The relative ratio of condensin I to II determines chromosome shapes

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To understand how chromosome shapes are determined by actions of condensins and cohesin, we devised a series of protocols in which their levels are precisely changed in Xenopus egg extracts. When the relative ratio of condensin I to II is forced to be smaller, embryonic chromosomes become shorter and thicker, being reminiscent of somatic chromosomes. Further depletion of condensin II unveils its contribution to axial shortening of chromosomes. Cohesin helps juxtapose sister chromatid arms by collaborating with condensin I and counter-acting condensin II. Thus, chromosome shaping is achieved by an exquisite balance among condensin I and II and cohesin.

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Upon entry into mitosis, a mass of chromatin distributed within the interphase nucleus is converted into a discrete set of rod-shaped structures. This process, often referred to as mitotic chromosome condensation, culminates in the assembly of metaphase chromosomes in which two sister chromatids are juxtaposed with each other along their entire lengths. It is thought that chromosome condensation is a critical preparatory step to ensure faithful segregation of duplicated chromosomes. Cohesin helps juxtapose sister chromatid arms by collaborating with condensin I and counteracting condensin II. Thus, chromosome shaping is achieved by an exquisite balance among condensin I and II and cohesin.

Results and Discussion

Condensin I and II differently contribute to the assembly of duplicated chromosomes in Xenopus egg extracts

To dissect spatiotemporal regulation of condensins and cohesin, we exploited a cell-free system derived from Xenopus eggs that recapitulate a series of cell cycle-regulated chromosomal events. It should be noted that this study is the first to describe the behaviors of condensin I and II in the cell-free system using duplicated DNA substrates (hereafter referred to as the duplicated chromosome assembly assay). In this assay, sperm chromatin was first duplicated by incubating with an interphase extract for 100 min, and converted into metaphase chromosomes by adding a nondegradable form of Xenopus cyclin B into the extract. At various time points after cyclin B addition, chromatin was isolated and analyzed by immunoblotting and immunofluorescence with antibodies for condensin I- and II-specific subunits, and cohesin subunits (see Supplemental Figs. S1A for the subunit composition of these complexes). Association of condensin I with chromosomes became prominent after nuclear envelope breakdown (NEBD), which occurred ~20 min after cyclin addition [Fig. 1A; Supplemental

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To evaluate how condensin I and II might contribute to the assembly of metaphase chromosomes in this assay, we performed immunodepletion experiments (Fig. 1C; see also Supplemental Fig. S1C). When both condensin I and II were depleted from an extract, no individual chromosomes were discernible and only fuzzy masses of chromatin were observed (Δcond I and II). An extract depleted of condensin I alone produced fuzzy, yet slightly ordered, masses of chromatin, in which condensin II were concentrated on axial structures as well as at the centromere/kinetochore regions (Δcond I). In striking contrast, individual chromosomes with a reasonably good morphology were assembled in a condensin II-depleted extract (Δcond II). Thus, unlike in HeLa cells (Ono et al. 2003), the functional contribution of condensin II was much smaller than that of condensin I in this cell-free system. Our previous study had estimated the relative ratio of condensin I to II to be ~5:1 in Xenopus egg extracts and ~1:1 in nuclear extracts from HeLa cells (Ono et al. 2003), providing a reasonable explanation for the differential contribution of condensin I and II to chromosome condensation between Xenopus egg extracts and HeLa cells.

The relative ratio of condensin I to II is a critical determinant of chromosome shapes

It is known that metaphase chromosomes in early embryonic cells tend to be longer and thinner compared with those in somatic cells (Belmont et al. 1987; Micheli et al. 1993), and the chromosomes assembled in Xenopus egg extracts recapitulate the embryonic properties (Fig. 1B,C). One attractive possibility would be that the relative ratio of condensin I to II contributes to such structural differentiation between embryonic and somatic chromosomes. We therefore sought to test what would happen if the ratio of condensin I to II in the extracts (5:1) were converted to 1:1. It should be emphasized that such precise manipulation of protein levels is virtually impossible in any other experimental systems. To achieve this in our cell-free assay, one volume of a mock-depleted extract was mixed with four volumes of a condensin I-depleted extract to reconstitute an extract containing the equal level of condensin I and II (Supplemental Fig. S2A,B). Remarkably, chromosomes assembled in the 1:1 extract indeed displayed a characteristic appearance reminiscent of somatic chromosomes, being shorter than chromosomes assembled in the mock-depleted extract (5:1) (Fig. 2A). The chromosomes assembled in the 5:1 and 1:1 extracts had lengths of 13.3 ± 4.9 μm [average ± standard deviation] and 9.3 ± 3.3 μm, respectively [Fig. 2B]. The width of the chromosomes was apparently greater in the 1:1 extract than in the 5:1 extract (Fig. 2A, right panels). One could argue, however, that the 1:1 extract produced the short and thick chromosomes simply because the absolute level of condensin I present in the extract was reduced (not because the relative ratio of condensin I to II was altered). To exclude this possibility, we prepared another extract in which the absolute levels of both condensin I and II were reduced to 20% of the original level while their relative ratio was kept...
condensin II primarily contributes to axial shortening of chromosomes, and that this action of condensin II is rather cryptic when the action of condensin I is predominant under the standard condition.

Functional interplay between cohesin and condensins in chromosome shaping

Despite some descriptive observations implicating cohesin in metaphase chromosome structures [Gimenez-Abian et al. 2004; Nakajima et al. 2007; Shintomi and Hirano 2009], no systematic approach had been taken to address this problem. In particular, we wished to test how cohesin’s action might potentially be coordinated with the action of condensins. To this end, we established a protocol in which the levels of cohesin and condensin I were partially reduced individually or in combinations [Supplemental Fig. S4A]. To clearly visualize the chromosome axes regardless of the level of condensins targeted, an antibody against topoisomerase II was used for immunofluorescence analysis in this experiment (Fig. 2A). We found that, under the condition where condensin I existed dominantly over condensin II (4:1), reducing the cohesin level from 90% to 10% increased the distance between sister chromatids only modestly (from 1.04 ± 0.13 μm to 1.18 ± 0.19 μm). A similarly modest increase in the distance between sisters was also observed when the relative ratio of condensin I to II was forced to be 1:1 with a near-original level of cohesin (1.21 ± 0.16 μm). In striking contrast, when the cohesin level was reduced to 10% in an extract with the 1:1 ratio of condensin I to II, the distance between chromatid arms (but not between sister centromeres) was drastically increased (1.62 ± 0.25 μm), resulting in the formation of highly characteristic, X-shaped chromosomes. Admittedly speculative, we interpret these observations as follows. Parallel juxtaposition of sister chromatid arms within a metaphase chromosome is sustained by two mechanistically distinct factors: One is cohesin-mediated cohesion between sister chromatid arms, and the other is condensin I-dependent structural rigidity of each chromatid arm (e.g., Almagro et al. 2004). Our current results clearly demonstrate that reducing either one of the two factors to 10%–20% affects the juxtaposition of sister arms only modestly, whereas reducing both of them has a synergistic effect, leading to hyperdissociation of sister chromatid arms. It should be noted that similar X-shaped chromosomes are often observed in somatic cells arrested at mitosis for a long time, during which cohesin is lost progressively from chromosome arms [Gimenez-Abian et al. 2004; Nakajima et al. 2007]. Although previous studies in budding yeast implicated possible contribution of cohesin to “condensation” [Guacci et al. 1997; Lavoie et al. 2002] and of condensin to “cohesion” [Lam et al. 2006], the inability to visualize individual chromosomes in yeast made it difficult to provide lucid explanations for the functional cross-talk between the two processes. Our current study helps clarify this tangled issue, demonstrating that cohesin and condensin I collaborate to shape and maintain metaphase chromosomes, yet through distinct mechanisms.
We next investigated whether more drastic manipulation of cohesin levels left on metaphase chromosomes might affect the behavior of condensins and, if so, how. To this end, we prepared duplicated chromosomes with no cohesin in a cohesin-depleted extract and those with an increased level of cohesin in a Wapl-depleted extract (Fig. 3B; Losada et al. 1998; Shintomi and Hirano 2009). It was noticed that the axial distribution of condensin II became more prominent in cohesin-free chromosomes (Δcohesin) compared with control ones (Δmock). Conversely, condensin II was hardly detectable in poorly resolved chromosomes with an unusually high level of cohesin (ΔWapl) [Fig. 3C]. On the other hand, the signal intensity of condensin I was indistinguishable among the three conditions examined, confirming our previous data that condensin I loading occurs independently of cohesin release in this cell-free system (Losada et al. 1998, 2002). The current results show instead that accumulation of condensin II on chromosome axes is counterbalanced by the amount of cohesin left on metaphase chromosomes (Supplemental Fig. S4B). Notably, cohesin is released from chromatin exactly when condensin II initiates chromosome condensation within the prophase nucleus in somatic cells [Waizenegger et al. 2000; Hirota et al. 2004; Ono et al. 2004; Gerlich et al. 2006a,b]. Thus, the balancing actions between cohesin and condensin II observed here would represent a “tug-of-war” between forces holding and resolving sister chromatids.

Additional evidence that balancing actions of condensin I and II are important for chromosome assembly

To gain additional insight into functional interactions between condensin I and II, we set up a different type of assay that induces chromosome assembly from an unduplicated substrate [the single-chromatid assembly assay]. In this experimental setup, sperm chromatin was directly incubated with metaphase-arrested egg extracts to produce single chromatids, independently of DNA replication or cohesin. This simple assay for chromosome assembly, which magnifies its individualization process (Swedlow and Hirano 2003), had been used effectively for the original identification of condensin I [Hirano and Mitchison 1994; Hirota et al. 1997]. In this assay, both condensin I and II started to associate with chromatin in a very short time window upon incubation [i.e., within 5 min], as judged by immunoblotting analysis [Supplemental Fig. S5A]. Consistently, immunofluorescence analysis showed that condensin I and II localized to the axis of each chromatid with distinct patterns at both early and late time points. After an 80-min incubation, condensed chromatids became individualized so that they were readily distinguishable from each other [Supplemental Fig. S5B].

During the course of this study, we noticed unexpectedly that a far larger amount of condensin II was loaded on the single chromatids than the duplicated chromosomes, whereas the signals of condensin I on the two types of chromosomes were comparable [Fig. 4A]. Although the absence of cohesion alone would not fully explain the apparent hyperloading of condensin II observed in the single-chromatid assembly assay [and the reason is currently unknown], we realized that this property might give us an opportunity to address balancing actions of condensin I and II in a different context. With this background information in our hand, the same ratio manipulation was applied to the single-chromatid assembly assay. To our surprise, we found that no discrete chromatid structures were assembled in the 1:1 extract; instead, masses of entangled chromatin fibers were observed, although both condensins bound to them [Fig. 4B, row c]. This characteristic morphology was distinct from that produced in extract depleted of either condensin I, II, or both [Fig. 4B, row b, Supplemental Fig. S5C; Ono et al. 2003]. Interestingly, however, when condensin II was further

Figure 3. Cohesin collaborates with condensins to shape chromosomes. (A) Chromosomes were assembled in extracts containing various levels of cohesin and condensin I [for details, see Supplemental Fig. S4A], and subjected to immunofluorescence analysis with anti-topoisomerase II (topo II). Blowup images of selected regions [indicated by the rectangles] are shown in the bottom row. Bar, 5 μm. The average distance between sister chromatids, together with standard deviation, is provided below each image [n = 22, 28, 21, 20, from left to right]. (B) Chromosomes were assembled in a control extract [Δmock], an extract depleted of cohesin [Δcohesin], or Wapl [ΔWapl]. To visualize cohesin left on these chromosomes, immunofluorescence was performed with an antibody against its SA1 subunit. Bar, 5 μm. (C) Chromosomes were assembled as described in B, and labeled with anti-CAP-G and anti-CAP-H2 antibodies. Bar, 5 μm.
between sister chromatids (Losada and Hirano 2001; Hirano and Mitchison 1994; Hirano et al. 1997; Ono et al. 2003) and cohesin (Guacci et al. 1997; Michaelis et al. 1997; Losada and Hirano 2005). Given the fixed volume of a chromatin fiber whose density within a chromosome has been shown to be constant from prometaphase through metaphase [Mora-Bermudez et al. 2007], axial shortening would create shorter and thicker chromatids, whereas lateral compaction would make chromatids longer and thinner. In fact, it has been demonstrated that chromosomes progressively get shorter and thicker during animal development [Belmont et al. 1987; Michel et al. 1993]. On the basis of the ratio manipulation experiments reported here, we propose that the primary actions of condensin I and II are aimed at lateral compaction and axial shortening, respectively, and that their intricate balance acts as one of the critical determinants in shaping chromatids [Fig. 4C]. While this is clearly an oversimplified model, it is consistent with the recent finding that caspase-mediated cleavage of a condensin I subunit results in the formation of short and thick chromosomes in cells arrested at metaphase for a long time [Lai et al. 2011]. On the other hand, our results suggest that cohesin contributes to chromosome shaping in two ways. First, release of cohesin itself from chromosome arms facilitates condensin II-mediated axis formation. Second, residual cohesin holds sister chromatid arms together, whose juxtaposition could further be reinforced by condensin I-mediated chromatid rigidity. In the future, it will be of great interest to combine the sophisticated cell-free protocols reported here with other assays such as micromanipulation of chromosomes [Marko et al. 2008]. Furthermore, it will be important to examine exactly how expression of condensins and cohesin might be under the control of developmental cues, and how their intricate balance might be established and fine-tuned among different organisms. Such considerations would undoubtedly help broaden our horizon, linking chromosome biology to other neighboring areas such as developmental biology and evolutionary biology.

Materials and methods

**Duplicated chromosome and single-chromatid assembly assays**

Interphase low-speed supernatants and cytostatic factor-arrested high-speed supernatants of *Xenopus* eggs were prepared as described previously [Hirano et al. 1997; Shintomi and Hirano 2009]. Throughout this study, we refer to them as interphase egg extracts and metaphase-arrested egg extracts, respectively. To prepare duplicated chromosomes, sperm chromatin was mixed with interphase egg extracts at a final concentration of 1000 per microliter. After a 100-min incubation at 22°C, the mixture was supplemented with a nondegradable cyclin B (60 nM) to induce mitosis, and incubated for another 80 min. To assemble single chromatids, sperm chromatin was directly incubated with metaphase-arrested egg extracts. Immunodepletion was performed as described previously [Shintomi and Hirano 2009]. Detailed protocols for depletion and manipulation are described in the Supplemental Material.

**Immunofluorescence**

Immunofluorescence was performed as described previously [Shintomi and Hirano 2009] with the following modifications. For double labeling with rabbit antibodies against condensin I- and II-specific subunits, chromosomes recovered on coverslips were first incubated with 2 μg/mL unlabeled anti-CAP-H2, followed by Alexa Fluor 594-labeled anti-rabbit H2.
IgG secondary antibody [Invitrogen]. The coverslips were soaked in TBS containing 1 mg/mL rabbit IgG to saturate the open IgG-binding site, and then incubated with anti-CAP-G (1 µg/mL) that had been covalently labeled with Alexa Fluor 488 (by using a kit from Invitrogen). After counterstaining with DAPI, the coverslips were mounted on slides with VectaShield mounting medium (Vector Laboratories) and examined with an epifluorescence microscope (BX51, Olympus).

Quantification of chromosome shapes

For quantification of chromosome lengths, chromosomes were randomly selected and the lengths of lines manually traced along CAP-G-positive structures were quantified with the measure function of the ImageJ software [http://rsb.info.nih.gov/ij]. An unpaired two-tailed t-test was used for statistical analysis. For quantification of chromosome arm distances, lines were drawn perpendicularly to the path of sister chromatids at a couple of points on each chromosome (with avoiding centromeric regions), and the distance between top II-peak positions on sister chromatids was measured.

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