMC complexes in which circular DNA can be entrapped (SI Appendix, Fig. S1A). These include the SMC-Kleisin (S-K) compartment and, when the head domains are engaged, the Engaged heads-Kleisin (E-K) and the Engaged heads-SMC (E-S) compartments. Sister chromatid cohesion is mediated by the coentrapment of sister chromatids within cohesin’s S-K compartment. In contrast, it is not currently clear in which compartment the DNAs reside during loop extrusion, or if in any (3, 16).

Cryo-EM structures of cohesin in the ATP-bound engaged (E) state, and bound to one strand of DNA, revealed that DNA is “clamped” on top of the dimerized SMC ATPase domains by Scc2 and the closed neck of Smc3 (17–19). Analysis of the same cohesin–DNA interaction using in vitro cysteine cross-linking revealed that the DNA is not entrapped within the S-K ring but threads through both the E-S and E-K compartments (17). This strongly suggests that in the clamped state of cohesin the kleisin runs over the top of the DNA and that the DNA has not passed through one of the trimer interfaces to reach S-K. Furthermore, a recent structure of MukBEF in the ATP and DNA-bound state conclusively shows that the clamped DNA passes through the E-K and E-S compartments because the entire kleisin MukP is resolved (15). Interestingly, in this structure, the ε-SMC-kleisin neck interface is not completely closed; however, the MatP DNA-unloading protein was also bound.

We aimed to investigate how condensin interacts with DNA in the ATP engaged state and how the DNA interaction might regulate the state of the neck interface. To address this, we started by using cryo-EM to solve the structure of condensin in the presence of DNA, ADP, and BeF3.

**Results**

**Cryo-EM Structure of a “Clamped” Condensin Complex.** To reveal the structure of the condensin–DNA complex we purified a “tetrameric” budding yeast condensin complex lacking the Ycg1 subunit (SI Appendix, Fig. S1B, containing subunits Smc2, Scm4, Brn1, and Ycs4). We mixed the condensin tetramer with 80-bp double-stranded DNA (dsDNA), ADP, and beryllium fluoride (BeF3−), applied the sample to cryo-EM grids, and collected images (SI Appendix, Fig. S2A). ABC-type ATPases bound to ADP,BeF3 often represent the enzyme–substrate complex (20). Due to the apparent flexibility of the SMC coiled coil and hinge parts in our data, we were unable to perform analysis of the entire complex (SI Appendix, Fig. S2B) but instead focused on the head module (Fig. 1B and SI Appendix, Fig. S3, Left). Following the picking of particles with a focus on the SMC head domains, 702,764 good particles showing densities of SMC heads, Ycs4, and DNA were selected with two-dimensional (2D) classifications. Through further processing we reconstructed two cryo-EM maps (Forms I and II) at resolutions of 2.95 and 3.05 Å, with 286,294 and 251,999 particles, respectively (Fig. 1C and SI Appendix, Fig. S2 C and D). The maps enabled us to build and refine reliable atomic models from both (SI Appendix, Table S1). A comparison of their conformations showed strong similarity between the two atomic models, with a root-mean-squared deviation of 1.2 Å (Fig. 1D and SI Appendix, Fig. S2E).
In both structures, Form I and II, the SMC head domains and proximal coiled coils, Ycs4 and Brn1 form a compact globular structure, the head module. The majority of the SMC coiled coils and hinge are unresolved, most likely due to flexibility with respect to the resolved parts (Fig. 1B and SI Appendix, Fig. S2B). In both structures, the DNA is held between the upper surface of the dimerized SMC head domains and a groove in Ycs4 (Fig. 1C). The map allowed us to trace a significant fraction of the Brn1 kleisin subunit (279/754, 37% of residues in Brn1), which binds to the neck region of Smc2, to Ycs4 via an extended interaction surface and the cap of the Smc4 head domain (Fig. 1D). The neck interface between Smc2 and the N-terminal domain of the kleisin is closed in our structure, possibly indicating that the DNA-bound clamped conformation prevents ATP-dependent opening of this interface, since it had been reported previously that the Smc2/kleisin neck gate opens upon ATP binding in the absence of DNA (8). Forms I and II differ in the DNA density where they are not clamped between Ycs4 and the SMC heads. While the DNA of Form I traverses the entire surface of the SMC heads, Form II shows less-well-resolved DNA density in the region distal to Ycs4 (SI Appendix, Fig. S2 C–E). Because Forms I and II therefore differ only slightly, we describe the structure based on Form I, unless otherwise noted (Fig. 1 C and D).

Structure of ATPase Head Domains and Interactions with DNA. Condensin contains two ATPase sites, with each SMC head domain possessing the Walker A and Walker B motifs from one shared or “sandwiched” active site and the “ABC signature motif” from the other. In the structure, the head domains are dimerized, with two ADP·BeF₃ sandwiched in the interface (Fig. 2A). Each active site contains a network of interactions between the ADP·BeF₃, a Mg²⁺ ion and highly conserved residues in Smc2 and Smc4. Engagement of the SMC head domains produces a pseudo-two-fold symmetrical DNA-binding surface on their upper side, which is bordered by the two coiled coils (Fig. 2B). Approximately 18 bp of the DNA traverse this surface, interacting with a large number of mostly positively charged residues (SI Appendix, Fig. S4A). Interestingly, the DNA is not exactly centered on the pseudo-two-fold axis of the ATPase heads (Fig. 2B) and therefore additionally interacts with the coiled coil Smc2 neck and not with the coiled coil of Smc4 (Fig. 2B). The N-terminal domain of Brn1, bound to the neck of Smc2, is also close to the DNA in our structure.

Ycs4 is a large hook-shaped protein consisting of 21 HEAT repeats and a protrusion called the “probosisc” (8) (SI Appendix, Fig. S4C). In our cryo-EM map, we observed two clear protein densities running along the inside and around the tip of the hook (SI Appendix, Fig. S4C). The first density clearly corresponds to a part of Brn1 that has been identified earlier in the cryo-EM structure of apo condensin and also a Ycs4–Brn1 crystal structure (residues 184 to 223) (8), while we assigned the second density to a region of Brn1 that had been identified biochemically but had not been resolved earlier (residues 275 to 323).

The DNA is clamped onto the top of the dimerized heads by Ycs4. On Ycs4, the DNA binds to a positively charged...
surface formed by the edge of a series of HEAT repeats near the middle of the protein, and the surface includes residues K211, K292, K300, K377, R416, K420, and R556 (Fig. 2C and SI Appendix, Fig. S4B). At the other end of Ycs4, and at the other end of the DNA, the tip of the Ycs4 hook also binds the backbone of the DNA through an arginine residue (R1122).

Ycs4 forms extensive contacts with Smc2 and Smc4 at four interfaces (Fig. 2D). The first is a tripartite interaction between Ycs4’s N-terminal HEAT repeats 3 and 4, the neck of Smc2, and the upper end of Brn1’s helix α3 (Fig. 2E). This arrangement traps Brn1’s α3 between Ycs4 and Smc2, possibly preventing opening of the Smc2–Brn1 interface, which, as has been mentioned, has been reported to occur during head engagement in the absence of DNA (8). The second interaction site of Ycs4 is with a loop emanating from the side of the Smc2 head domain (residues 69 to 73) that also contacts an alpha helix from Brn1 (residues 189 to 196) (Fig. 2F). The third interaction is large and is between a region in Ycs4 toward the C terminus (residues ~1010 to 1094) and a beta-sheet on the back of the Smc4 head domain (Fig. 2G). Finally, an unstructured and not-well-conserved region near the N terminus of Smc4 (127 to 131) forms the fourth Ycs4 interface, using the outside of the Ycs4 hook (Fig. 2H and SI Appendix, Fig. S4F).

Having structures of condensin in multiple states (apo, apo-bridged, and ATP-clamped) (7) (Protein Data Bank [PDB] ID codes 6YVU, 6YVV, and this study) allowed us to analyze the conformational changes that accompany them. The structure of Ycs4 is similar in the apo and clamped structures, but with a 10-Å outward movement of the C-terminal HEAT repeats, which results in a slight widening of the hook (SI Appendix, Fig. S4E, Left). In the apo and apo-bridged conformations, the association between Smc4 and Ycs4 is similar, but the distance between the SMC heads is enlarged in the apo-bridged state, and Smc2 binds to the N-terminal HEAT repeats of Ycs4 (SI Appendix, Fig. S4E, Middle). The relative orientation of Ycs4 and the SMC heads is similar. However, in the clamped state there is a 55° rotation of Ycs4 relative to the SMC heads when compared to both apo and apo-bridged (SI Appendix, Fig. S4E, Right).

**In the Clamped State, Ycg1 Binds DNA but is Only Loosely Attached to the Head Module of Condensin.** A previous cryo-EM structure of condensin “pentameric” holocomplex (subunits Smc2, Smc4, Brn1, Ycs4, and Ycg1) showed that in the absence of DNA Ycg1 binds to Smc2 in the ATP-engaged state (7). To determine whether Ycg1 also interacts with the head module in the DNA-bound, clamped state we mixed the condensin pentamer (SI Appendix, Fig. S1C) with 80-bp DNA, ADP, and BeF3 as before and examined by cryo-EM (SI Appendix, Fig. S5A). Using our tetramer structure to guide the analysis, we were able to solve two separate structures after extensive classifications (SI Appendix, Figs. S3, Right and S5B).

First, we obtained a map of the larger subcomplex comprising the condensin head module, which was almost indistinguishable from the one determined for the tetramer. This 3.7-Å resolution map (processed as a single map due to particle number limitations: 45,112 particles; SI Appendix, Fig. S3, Right)
demonstrated clearly that Ycg1 does not associate rigidly with Smc2 or any other subunit within the globular head module of the complex (SI Appendix, Fig. S5 B and C).

Second, from the same dataset, we solved the structure of Ycg1 bound to DNA (Fig. 3 A and B) at 3.2-Å resolution. In this structure, the region of Brn1 that binds Ycg1 and makes up the “safety belt” (residues 387 to 524) (21) was mostly resolved, with the exception of a disordered connecting loop (residues 411 to 458) (Fig. 3C). The DNA is somewhat bent as it passes over the DNA-binding surface of Ycg1 and through the safety belt. The DNA covers most of Ycg1’s positively charged surface, and we could identify more Ycg1–DNA interactions than a previous study using X-ray crystallography, which used a shorter DNA (SI Appendix, Fig. S5 D and E). The fact that we observed Ycg1 bound to Brn1 while not being bound to the head module of condensin strongly suggests that in condensin’s clamped state Ycg1 remains attached to the rest of the complex via the flexible linkers provided by Brn1 either side of its interaction site with Ycg1 (Fig. 3A).

The Path of the Brn1 Subunit Relative to DNA. The path of DNA through SMC complexes is important because it determines whether circular (or very long) DNA is topologically entrapped. In previous clamped structures of cohesin, little of the kleisin subunit Scc1 was resolved and the path was inferred and validated using cysteine cross-linking experiments (17). In the present structure, substantially more of the kleisin chain is visible, which helps to determine whether or not the DNA runs through, and is entrapped by, the tripartite ring of Smc2–Smc4–Brn1 (S-K) (Fig. 3D). As described above, in our structure the N-terminal domain of Brn1 (22 to 106) is bound to the neck of Smc2. Brn1 then snakes along the concave surface of Ycs4 (Brn1: 163 to 223) and along the C terminus of the protein (Brn1: 275 to 323). The next resolved Brn1 residues bind to Ycg1 and form the safety belt around DNA (Brn1: 459 to 525), before Brn1’s C-terminal winged-helix domain binds to the cap of Smc4 (Brn1: 526 to 747). While several regions remain unresolved, the position of Brn1 residues 106 and 163 indicate that the Brn1 chain most likely passes above the DNA. This means that the DNA runs at least through the E-K compartment (SI Appendix, Fig. S1A).

Interactions between Condensin and Circular DNA in the Clamped State. The condensin–DNA complexes investigated so far in this study were assembled using 80-bp double-stranded linear DNA. Therefore, the clamped head module and Ycg1 were possibly bound to separate DNA molecules and any topological constraints were overridden by the fact that linear DNA was used. To examine the arrangement of the condensin holocomplex when all subunits have the opportunity to bind to the same strand of DNA, we mixed the condensin pentamer, nicked (relaxed) plasmid DNA (1.7 kb), ADP, and BeF3, and examined the sample using cryo-EM with a Volta phase plate (VPP) to increase imaging contrast (Fig. 4A and SI Appendix, Fig. S6). In the individual cryo-EM images, without averaging, we observed complexes of condensin where the main body and Ycg1 were most likely bound to the same plasmid. Even without averaging, the plasmid DNA could be seen passing though the head module and also through Ycg1 in the same manner as revealed by our higher-resolution structures (Figs. 1C and 3B, respectively).

We then generated 2D class averages and three-dimensional (3D) models of the clamped condensin head module and Ycg1, each bound to circular DNA (Fig. 4 B and C). While these structures are at lower resolutions due to smaller particle numbers, they showed the same conformations and that the DNA is also entrapped within the E-K compartment. Given that the DNA is circular, it is not possible that the DNA slid into the E-K compartment and therefore there are two possibilities for how the DNA enters: 1) One of the three interfaces in the SMC-kleisin ring opens and the DNA is topologically entrapped within this tripartite ring (E-K and S-K) or 2) the DNA passes between the disengaged heads before they engage ATP (E-K and E-S). In this second model, the DNA becomes entrapped within both the E-S and E-K compartments simultaneously but never enters the S-K ring. It is important to point out that since the position of the coiled coils and hinge domains could not be resolved in the clamped state, E-S and S-K entrapment cannot be deduced from our head module structure alone (Fig. 4D) and additional topological interactions with either E-S (as deduced for cohesin) (17) or with S-K are conceivable. S-K entrapment could be indicated by the opening of the coiled coils in our clamped structure, but it requires gate opening as has been proposed before (18).

The Smc2–Brn1 Neck Interface Remains Shut in the Clamped State. It has been reported previously that ATP binding, in the absence of DNA results in opening of the Brn1 (N-terminal domain) and Smc2 neck interface (8), and also the equivalent interface between N-Scc1 and the Smc3 neck in cohesin (9, 10). In support of this, in nucleotide-bound and SMC head-engaged structures of cohesin (22) and condensin (7), both in the absence of DNA, the N-terminal domain (NTD) of the kleisin is not present, whereas in nucleotide-free and non-engaged structures the kleisin’s NTD is bound (7, 23).

Furthermore, a biochemical experiment with recombinant condensin, known as the N-terminal kleisin release assay, has been used to demonstrate that ATP binding opens this interface (8). Recognition sequences for Tobacco Etch Virus (TEV) protease were inserted into an unstructured region of Brn1 (after residue 141). After cleavage of Brn1 with TEV protease, in the absence of ATP, the Brn1 N-terminal fragment coimmunoprecipitated with the rest of the complex, reporting a closed interface. However, when ATP was included during the wash steps, the N-terminal fragment did not coimmunoprecipitate with the complex, indicating that ATP stimulates the disengagement of the interface. Furthermore, the N-terminal fragment was released in a mutant that can bind but not hydrolyze ATP but was abolished in a mutant that cannot bind ATP. This showed that ATP binding, but not its subsequent hydrolysis, stimulates opening of the Smc2–Brn1 neck interface. Similar experiments have demonstrated that this mechanism is conserved in cohesin at the Smc3–Scc1 neck interface (10).

Surprisingly, in our clamped structure the Smc2–Brn1 interface is closed despite engagement of the ATPase heads and nucleotide binding (Fig. 4E, Left). Two possibilities could explain this finding: 1) Unlike ATP, binding of ADP·BeF3 does not stimulate release or 2) binding of DNA and being in the clamped conformation inhibits release. To test these hypotheses, we used the same N-terminal Brn1 release assay (Fig. 4E, Right). We found that, as previously described, in the absence of nucleotide the N-terminal fragment of Brn1 is efficiently coimmunoprecipitated. When ATP was included during the immunoprecipitation wash steps, the N-terminal fragment no longer coimmunoprecipitated, confirming that ATP stimulates release. Likewise, when we washed the beads with buffer containing ADP·BeF3 the N-terminal fragment did not coprecipitate effectively. This demonstrates that as with ATP head
engagement stimulated by ADP·BeF₃ results in the disengagement of the Smc2-Brn1 neck interface. In contrast, when the beads were washed in buffer containing both 80-bp dsDNA and ADP·BeF₃, the N-terminal fragment was efficiently coprecipitated. Together, these experiments showed that condensin can adopt two conformations during nucleotide-driven head engagement: In one the Smc2-Brn1 interface is open, as shown in the cryo-EM structure of condensin bound only to ATP or AMP-PNP (7), and in the other, reported here, it is firmly shut (SI Appendix, Fig. S7 A and B). It seems reasonable to state that the only determinant of which conformation it adopts appears to be the presence or absence of DNA.

Discussion

A mechanism by which condensin transforms interphase chromosomes into mitotic chromatids has recently been proposed: Condensin extrudes chromosomal fibers, while hydrolyzing ATP, to create an array of DNA loops spanning the entire chromosome (4). However, the molecular basis for this extraordinary motor activity remains enigmatic, despite a significant number of experiments and models (24). In this paper we present the structure of condensin in complex with DNA and ADP·BeF₃, a slowly hydrolyzable mimic of ATP. This has revealed several important behaviors of condensin during the ATPase cycle. First, condensin adopts...
a “clamped” conformation in which DNA is bound between the engaged SMC head domains, the Ycs4 subunit and the Smc2 neck domain. Second, the DNA passes through the E-K compartment between the kleisin and the head domains. Third, the interface between Smc2 and Brn1 is closed. The latter is surprising because head engagement was thought to open this interface (8). We also present biochemical evidence that this ATP-stimulated interface opening is inhibited by the presence of DNA.

The clamped state of condensin is closely related to the clamped state of cohesin ([SI Appendix, Fig. S7 C and D]) (17–19), highlighting potential deep mechanistic similarities between the two protein complexes. In cohesin, the Scc2 cohesin “loader” subunit takes the position of Ycs4 in clamped condensin and both complexes show very similar arrangements around the neck gates (Smc2–Brn1; Smc3–Scc1), with the kleisins’ N-terminal domains wedged in between the SMC neck, the DNA and the clamping subunit. It is therefore not surprising that neck gate opening seems to depend on the absence of DNA and the presence of ATP in both complexes (8, 22). Because the clamped state of MukBEF is analogous to the clamped state of cohesin ([SI Appendix, Fig. S7E]) (15), it is also related to condensin and together with the finding that SbcCD (Mre11–Rad50), an SMC-like protein involved in DNA repair also binds DNA in this way (PDB ID code 6S85) (25), it seems that DNA clamping is a conserved state of these complexes and is likely involved in their most fundamental functions.

Taken together with published structures (7), our data demonstrate that condensin can adopt at least four stable conformations: two very different conformations in the presence of nucleotide (ATP-engaged [SI Appendix, Fig. S7A, Right] and DNA-clamped), depending on the presence of DNA, and the two nucleotide-free apo conformations (apo and apo-bridged [SI Appendix, Fig. S4E, Left and Middle]) (7). In the apo conformation, Ycs4 is bound to the SMC heads, which are juxtaposed with their ATPase active sites facing away from each other. In the apo-bridged conformation, the heads are forced apart by Ycs4 which sits between them. In both states Ycg1 is bound to the complex only via the flexible Brn1 linkers. In the ATP-bound structure without DNA (ATP-engaged), the heads are engaged but now Ycs4 is no longer bound to the main body whereas Ycg1 tightly binds to the Smc2 head and the Smc2-Brn1 neck interface is open ([SI Appendix, Fig. S7A, Right]). Finally, in the ATP- and DNA-bound clamped structure, the head domains are engaged around nucleotides, however Ycs4 is associated with the main body rather than Ycg1—and the Smc2-Brn1 neck interface is closed (this study).

It is conceivable that the different conformations between the ADP-BeF3 + DNA and the ATP or AMP-PNP structures (Smc2–Brn1 interface opening and Ycg1/Ycs4 binding to SMC heads) are a result of the different nucleotides that were used to engage the head domains rather than the presence of DNA. This is unlikely for three reasons. First, in previous structural studies of ABC-type ATPases, heads engaged with different nucleotide analogs result in almost identical conformations of these domains (26). Second, in the equivalent clamped structure of cohesin, ATP was used in combination with mutations (EQ Walker B) that result in defective ATP hydrolysis but not binding (17). Despite using ATP rather than ADP-BeF3, the proteins adopted the same clamped conformation and the Smc3–Scc1 interface was closed. Third, our Brn1 release assay showed no obvious difference between the effects of ATP or ADP-BeF3 on neck gate opening.

Our structure raises an interesting question about the significance of the previous ATP-bound structure of condensin without DNA. Does condensin adopt two distinct ATP-bound conformations as part of the loop extrusion process? If not, what is the relevance of ATP binding when not in the clamped state? One possible explanation is indicated by the state of the Smc2–Brn1 neck interface in each structure. In the clamped state this interface is shut, whereas in the DNA-free structure the Brn1 N-terminal domain cannot be resolved, likely indicating that it is open. This finding is further supported by our biochemical evidence showing that ATP binding releases the Brn1 N-terminal domain, unless DNA is included in the reaction. Because opening of this interface compromises the integrity of the tripartite condensin ring, it is possible that the ATP-bound state represents a hypothetical unloading reaction. Condensin needs to be able to dissociate from chromatin in a regulated manner, and opening of the Smc2–Brn1 interface is one likely mechanism for that release, especially if loop extrusion or other cellular activities of condensin involve either topological or pseudo topological entrapment of DNA (16, 24). Indeed, unloading through the Smc3–Scc1 interface in cohesin (9, 10), whose topological entrapment in cells is firmly established (11), is similarly driven by ATP binding (10, 22). Finally, a recent structure of MukBEF, in complex with the MatP unloader, which is believed to show the complex poised for topological unloading of DNA, shows the analogous N-kleisin neck gate ([MukF-MukB] in an open state ([SI Appendix, Fig. S7B]) (15).

It is also conceivable that ring opening at the neck interface is an integral part of the DNA motor activity since DNA may not stay bound to the head domains during the entire translocation cycle. However, recent in vitro single-molecule experiments have demonstrated that both condensin and cohesin are capable of loading onto DNA and loop extruding when the neck interface is permanently closed (3, 27). This is consistent with the finding that closure of the equivalent cohesin or SMC–Scc1D interfaces results in functional complexes in yeast and Bacillus subtilis, respectively (11, 28).

Our cryo-EM data provide clear evidence that condensin can bind to DNA via Ycg1 and the clamped head module simultaneously. Based on our structures and published single-molecule experiments (4) we propose that this is an intermediate step in the loop extrusion process and the variable distance of Ycg1 with respect to the head module might determine or allow condensin to make steps on DNA that are needed to extrude loops and to enlarge them. It seems also worth pointing out that the two binding sites in the clamped head module and on Ycg1 enable the trapping of preformed DNA loops as shown directly in Fig. 4A, and this would be a convenient initial reaction to start loop extrusion. In this scenario, Ycg1 is likely to act as a static anchor on the DNA, which implies that the clamped DNA is being translocated during loop extrusion, consistent with the regulation of DNA clamping by ATP turnover.

However, it is clear that a precise understanding of the order of the conformations during the loop extrusion cycle will be required to be able to describe the molecular mechanism for DNA translocation and extrusion. This may be achieved using single-molecule approaches such as fluorescence resonance energy transfer (FRET), especially for the presumably very flexible SMC arms (29), but we envisage that resolving these structures of SMC complexes during the loop extrusion process by cryo-EM may ultimately be needed to understand the mechanisms involved at the same level of detail that has been achieved for cytoskeletal or other nucleic acid motor proteins, as long as the important and relevant parts of the complex retain enough rigidity for this to be achievable or can be made visible at least partly.
Materials and Methods

Plasmids and Protein Expression. Wild-type condensin holocomplex (pentamer) was overexpressed in budding yeast from two 2 μm high-copy-number plasmids (pGAL7 SM4-3xStrepII pGAL10-SMC2 pGAL1 BRN1-HA3-His12 URA3 and pGAL1 YCG1 pGAL10 YCS4 TRIP1) (30). Condensin holocomplex containing TEV cleavable Bm1 was overexpressed from pGAL7 SM4-3xStrepII pGAL10-SMC2 pGAL1 BRN1 Ybjb5 tag replacing residues 13 to 23, 3 x TEV site inserted at residue 141)HA3-His12 URA3 and pGAL1 YCG1 pGAL10 YCS4 TRIP1.

For expression of the condensin tetramer, YCG1 was deleted by Gibson assembly to produce pGAL1 YCG1 TRP1. Cultures were grown in -URA -TRP dropout medium + 2% raffinose until optical density at 600 nm (OD600) of 0.8 to 1. Protein expression was induced by adding 2% galactose to the medium and incubating overnight at 30 °C.

Protein Purification. Recombinant condensin complexes were purified as previously described, with minor modifications (30, 31). Induced yeast extracts were centrifuged, washed once in phosphate-buffered saline, and centrifuged again. The cell pellet was resuspended in 1x protein volume of Buffer A (50 mM Tris·HCl, pH 8.0, 200 mM NaCl, 5% glycerol, and 5 mM 2-mercaptoethanol) with protease inhibitors (Roche) and 300 μL benzonase (Sigma). The cell suspension was then lysed in a Spex Freezer Mill (Cryogenic) with protease inhibitors (Roche) and 300 U/L benzonase (Sigma). The cell suspension was then lysed in a Spex Freezer Mill (Cryogenic) and incubating overnight at 30°C to 1. Protein expression was induced by adding 2% galactose to the medium and incubating overnight at 30 °C.

Assembling Clamped Condensin and Cryo-EM Grid Preparation. Purified condensin samples were buffer-exchanged using Zeba Micro Spin 7K MWCO columns (Thermo Fischer Scientific) in TNT buffer (30 mM Tris·HCl, 60 mM NaCl, 1 mM TCEP, and 2 mM MgCl2, pH 7.5). Then, 0.5–1 mg/mL samples were incubated with 1–2 μM 80-bp dsDNA (5’-GAATTCGGTGCGCATAATGTATAATAAGATAAA-TAAGCTTAATGGTCTCCGGATCTAATACGACTCACTACGTTCTAC-3’, and 5’-GGAAATCCCTATACCCTGTAGATGAAGAAGGATATACGTTCTAC-3’). EDTA was added to a final concentration of 75,000 μM, followed by 50 μL Buffer SB. The protein was then eluted in buffer SB containing 5 mM deoxyribonuclease (DNase) (IBA). The peak fractions were pooled and concentrated with a Vivaspin 20 100,000 Da centrifugal concentrator (Sartorius).

For fluorescent labeling of TEV cleavable condensin, at this stage CoA-LD555 (Lumdyne) was conjugated to Bm1-ybjb5 with recombinant SYP synthase (Addgene Plasmid 75015) as previously described (32). All proteins were further purified by size-exclusion chromatography on a Superose 6 Increase 10/300 GL column (Cytiva), preequilibrated with Buffer SB containing 1 mM MgCl2. The peak fractions were concentrated using a Vivaspin 20 100,000 Da centrifugal concentrator and frozen.

Cryo-EM Image Acquisition. All cryo-EM datasets were acquired at 300 kV on a FEI Titan Krios electron microscope (Thermo Fischer Scientific). EPU was used for automatic data collection, and for non-VPP datasets AFIS (aberration-free image shift) was used to increase throughput. For the condensin tetramer:80-bp DNA dataset, images were acquired using a GIF Quantum energy filter with 20-eV slit width and a 100-μm objective aperture; 8,784 movies were recorded with a Gatan K3 summit direct electron detector in supersolution mode, with a nominal magnification of 81,000x, corresponding to a pixel size of 1.07 Å per pixel (0.535 Å per pixel in superresolution) and a nominal defocus range of 1.5–3.3 μm. Each movie was dose-fractionated into 55 frames, with a total dose of 55 e−/Å2. For the pentamer:80-bp DNA dataset, 4,302 images were collected using a Falcon 4 detector, with a nominal magnification of 75,000x (calibrated pixel size 1.08 Å) and without any objective aperture. The nominal defocus range was set to 1.5–3.0 μm. Each image was recorded in EER (electron event representation) format with a total dose of 40 e−/Å2. For the pentamer:circular DNA dataset, 3,179 images were acquired with a VPP (defocus range: 0.6–1.1) using a Falcon 4 detector, with a nominal magnification of 75,000x (pixel size = 1.08 Å). Since the number of condensin particles in this VPP dataset was not sufficient to obtain good 3D classes, an additional 1,016 non-VPP images were acquired with a 100-μm objective aperture (defocus range: 1.5–3.0 μm), using a grid obtained under the same conditions but with a more concentrated sample. Both datasets were recorded in EER format with a total dose of 32 e−/Å2.

Cryo-EM Image Processing. The cryo-EM data processing workflow is summarized in SI Appendix, Fig. S3. Processing was performed with RELION 3.1, CtfFind4, cryOLO, and cryoSPARC v3.2 (34–37), and RELION was used, unless otherwise specified. Overall resolutions were determined based on Fourier shell correlation gold-standard criteria (0.143) (38). All images were subjected to beam-induced motion correction as implemented in RELION 3.1. Movie frames were aligned and combined with dose weighting using 7 × 5 patches (for K3 datasets) or 5 × 5 patches (for Falcon 4 datasets). The Falcon 4 movies in EER format were dose-fractionated into groups of 40 frames, corresponding to a dose of 1 e−/Å2 or 0.8 e−/Å2 per frame for the pentamer:80-bp DNA or pentamer:circular DNA datasets, respectively. CTF parameters were estimated with CtfFind4.

For the condensin tetramer:80-bp DNA dataset, ~3 M particles were picked with a Laplacian-of-Gaussian blob as template and subjected to 2D classification. An initial 3D model of the head complex was generated using particles from selected 2D class images showing different orientations. Then, to obtain more particles accurately using cryOLO, a model was trained with the coordinates from the images that formed the selected 2D classes. The resulting ~2 M particles were extracted using a box size of 3002 pixels (pixel size = 1.07 Å), followed by 2D classification; 702,764 accepted particles were subjected to further 3D classification and resulted in two 3D classes showing secondary structure features. Further processing was performed separately for each class, but in the same way. First, 3D autorefinement was performed and resulted in maps of 3.27 Å (Form I) and 3.24 Å (Form II), followed by CTF refinement for magnification anisotropy, per-particle defocus, per-micrograph astigmatism, and beam tilt. For beam tilt and CTF refinement, the particles were divided into optics groups according to their hole positions (because of AFIS data collection). Then, Bayesian polishing was performed, followed by another round of 3D refinement to generate final maps with 2.95 and 3.05 Å resolutions for Forms I and II, respectively.

For the condensin pentamer:80-bp DNA dataset, particles were initially picked using cryOLO with a general model. Approximately 600,000 particles were picked and a subset of 50,000 particles was subjected to a 3D classification. An initial 3D model of the head complex was generated using particles from selected 2D class images showing different orientations. Then, to obtain more particles accurately using cryOLO, a model was trained with the coordinates from the images that formed the selected 2D classes. The resulting ~2 M particles were extracted using a box size of 3002 pixels (pixel size = 1.07 Å), followed by 2D classification; 702,764 accepted particles were subjected to further 3D classification and resulted in two 3D classes showing secondary structure features. Further processing was performed separately for each class, but in the same way. First, 3D autorefinement was performed and resulted in maps of 3.27 Å (Form I) and 3.24 Å (Form II), followed by CTF refinement for magnification anisotropy, per-particle defocus, per-micrograph astigmatism, and beam tilt. For beam tilt and CTF refinement, the particles were divided into optics groups according to their hole positions (because of AFIS data collection). Then, Bayesian polishing was performed, followed by another round of 3D refinement to generate final maps with 2.95 and 3.05 Å resolutions for Forms I and II, respectively.

For the condensin pentamer:80-bp DNA dataset, particles were initially picked using cryOLO with a general model. Approximately 600,000 particles were picked and a subset of 50,000 particles was subjected to a five-class ab initio initial model reconstruction in cryoSPARC v3.2. Using the resulting five models, heterogeneous refinement was performed for all picked particles in cryoSPARC. The classes showing features of “clamped” head modules or Ycg1–DNA complexes were selected, and the corresponding particles for each class were subjected to cryoSPARC model training separately, followed by automated particle picking. By cryOLO picking with each model, 476,997 particles for the head module and 640,437 particles for Ycg1–DNA were obtained, respectively. The particles for each model were first cleaned up by cryoSPARC heterogeneous refinement using five models from the first ab initio reconstruction. Additional 3D and 2D classifications without alignment in RELION resulted in 45,112 particles for the head module and 91,024 for Ycg1–DNA. The particles for each class were reextracted with box sizes of 3202 and 3002 pixels (pixel size = 1.08 Å), respectively. Three-dimensional autorefinement performed for each particle followed by CTF refinement (magnification anisotropy, per-particle defocus, per-micrograph astigmatism, and beam tilt) and Bayesian polishing. After 3D refinement, maps at 3.68 Å and 3.20 Å resolutions were obtained for the head complex and Ycg1–DNA, respectively.

For condensin pentamer:circular DNA, particle picking was separately done for the two datasets, VPP and non-VPP, using cryOLO with a general model, followed by extraction with a box size of 3602 pixels with 2x binning (1802-pixel...
size, pixel size = 2.16 Å). The particles from the two datasets were then merged, an ab initio reconstruction was obtained, followed by multiple rounds of heterogeneous refinement in cryoSPARC. 36,588 and 27,040 particles corresponding to the head module and Ycg1-DNA, respectively, were selected, and subjected to 3D homogeneous refinement in cryoSPARC to generate 8.75- and 8.95-Å maps for the head complex and Ycg1-DNA, respectively.

During processing of the dataset, we also discovered 2D classes presumed to be SMc coiled-coil arms or the hinge (SI Appendix, Fig. S2B), however, we were not able to obtain reliable 3D maps of these parts, presumably due to their flexibility.

Model Building and Refinement. For atomic model building, the maps of the two Forms I and II of the head module from the tetramer dataset and the Ycg1-DNA complex from the pentamer dataset were further improved using DeepEMHancer (39). Model building was carried out in Coot (40) and ISOLDE (41). Coordinates were refined using phenix.real_space_refine (42). Model validation was performed with MolProbity (43). The atomic model for the head module was first built into the 2.95 Å cryo-EM density of Form I from the tetramer:30-bp DNA dataset. The models of Smc2 head-Brn1(125–108), Smc4 head-Brn1(43–68,685–747), and Ycs4-Brn1(156–174,184–195,199–220) were taken from the apo-condensin cryo-EM structure from Saccharomyces cerevisiae (PDB ID code 6YVU) (7) and used as templates. The DNA model was initially taken from the S. cerevisiae cohesin cryo-EM structure (PDB ID code 6Z26) (17). Each atomic model was docked into the EM map using UCSF Chimera (44). Coordinates were then manually adjusted and rebuilt in Coot. The DNA model was further refined using the annealing function in ISOLDE. Clear map densities in the vicinity of HEAT repeats 14 to 18 of Ycs4 were apparent after building using the secondary structure predictions and cryoID (45), and then built de novo in Coot. The N- and C-terminal low-resolution regions of Ycs4(26–92 and 119–116)–Brn1 were built using ab initio models of S. cerevisiae Ycs4 generated by AlphaFold2 (https://alphafold.ebi.ac.uk/entry/Q06156) as a template (46). For Form II of the head module from the tetramer dataset the refined model of Form I was docked into the 3.05 Å map and rigid-body-refined in PHENIX, followed by manual adjustments in Coot. For the Ycg1-DNA atomic model, the crystal structure of the Ycg1-Bm1 complex (PDB ID code 5OOQ) (21) was docked into the Ycg1-DNA map from the pentamer dataset in Chimera, and a DNA model was flexibly fitted using ISOLDE. The model was then manually adjusted and rebuilt in Coot, followed by real-space refinement in PHENIX. Figures and movies were generated with PyMOL 2.5 (Schrodinger), UCSF Chimera, and ChimeraX (47). Electrostatic potentials were calculated and displayed in PyMOL.

Bm1 N-Terminal Release Assay. The N-terminal release assay was performed as previously described with modifications (8). Twenty-five micrograms of

condensin holocomplex containing the Bm1 TEVs (after residue 141) and labeled with LDS55 (Lumidone) (1 μg/mL), 1x acEV buffer (Thermo), and 20 U acEV protease (Thermo) was incubated at 4 °C for 16 h to cleave Bm1.

For each nucleotide condition, 20 μL of protein A magnetic dynabeads (Thermo) were washed twice in 500 μL wash buffer (50 mM Tris·HCl pH 7.5, 125 mM NaCl, 50 mM KCl, 5 mM MgCl2, and 5% glycerol, 1 mM DTT, 0.2 mM phenylmethylsulfonyl fluoride, and 0.01% [vol/vol] Tween-20). The beads were resuspended in 300 μL wash buffer and 3 μL of anti-HA tag antibody (Roche) was added and mixed at 4 °C for 1 h. The antibody-bound beads were washed two times and resuspended in 300 μL wash buffer. Five micrograms of cleaved Bm1-condensin was added to the beads per condition and mixed for 1 h at 4 °C. The beads were washed three times in wash buffer followed by three washes in 300 μL wash buffer, wash buffer with 1 mM ATP, wash buffer with ADP·BeF3 (0.5 mM ADP, 0.5 mM BeSO4, and 10 mM NaF), or wash buffer with 80–bp dsDNA and ADP·BeF3 (10 μM DNA was added and mixed before addition of 0.5 mM ADP, 0.5 mM BeSO4, and 10 mM NaF). The beads were then suspended in 50 μL 2× sodium dodecyl sulfate (SDS) (100 mM Tris·HCl, pH 6.8, 4% [wt/vol] SDS, 20% glycerol [vol/vol], 0.2% [wt/vol] bromophenol blue, and 0.2 M DTT) buffer and incubated at 65 °C for 5 min to elute the proteins. The eluate was resolved by SDS polyacrylamide gel electrophoresis, the retained Bm1 N-terminal fragment was visualized on a Typhoon Scanner (Amersham), and immunoprecipitation of condensin and equal loading was confirmed by Coomassie staining of the eluate.

Data Availability. The model coordinates and cryo-EM density maps were deposited in the Protein Data Bank (PDB) and Electron Microscopy Data Bank (EMDB) with the following accession codes: cleaved head module from tetramer dataset, Form I (PDB ID 7Q2X and EMD-13783) and Form II (PDB ID 7Q2Y and EMD-13784); cleaved head module from pentamer dataset (EMD-13785) and Ycg1-DNA (PDB ID 7Q2Z and EMD-13786); cleaved head module from circular DNA dataset (EMD-13787) and Ycg1-DNA (EMD-13788). See SI Appendix, Table S1 for details.

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