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A super-resolution map of the vertebrate kinetochore

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A longstanding question in centromere biology has been the organization of CENP-A–containing chromatin and its implications for kinetochore assembly. Here, we have combined genetic manipulations with deconvolution and super-resolution fluorescence microscopy for a detailed structural analysis of chicken kinetochores. Using fluorescence microscopy with subdiffraction spatial resolution and single molecule sensitivity to map protein localization in kinetochore chromatin unfolded by exposure to a low salt buffer, we observed robust amounts of H3K9me3, but only low levels of H3K4me2, between CENP-A subdomains in unfolded interphase prekinetochores. Constitutive centromere–associated network proteins CENP-C and CENP-H localize within CENP-A–rich subdomains (presumably on H3-containing nucleosomes) whereas CENP-T localizes in interspersed H3-rich blocks. Although interphase prekinetochores are relatively more resistant to unfolding than surrounding pericentromeric heterochromatin, mitotic kinetochores are significantly more stable, reflecting mitotic kinetochore maturation. Loss of CENP-H, CENP-N, or CENP-W had little or no effect on the unfolding of mitotic kinetochores. However, loss of CENP-C caused mitotic kinetochores to unfold to the same extent as their interphase counterparts. Based on our results we propose a new model for inner centromeric chromatin architecture in which chromatin is folded as a layered boustrophedon, with planar sinusoids containing interspersed CENP-A–rich and H3-rich subdomains oriented toward the outer kinetochore. In mitosis, a CENP-C–dependent mechanism crosslinks CENP-A blocks of different layers together, conferring extra stability to the kinetochore.


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Centromeric Region Unfolds to Higher Extents in Interphase than in Mitotic Cells. We previously showed that exposure of chromosomes to low ionic strength TEEN buffer [a low-salt buffer in which chromatin higher-order structures are destabilized (20)] causes chromatin to unfold to beads-on-a-string nucleosomes while retaining kinetochore protein binding (15, 20). Although interphase prekinetochores are much more resistant to unfolding in this buffer than pericentromeric heterochromatin (15), they did unravel

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during longer incubations in TEEN buffer, producing fibers in which patches of CENP-A and H3-containing chromatin alternated (Fig. 2B). The intercalated H3-containing chromatin was rich in H3K9me3 (mean occupancy, 56 ± 20% of the CENP-A-containing region; n = 11). In other experiments, H3K4me2 appeared to be present at lower levels in the intercalated chromatin (mean occupancy, 18 ± 12%; n = 9; Fig. 2D). A similar distribution of histone modifications has been observed in centromeres in maize and rice (10, 12).

Mitotic kinetochore chromatin is much more resistant to unfolding in TEEN buffer than its interphase counterpart, presumably because of structural maturation as cells enter mitosis (23, 24). The mean length (±SD) of unfolded kinetochores (determined by the borders of the GFP-CENP-A domains) derived from mitotic centromeres was 1.1 ± 0.44 μm (n = 98), compared with 2.47 ± 1.31 μm (n = 164) for interphase prekinetochores (Fig. 2C and E). Considering that the kinetochore diameter determined by serial-sectioning electron microscopy is 145 ± 27 nm (n = 48; Fig. S1) in SMC2ON DT40 cells at metaphase, the average mitotic kinetochore fiber undergoes an eightfold extension in TEEN buffer. This contrasts with an average 17-fold extension of interphase prekinetochore fibers.

In mitosis, 95% of the extended CENP-A domains measured less than 3 μm (Fig. 2E). Similarly, most (60%) unfolded interphase prekinetochores yielded single CENP-A blocks less than 3 μm long. We refer to these interphase prekinetochores and normal mitotic kinetochores as having undergone “stage 1” unfolding. The remaining 40% of unfolded interphase kinetochores displayed multiple interspersed CENP-A subdomains stretching over more than 3 μm, up to a maximum of 13.4 μm. We refer to this as “stage 2” unfolding.

CENP-C Is Required for the Increased Structural Integrity of Mitotic Kinetochore. In a genetic approach to identify proteins specifically required for the increased stability of mitotic kinetochores, we performed our kinetochore unfolding assay using mitotic cells depleted of CENP-C (18), CENP-H (16), CENP-N (17), and CENP-W (19). Remarkably, no significant difference was observed between unfolded WT kinetochores and those prepared from cells lacking CENP-H, CENP-N, or CENP-W (Fig. 3A and B). Thus, we observed stage 2 CENP-A domain unfolding in 4%, 3%, and 8% of unfolded CENP-HOFF, CENP-NOFF, and CENP-WOFF kinetochores. This was not significantly different from the values obtained with CENP-HON, CENP-NON, and CENP-WON kinetochores in this assay, although following loss of CENP-W a few more unfolded prekinetochores were seen.

In contrast, the loss of CENP-C caused a significant destabilization of the mitotic kinetochore chromatin in the TEEN assay, as the percentage of kinetochores undergoing stage 2 unfolding increased from 5% to 25% (Fig. 3C and D). As with interphase prekinetochores, many CENP-A domains underwent only stage 1 unfolding following CENP-C depletion. However, the maximal lengths of stage 2-unfolded CENP-A domains achieved after CENP-C depletion approached those observed for interphase kinetochore fibers (Fig. 2E).

Depletion of condensin yielded intermediate results in this unfolding assay. The number of extended CENP-A domains larger than 3 μm increased from 4% to 14%, but domains larger than 6 μm were never observed (Fig. 3B). This suggests that condensin in the underlying heterochromatin may contribute in part to the structural integrity of kinetochore chromatin during mitosis, but that loss of condensin does not lead to full stage 2 unfolding.

Thus, of the four CCAN proteins tested, only CENP-C is required for the enhanced structural integrity of mitotic kinetochore chromatin.

Mapping Unfolded Interphase Prekinetochores Using Super-Resolution Microscopy. Fluorescence microscopy with subdiffraction limit spatial resolution yielded further insights into the organization of
kinetochore chromatin. Our method resembles (fluorescence) photostimulation-localization microscopy and stochastic optical reconstruction microscopy [reviewed by Lippincott-Schwartz and Manley (25)], which are based on single-molecule detection of switchable fluorophores and provide single-molecule sensitivity with a spatial resolution of tens of nanometers. Cycles of stochastic switching, detection, and localization of single molecules on a wide-field microscope are used to reconstruct super-resolution images (Materials and Methods). A related technique was recently used to localize chromosomal proteins in fixed cells with a precision of 20 to 30 nm (26).

We tagged CENP-A with the reversible fluorescent protein Dronpa, which photo-switches between dark and bright states after irradiation at 488 and 405 nm, respectively (27, 28). Chromatin from cells stably expressing Dronpa-CENP-A was unfolded as described earlier and immunostained for detection of different histone modifications using Alexa Fluor 647-labeled secondary antibodies. In the presence of a “switching buffer” containing an oxygen scavenger and reducing agent (Materials and Methods), Alexa Fluor 647 cycles reversibly between fluorescent and dark metastable states (29, 30). By sequential imaging with 488-nm and 633-nm irradiation, a two-color super-resolution map revealing the distribution of CENP-A relative to other kinetochore components could be constructed.

Fig. 4 A shows an extended centromeric region stretching over 13 μm in which alternating CENP-A and H3K9me3 domains are readily apparent. Super-resolution microscopy also detected H3K4me2 between CENP-A domains, but whereas H3K9me3 was present as discrete blocks, H3K4me2 showed a more scattered distribution (Fig. 4B). Chromatin regions unlabeled with anti-H3K9me3 or anti-H3K4me2 may contain nucleosomes with other (or no) modifications.

A rough estimate of the amount of CENP-A in the fiber in Fig. 4 A can be obtained from the number of localizations (a total of 123). Assuming that, on average, each Dronpa molecule switches approximately three to five times (31), the number of labeled CENP-A molecules is approximately 25 to 40. This value should be taken with caution, as Dronpa can switch as many as 170 times (28). For this estimation, we also assume that most CENP-A in the fibers is labeled with Dronpa and that the amount of endogenous CENP-A is not significant (Fig. S2A).

By reconstructing super-resolution images of the density of single-molecule localizations (Materials and Methods), the width could be estimated for CENP-A- and H3K9me3-labeled fibers (Fig. 4 C and ). Both exhibited a bimodal distribution centered at approximately 40 nm and 60 nm. Taking into account that the values are convolved with a 37-nm error (Fig. S2 B–D), these measurements likely correspond to two populations of chromatin fibers that differ in diameter by 20 nm. For chromatin fibers in this size range, the most likely candidates are the canonical 10- and 30-nm fibers. This important result suggests the possibility to distinguishing between different levels of chromatin organization by super-resolution fluorescence microscopy.

**Localization of CCAN Proteins in Stretched CENP-A–Containing Chromatin Using Super-Resolution Microscopy.** To determine whether components of the CCAN are associated with CENP-A or H3 blocks in kinetochore chromatin, we localized CENP-C, CENP-H, and CENP-T by super-resolution microscopy of unfolded kinetochores from cells expressing Dronpa-CENP-A (Fig. 5). All three CCAN components colocalized with CENP-A in shorter compact fibers (Fig. 5 A–C). However, when longer fibers were unfolded to reveal multiple CENP-A subdomains CENP-C and CENP-H remained closely associated with the CENP-A–rich subdomains, but CENP-T clearly localized between those subdomains (Fig. 5 A–C). Thus, the distribution of CENP-T resembles that of H3K9me3 and H3K4me2 in unfolded prekinetochores, and suggests the presence of blocks of canonical histone H3 in the outer surface of kinetochore chromatin.

**Discussion.**

We have combined gene targeting with biochemical manipulation of chromatin higher-order structure and super-resolution microscopy to characterize a vertebrate kinetochore. Kinetochores persist during interphase as locally condensed chromatin domains known as prekinetochores (32) that undergo a program of structural (23) and biochemical (24) maturation as cells enter mitosis. By using a protocol in which kinetochore chromatin is unfolded in vitro with TEEN buffer (20), we show that this maturation renders mitotic kinetochore chromatin considerably more robust than that of interphase prekinetochores. This presumably helps to give kinetochores the structural rigidity required to withstand pulling forces within the mitotic spindle.

Genetic analysis using DT40 conditional knockouts reveals that this mitotic stabilization of kinetochore chromatin requires CENP-C but not CENP-H, CENP-N, or CENP-W. This was surprising, as CENP-H is required for CENP-C accumulation at interphase prekinetochores (16, 18). However, a role for CENP-C...
in stabilization of the mitotic kinetochore is consistent with previous observations that CENP-C determines the size and continuity of the kinetochore plate (24, 33). CENP-C could perform a scaffolding role by interacting directly with DNA (19, 34) or RNA (35, 36) or with proteins such as CENP-L or Pcs1 as shown in Schizosaccharomyces pombe (37).

Interphase prekinetochore chromatin unfolded with TEEN buffer consists of extended fibers in which multiple CENP-A subdomains alternate with subdomains containing H3K9me3. This heterochromatin-associated modification was also seen to abut CENP-A domains in stretched pericentromeric chromatin trailing behind kinetochores undergoing poleward excursions in condensed-depleted cells. Furthermore, the pericentromeric chromatin also consistently displayed a gap between H3K4me2 staining and the kinetochore. However, further analysis of the histone modification pattern of unraveled centromeric fibers, using both deconvolution and super-resolution imaging, suggested that, as in rice centromeres (10), both histone H3 modifications (detected in independent experiments) are present in the centromeric region. Comparison of the two sets of experiments suggested that H3K4me2 may be present at lower levels than H3K9me3 in this region.

Although it is now accepted that CENP-A domains alternate with canonical H3 blocks within kinetochore chromatin (6, 7), the modification pattern of the canonical histones blocks seems to be less conserved. In human and Drosophila interphase prekinetochores, H3K4me2, but not H3K9me3, was found to be intercalated between CENP-A subdomains (8). H3K4me2 and lower levels of H3K9me3 were readily detected within the kinetochore of a human artificial chromosome by ChIP (38). In contrast, levels of H3K4me2 were much lower in one human neocentromere (39) and in maize centromeres (11, 12). Clearly, more work is required to understand the role of particular histone modifications in kinetochore chromatin structure and function.

Here we examined the distribution of members of the CCAN relative to CENP-A along the chromatin fiber. CENP-C, CENP-H, CENP-I, CENP-K-U, and CENP-W all coimmunoprecipitate with CENP-A following partial digestion of chromatin by micrococcal nuclease (40, 41), and ChIP studies showed colocalization of CENP-C and CENP-H with discontinuous domains of CENP-A in human neocentromeres (39, 42, 43). CENP-N is the only CCAN protein that has been shown to interact directly with CENP-A nucleosomes (44). CENP-C, CENP-T, and CENP-W coimmunoprecipitate with H3 nucleosomes after extensive nuclease digestion (19, 45). This suggests that some canonical H3 nucleosomes must be close to CENP-A–rich subdomains in the inner kinetochore. We confirmed this by super-resolution light microscopy, finding that CENP-C and CENP-H colocalized with CENP-A–rich subdomains in unfolded kinetochores. This suggests the presence of some canonical H3 nucleosomes within the CENP-A–rich subdomains. In contrast, CENP-T, H3K4me2, and H3K9me3 were interspersed between the CENP-A subdomains.

Detailed maps of kinetochore proteins in fixed chromosomes derived from measurements using two-color fluorescence light microscopy place CENP-C external, but very close to, CENP-A, with CENP-T slightly external to CENP-C (46, 47). Current kinetochore chromatin folding models based on data available from localization of canonical histones and CENP-A on unfolded chromatin fibers propose that CENP-A and H3 coexist in the same fiber and are sorted on different faces of an “amphipathic” superhelix, in which CENP-A faces the outer kinetochore and the H3-containing blocks are embedded in the centromere (7, 13). The data here presented extend this mapping of

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**Fig. 4.** Characterization of unfolded prekinetochores using fluorescence microscopy with subdiffraction spatial resolution. (A) Example of a 13.4-μm interphase fiber in which H3K9me3 blocks are clearly observed between CENP-A arrays. (B) Example of a 15.1-μm fiber in which H3K4me2 is also detected between CENP-A arrays, but with a more diffuse distribution. Dronpa-CENP-A is represented as green, H3K9me3 and H3K4me2 labeled with Alexa647 are represented as red. Yellow represents colocalization. Each dot corresponds to the localization of a single molecule switching event. (C) Frequency distribution histogram of stretched fiber widths measured from super-resolution reconstructed images of (C) Dronpa-labeled CENP-A and (C) Alexa Fluor 647–labeled H3K9me3. The solid lines represent the best fit to bimodal Gaussian distributions centered at 46 and 67 nm for CENP-A (n = 53) and 40 and 57 nm for H3K9me3 (n = 30).

**Fig. 5.** Localization of CCAN components in interphase CENP-A chromatin fibers. CENP-H (A) and CENP-C (B) in red colocalize with CENP-A (green) both in short (A, B) and more extended (A’, B’) fibers. (C) CENP-T (red) colocalizes with CENP-A (green) in short fibers (C); however, in more extended fibers (C), CENP-T is interspersed between CENP-A domains.
CENP-A relative to the CCAN proteins (most of which were still unidentified at the time the existent models were proposed) and require changes to the original solenoid model to take into account the fact that H3 domains with associated CENP-T are present on the outer face of the kinetochore chromatin.

Our data could be explained by a modification of the “amphipathic” superhelix model if helical segments were oriented radially with their long axes perpendicular to the chromosome axis. Such an orientation would expose some CENP-A and H3 on the outer surface of the chromosome, but is difficult to reconcile with immunoelectron microscopy and super-resolution colocalization of other kinetochore proteins with CENP-A by fluorescence microscopy, all of which indicate that CENP-A seems to provide a basal layer to the kinetochore that does not penetrate significantly into the chromosome interior (13, 46, 47).

Based on our data, we suggest a simple alternative model for the topology of chromatin fiber folding in regional kinetochores. We propose that alternating CENP-A and H3 domains fold into a planar sinuosoidal patch, or boustrophedon (Greek: “ox-turning”; Fig. 6). Such a topology would allow kinetochore size to vary according to the number of microtubules bound with minimal perturbation of local packing. Each kinetochore could be composed of several such patches stacked on top of one another as shown in Fig. 6A.

This organization can explain the two stages of kinetochore chromatin unfolding observed in the present study. Stage 1 unfolding might correspond to the “straightening out” of the boustrophedon folds, with the different layers remaining held together laterally by structural crosslinks dependent at least in part on CENP-C. Stage 2 unfolding would entail the loss of connections between layers and stretching out of the whole kinetochore into a single contiguous linear segment. As we have shown, interphase prekinetochores frequently undergo stage 2 unfolding in TEEN buffer, whereas this is seldom observed for mitotic kinetochores.

Given that CENP-C is required for this stability of the mitotic kinetochores, it is interesting to note that the genetic requirements for stable association of CENP-C with kinetochores differ between interphase and mitosis, with the former, but not the latter, requiring CENP-H (16, 18). This suggests that at least one aspect of kinetochore maturation may involve a change in the detailed mechanism of CENP-C association with kinetochore chromatin.

In surface view, our proposed model fits well with the recent proposed patterning of kinetochore proteins based on known protein associations in a “horizontal view” of the outer kinetochore (1). That model did not suggest a topological path for the chromatin fiber, which we propose here. In both models, CENP-A and H3 nucleosomes face the external surface, enabling the binding of all CCAN proteins. CENP-C could bind to the more internal CENP-A blocks, crosslinking several layers and explaining the similar oscillations undergone by CENP-A and CENP-C when kinetochores are under tension exerted by microtubules (47). The KMN network assembles in mitosis on top of the CCAN and binds microtubules. KMN binding may confer stability to the kinetochore by crosslinking the CENP-C chromatin either directly or indirectly.

Materials and Methods

DT40 conditional knockout cell lines were cultured as described before (22). Transfection of cells with a construct expressing GFP-CENP-A, antibody staining conditions, the quantification of DNA amounts in CENP-A and CENP-H kinetochore domains and detailed electron microscopy analysis of kinetochore sizes are described in SI Materials and Methods.

Super-Resolution Microscopy with Single Molecule Sensitivity. Cells expressing Dronpa-CENP-A were plated on clean 22 × 22 mm coverslips for 20 min. Dronpa cDNA was provided by J. Lippincott-Schwarz (Bethesda, MD). The TEEN assay and antibody staining were performed as described in SI Materials and Methods using Alexa Fluor 647 (Invitrogen)-coupled secondary antibodies at 1:200 dilution. Coverslips were then attached to a CoverWell imaging chamber (Grace Bio Labs), containing “switching buffer” that promotes photo-induced blinking of Alexa Fluor 647: 10 mM PBS solution (pH 7.4) with an oxygen scavenger (0.5 mg/mL glucose oxidase; Sigma), 40 μg/mL catalase (Sigma), 10% wt/vol glucose (Fisher Scientific), and 50 mM β-mercaptoethanol (Fluka) (29). Single-molecule fluorescence imaging was performed on a Nikon Eclipse TE2000 inverted microscope, equipped with a total internal reflection fluorescence oil-immersion objective (apochromat, magnification ×100; N.A., 1.49; Nikon). Wide-field illumination was achieved by focusing the expanded and collimated laser beam onto the back-focal plane of the objective. The resulting illuminated area was approximately 60 μm in diameter. Excitation was provided by a 488 nm CW Ar+ laser (163-C, 0.5 kW/cm2; Spectra-Physics) or 633 nm He/Ne CW laser (model 31–2140–000, 1 kW/cm2 at the sample; Coherent) passing through appropriate band-pass filters (Chroma Technology). Pulses for Dronpa photoactivation (2 Hz, 5 ms, 1 W/cm2) were provided by a CW 405-nm laser (Cube; Coherent) controlled by a function generator (USB-6218, National Instruments).

Emission fluorescence was collected by the same objective and imaged by an Andor Luca S electron-multiplying CCD camera after passing through a dichroic mirror (z488rdc or z633rdc; Chroma Technology), additional spectral filters (HQ500LP and HQ530/50, or HQ645LP and HQ700/75; Chroma Technology), and lenses resulting in a final pixel size of 74 nm. Integration time per frame was 100 ms. Typically 500 to 1,000 frames were collected. Two-color imaging of Dronpa and Alexa Fluor 647 was performed sequentially. Chromatic shifts were corrected by localizing immobilized 0.1 μm Tetraspeck beads (Invitrogen) with both colors.

Density super-resolution images like that in Fig. 5 were reconstructed by dividing each pixel into 16 subpixels and assigning each localization to a subpixel. The image brightness thus represents the density of localizations

Fig. 6. Model showing proposed boustrophedon arrangement of kinetochore chromatin. (A) A single continuous chromatin segment is arranged in a sinuosoidal wave in a series of layers linked at both ends to heterochromatin. (B) Top and side view as indicated in A. (C) CCAN protein distribution in the kinetochore. (D) The KMN network assembles in mitosis on top of the CCAN and may confer stability to the mitotic kinetochore by crosslinking the CENP-C either directly or indirectly (see text for details).
in a subpixel. The FWHM values were estimated from line cross-sections of density super-resolution images of fibers drawn with ImageJ by fitting a Gaussian function in GraphPad Prism. A spatial resolution of 37 nm was estimated by multiple localizations of the same single molecule of AlexaFluor647 analyzed under identical conditions, and fitting a Gaussian function to its cross section in a density reconstructed image (Fig. S2).

Movies were analyzed with Igor Pro by fitting Gaussian functions to individual molecules and localizing their centers.

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