Temperature-dependent regulation of rDNA condensation in Saccharomyces cerevisiae

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ABSTRACT
Chromatin condensation during mitosis produces detangled and discrete DNA entities required for high fidelity sister chromatid segregation during mitosis and positions DNA away from the cleavage furrow during cytokinesis. Regional condensation during G1 also establishes a nuclear architecture through which gene transcription is regulated but remains plastic so that cells can respond to changes in nutrient levels, temperature and signaling molecules. To date, however, the potential impact of this plasticity on mitotic chromosome condensation remains unknown. Here, we report results obtained from a new condensation assay that wildtype budding yeast cells exhibit dramatic changes in rDNA conformation in response to temperature. rDNA hypercondenses in wildtype cells maintained at 37°C, compared with cells maintained at 23°C. This hypercondensation machinery can be activated during preanaphase but readily inactivated upon exposure to lower temperatures. Extended mitotic arrest at 23°C does not result in hypercondensation, negating a kinetic-based argument in which condensation that typically proceeds slowly is accelerated when cells are placed at 37°C. Neither elevated recombination nor reduced transcription appear to promote this hypercondensation. This heretofore undetected temperature-dependent hypercondensation pathway impacts current views of chromatin structure based on conditional mutant gene analyses and significantly extends our understanding of physiologic changes in chromatin architecture in response to hypothermia.

Introduction
Chromatin condensation may have been the first dramatic cell cycle change observable to early microscopists and remains a topic both of fascination and clinical relevance. Chromatin condensation and the arrangement of chromosomes within the nuclear volume are collectively mediated by cohesin- and condensin-dependent DNA segment tetherings that occur in either cis (within a single DNA molecule) or trans (between separate DNA molecules) conformations. Recent evidence is consistent with the notion that cis DNA tetherings established during interphase are critical for the proper deployment of developmental transcription programs. Intriguingly, proper development likely also requires G1 trans tethers (here, between non-homologous chromosomes) that link together domains of either repressed or induced transcriptional activities termed TADs (Topologically Association Domains). The combination of cis and trans tethers during G1 that mediate regionalized chromatin condensation and nuclear architecture must remain plastic if cells are to respond appropriately to external cues. Numerous studies across several species document that the chromosome condensation state during interphase increases in response to heat-stress. For instance, hyperthermia induces premature chromosome condensation during S phase in CHO, HeLa and S3 cells and hypercondensation in Achlya ambisexualis hypha. Importantly, little is known regarding the mechanisms through which these induced and premature condensation reactions occur. Moreover, the extent to which cells respond to hyperthermia during mitosis, when chromosomes are already condensed, remains largely untested. Addressing these deficiencies becomes important since physiologic changes induced by heat stress are wide-ranging and include expression of heat shock proteins (HSPs), protein synthesis repression and altered rates of both transcription and mRNA turnover.

Budding yeast remains a mainstay model organism from which numerous aspects of chromatin condensation have come to light. Common to most studies is the testing of temperature-dependent gene product inactivation on rDNA structure. This is because rDNA undergoes cell cycle-specific structural changes: forming diffuse puff-like structures during G1 and condensing through multiple stages that include cluster and line formations before coalescing into discrete loop-like structures during mitosis. When either SMC (structural maintenance of chromosomes) cohesin or condensin complexes are impaired (for instance through the use of conditional alleles), mitotic cells contain puff-like rDNA loci instead of tightly condensed rDNA loops. Since the rDNA locus is a highly-specialized domain comprising iterative repeats that are repressed for Homologous
Recombination (HR) and under tight transcriptional regulation,29-31 parallel strategies were developed in yeast to assess changes in chromosome arm condensation.24,32 All of these studies operate under the assumption that mitotic chromatin structures in wildtype cells are refractory to the temperature shift required to inactivate conditional allele gene products. Conversely, a growing body of evidence suggests that DNA segment cis and trans tethers of interphase cells remain dynamic to respond to external cues.1-5,33 Here, we developed a streamlined rDNA condensation assay and report that wildtype budding yeast exhibit a mitotic chromatin hypercondensation activity that is induced at temperatures typically used to inactivate conditional alleles. Moreover, this hypercondensation activity predominantly targets the rDNA locus, revealing a heretofore unexplored and targeted physiologic response to hyperthermia.

Results

Temperature-dependent regulation of mitotic rDNA hypercondensation

Despite technical advances that include immunodetection, GFP tagging and Fluorescence in situ hybridization (FISH), barriers to detecting changes in chromatin architecture in yeast cells include small cell and nuclear sizes and limited longitudinal chromatin compaction relative to other eukaryotic cells.16,17,24-26,34,35 Here, we report on a streamlined procedure that provides for exquisite imaging of the yeast rDNA locus in the absence of GFP, stacked antibody complexes and hybridization of labeled nucleotide probes. To validate this procedure, we exploited a well-established observation that mutation of the cohesin subunit Mcd1/Scc1 results in condensation defects.17,19,24,26-38 Log phase wildtype and mcd1-1 mutant cells were synchronized in G1 (a factor) at 23°C and then released into 37°C (non-permissive temperature for mcd1-1) fresh medium supplemented with nocodazole to arrest cells preanaphase. Cell cycle progression was confirmed by detection of DNA content using flow cytometry (Figure S1A). The resulting preanaphase wildtype cells contained predominantly puff-like rDNA structures while preanaphase mcd1-1 mutant cells failed to condense their chromatin such that the rDNA formed predominantly puff-like structures (Fig. 1A). We quantified these differences and found that our procedure detects similar levels of condensation defects in mcd1-1 mutant cells as those obtained using FISH and GFP-based methodologies (Fig. 1B).17,24,26,27

During this investigation, we discovered a robust temperature-dependent effect that occurs in wildtype cells, independent of gene mutation. We modified our prior experimental strategy to further analyze this hypercondensation activity by releasing G1 arrested cells into either 23°C or 37°C medium supplemented with nocodazole (Fig. 1C). Cell cycle progression was confirmed by detection of DNA content using flow cytometry (Figure S1B). Our condensation assay revealed that preanaphase wildtype cells condense rDNA loops to form tight and discrete loops at both 23°C or 37°C. Surprisingly, however, rDNA loops were condensed but appeared significantly shorter in wildtype cells shifted to 37°C upon release from G1, compared with loops in wildtype cells maintained at 23°C (Fig. 1D). We thus quantified the loop lengths in wildtype cells at both temperatures by tracing the entire length of the loop structures. The results clearly document that rDNA loop lengths are significantly shorter (herein termed hypercondensed) in cells that progress from G1 into mitosis at 37°C (Fig. 1E).

Our condensation assay provides for detailed images of rDNA structure, but we wondered if other procedures could detect temperature-sensitive hypercondensation of rDNA. Net1 is an rDNA binding protein and represents one current standard by which rDNA structure is assessed.24,26-28 Wildtype cells expressing GFP-tagged Net1 were synchronized in G1, released into either 23°C or 37°C rich medium supplemented with nocodazole, and the resulting preanaphase cells assessed for rDNA loop lengths. Cell cycle progression was confirmed by detection of DNA content using flow cytometry (Figure S1C). Wildtype cells that progressed from late G1 into mitosis at 23°C exhibited significant longer rDNA loop lengths than cells released from G1 into 37°C medium (Fig. 1F, G). In combination, these results reveal that rDNA condensation is exquisitely sensitive to temperature changes in wildtype cells with elevated temperatures leading to robust hypercondensation that is easily measurable using well-established procedures.

Temperature-induced hypercondensation is not based on increased condensation rates

What is the basis for this heretofore unreported temperature-dependent effect on rDNA condensation? We speculated that the decreased loop size that occurs in wildtype cells maintained at 37°C might reflect an accelerated rate of condensation, relative to that which occurs at 23°C. If this model is correct, then longer incubations at 23°C should ultimately lead to hypercondensed rDNA. To test this prediction, cultures released from G1 were maintained in fresh medium supplemented with nocodazole for either 2.5 or 5 hours at 23°C and compared with cultures instead maintained for 2.5 or 5 hours at 37°C (Fig. 2A). Cell cycle progression was confirmed by detection of DNA content using flow cytometry (Figure S1D). The results reveal that rDNA loop lengths in cells maintained in preanaphase at 23°C for 5 hours failed to approach the loop lengths of cells maintained in preanaphase at 37°C for only 2.5 hours. Moreover, rDNA loop lengths in cells maintained in preanaphase at 23°C for 2.5 hours are identical to rDNA loop lengths in cells maintained in preanaphase at 23°C for 5 hours (Fig. 2B). The same is true for loop lengths in cells maintained at 37°C, regardless of incubation time. In all cases, rDNA loop lengths were significantly shorter in cells maintained at 37°C compared with cells maintained at 23°C (Fig. 2C). Thus, changes in rDNA architecture occur independent of condensation rates.

The rDNA hypercondensation machinery is active during mitosis

In most eukaryotes, chromosome condensation is a multi-step process that requires factors (such as core histones and...
cohesins) that are deposited during S phase and additional factors (such as linker histones and condensins) that associate or become activated later during the cell cycle.\textsuperscript{39,40} It thus became important to map the timing of this rDNA hypercondensation during the cell cycle. To test whether rDNA hypercondensation can be induced by elevated temperatures during M phase (after normal levels of condensation are already established), wildtype cells were synchronized in G1 at 23°C for 2.5 hours, then...
released at 23°C into fresh medium supplemented with nocodazole for 2.5 hours. Half of the resulting preanaphase cells were then shifted to 37°C for an additional hour and the other half maintained at 23°C for the same time period (Fig. 3A). Cell cycle progression was confirmed by changes in DNA content using flow cytometry (Figure S1E). As expected, rDNA in cells maintained at 23°C throughout the preanaphase arrest appeared as long loops that extend away from the DNA mass. In contrast, rDNA in cells arrested in mitosis at 23°C but then shifted to 37°C while maintaining the preanaphase arrest exhibited significantly hypercondensed rDNA loop lengths (Fig. 3B, C). Thus, cells exhibit an rDNA-directed hypercondensation activity during preanaphase that induces condensation beyond that which promotes entry into mitosis.

**Temperature-induced hypercondensation is reversible and occurs independent of changes in rDNA repeat number**

Budding yeast rDNA comprises approximately 150 repeats and is a hotspot for homologous recombination. This raised the possibility that the apparent hypercondensation is in reality a reduction in rDNA repeats through permanent recombination-based excisions and loss from the chromosome. We thus tested whether we could revert a hypercondensed rDNA loop back to an extended loop within a single preanaphase arrest. Wildtype cells were synchronized in G1 at 23°C, then released at 23°C into fresh medium supplemented with nocodazole. The resulting preanaphase arrested cells were then shifted to 37°C for 1 hour (which produces rDNA hypercondensation) before shifting back down to 23°C for one hour (Fig. 4A). Cell cycle progression was confirmed by detection of DNA content using flow cytometry (Figure S1F). Similar to our earlier results, cells shifted up to 37°C during preanaphase contain hypercondensed rDNA (Fig. 4B). Importantly, when these cells were shifted down to 23°C for one hour, the rDNA recovered to a less condensed state – nearly matching the extended loop lengths of cells maintained at 23°C throughout the time course (Fig. 4B, C). Thus, rDNA hypercondensation is reversible and occurs independent of changes in the number of rDNA repeats – a revelation supported by
the absence of Net1-GFP decorated excised rDNA circles under conditions that similarly produce short rDNA loops (Fig. 3C).

**Temperature-induced hypercondensation occurs independent of rDNA transcriptional inhibition**

rDNA transcription, ribosome biogenesis and translational outputs are all upregulated during periods of accelerated growth. This increased rDNA transcription requires a relaxed DNA conformation and RNA polymerase accessibility.\textsuperscript{23,29,30,42-44} In contrast, yeast cell rDNA transcription levels decrease to basal line during mitosis that is coordinated with rDNA condensation.\textsuperscript{45,46} One prediction from these findings is that wildtype cells shifted to 37°C and that contain hypercondensed rDNA would exhibit slower growth kinetics (reduced rDNA transcription) compared with cells incubated at 23°C. To test this prediction, both S288C and W303 wildtype strains were sub-cultured over a 2 day period to ensure log phase growth before monitoring cell growth by spectroscopy. The results show that both yeast strains exhibit significantly increased growth rates at 37°C compared with 23°C (Fig. 5A, B). These findings negate the model that wildtype yeast cells exhibit hypercondensed rDNA due to slower growth at 37°C and instead are consistent with previous finding that wildtype yeast
significantly increase ribosomes synthesis when shifted to 36°C from 23°C growth conditions. The mechanism through which yeast cells exhibit increased growth rates that require elevated transcription from the rDNA locus, despite the hypercondensation that occurs at 37°C (Figs. 1–4), remains unknown.

**Temperature-induced hypercondensation is rDNA specific**

rDNA is unique in terms of architecture, binding factors, transcription regulation, and altered levels of recombination, compared with the remainder of the genome. This raises the possibility that the temperature-dependent hypercondensation of rDNA is locus specific. To test this possibility, we obtained a chromosome arm condensation assay strain, kindly provided by Dr. Frank Uhlmann, that contains 2 lacO cassettes spaced 137 kb from each other on chromosome XII. Each LacO cassette is detected by lacR-GFP such that the inter-GFP distance allows for quantification of chromosome arm condensation. To validate this procedure, we placed through mating and dissection the LacO/lacR-GFP condensation cassette into mcd1– strains previously demonstrated as exhibiting condensation defects at both rDNA and chromosome arms. Log phase wildtype and mcd1– cells were synchronized in G1 and released into either 23°C or 37°C fresh medium supplemented with nocodazole to arrest cells preanaphase (Figure S2A). The resulting pre-anaphase synchronized cells were then fixed in paraformaldehyde and the disposition of arm condensation quantified by measuring the distance between GFP loci (Figure S2B). The results reveal a significant increase in the inter-GFP distance in mcd1– cells compared with wildtype cells (Figure S2C,D), confirming the efficacy of this arm condensation assay.

To test whether temperature-dependent hypercondensation extends to chromosome arm loci beyond that of rDNA, wildtype cells that harbor the arm condensation assay cassette were synchronized in G1 at 23°C and released into 23°C fresh medium supplemented with nocodazole for 2.5 hours. Half of the resulting preanaphase cells were shifted to 37°C for 1 hour while the other half was maintained at 23°C for 1 hour (Fig. 6A). Cell cycle progression and synchronization was confirmed by DNA content using flow cytometry (Figure S1G). The resulting preanaphase synchronized cells were then fixed and the distance between GFP loci quantified (Fig. 6B). The results reveal that preanaphase cells maintained at either 23°C or 37°C exhibit nearly identical inter-GFP distances (Fig. 6C, D). Thus, chromatin along the arm appears insensitive to temperature-induced hypercondensation that predominantly targets the rDNA locus.

**Figure 5.** Temperature-induced hypercondensation occurs independent of rDNA transcription. (A) Growth curve of wildtype S288C background strain (YBS1019). (B) Growth curve of wildtype W303 background strain (YBS2020).

**Figure 6.** Temperature-induced hypercondensation is rDNA specific. (A) Schematic of synchronization and experimental procedure performed on wildtype cells (YBS2078). (B) Micrographs of DNA masses (DAPI), chromosome arm loci (GFP) and cell morphology (DIC) in preanaphase arrested cells at 23°C and 37°C. (C) Quantification of the distribution of measured distances between 2 arm loci GFP dots (N = 111 cells at 23°C and 106 cells at 37°C). (D) Quantification and statistical analyses of the GFP arm inter-loci distances in each experimental group (N = 111 cells at 23°C; N = 106 cells at 37°C; P-value = 0.812).
Discussion

Chromosome condensation ensures appropriate chromosome segregation but also influences transcriptional programs and nuclear architecture. Thus, it is not surprising that mutation of condensation pathways results in mitotic failure, aneuploidy and is further linked to developmental maladies such as microcephaly. A major revelation of the current study is that yeast cells contain hypercondensation machinery that is induced specifically during mitosis in response to elevated temperatures. This newly observed phenomenon is due neither to accelerated condensation reactions (hypercondensation at 37°C cannot be balanced by longer incubation time at 23°C), nor rDNA repeat reduction through recombination (hypercondensed rDNA loop lengths re-extend quickly upon shifting to 23°C and micrographs thus fail to detect extrachromosomal rDNA signals in Net1-GFP cells that contain hypercondensed loops). Note that the homologous recombination machinery in general is suppressed during M phase such that changes in rDNA repeat numbers typically are monitored over several generations. In contrast, we observe inducible and reversible rDNA hypercondensation within a single M-phase arrest. Intriguingly, yeast hypercondense specific DNA loci in response to nutrient starvation, although the extent that these represent similar mechanisms in hypercondensation remains unknown.

What are the mechanisms through which rDNA hypercondensation at elevated temperature occurs? Early analyses of condensin mutants revealed reversibility in normal levels of chromosome condensation during mitosis: transient inactivation of temperature sensitive Brn1-9 condensin subunit during M phase resulted in rDNA decondensation that recondensed upon a shift back to permissive temperature. This reversibility of normal condensation levels during mitosis depends on both condensin and cohesin, although cohesin mutants thus far do not exhibit reversibility condensation - suggesting that cohesin promotes condensin activation. These observations raise the possibility that the reversible rDNA hypercondensation documented here requires similar molecular mechanisms that include cohesin and condensin - which are critical for mitotic chromosome condensation, stabilization of rDNA architecture and transcription regulation. Condensins in addition introduce positive supercoils into chromatin, promotes complementary single strand DNA reannealing and facilitates DNA catenation resolution. Cohesin-dependent tethering of DNA segments is also critical for transcription regulation, nuclear architecture and sister chromatid tethering. A final mechanism of hypercondensation could involve heat shock proteins, which may act in coordination or in parallel to the recruitment of condensins and cohesins (Fig. 7). The extent to which elevated temperature alters condensin, cohesin or heat shock protein deposition/activation onto rDNA remains an important goal of future research.

A second revelation of the current study is the apparent specificity of this hypercondensation activity. Using both our streamlined condensation assay and a well-established Net1-GFP strategy, we found that compaction along the longitudinal axis of rDNA is dramatically increased in response to elevated temperature. Conversely, genome-wide compaction along the chromosome arms is resistant to temperature-induced hypercondensation. In the context of cell physiology, mechanisms that selectively target rDNA are of critical interest. For instance, nutrient starvation induces rDNA hypercondensation and limits transcription factor accessibility which results in slow growth. On the other hand, increased cell growth kinetics correlate with elevated rates of rDNA transcription and ribosome assembly/maturaiton. In the current study, shifting wildtype cells to an elevated temperature (23°C to 37°C) produced an increased growth rate, consistent with previous findings that transcriptional silencing at rDNA loci is decreased and rRNA levels are increased at elevated temperatures. These observations raise a paradox: wildtype yeast at elevated temperatures exhibit both increased rDNA transcription and growth rates despite the hypercondensation of rDNA. Of the many possibilities, one model that resolves this apparent contradiction is that the shorter rDNA loop axis reflects significantly increased lateral loop extensions that accommodate elevated transcription during robust cell growth (Fig. 7). This transcriptional model is consistent with our results that exclude mechanisms of hypercondensation through accelerated condensation reactions and rDNA repeat reduction through recombination. Future efforts will be required to expose the mechanisms through which these potential longitudinal and lateral segment tetherings are regulated and document the extent to which transcriptionally active loops emerge laterally from the chromosomal axis.

Table 1. Yeast strains used in this study.

| Strain name | Genotype | Reference |
|-------------|----------|-----------|
| YBS3017     | mcd1-1, MATa; lacOs::YLR003c-1; lacOs::MMP1; Lac1-GFP; w303 | For this study |
| YBS2078     | MATa: lacOs::YLR003c-1; lacOs::MMP1; Lac1-GFP; w303 | 24 |
| YBS2020     | MATa; NET1::GFP::HIS3; w303 | 70 |
| YBS1019     | MATa; S288C | 17 |
| YBS982      | MATa trp1 ura3 bar1 gal1 | 17 |
| YBS555      | MATa mcd1-1 trp1 leu2 bar1 gal1 | 17 |

Figure 7. Model of rDNA transcription loop extension and possible mechanisms of hypercondensation. rDNA hypercondensation occurs in cells exposed to elevated temperature, potentially through transcription-dependent increase in axial loop extension (below). Candidates required to drive mitotic hypercondensation may include temperature-dependent enhanced activation of cohesin, condensin, or heat shock proteins (either directly or through condensation inhibitor inactivation) that specifically target the rDNA loci (above).
The utilization of temperature-sensitive mutants remains a mainstay of yeast research. In particular, almost every condensation assay performed to date uses a temperature shift from permissive to non-permissive temperatures.\textsuperscript{16,17,19,24-27} Thus, our current study formally raises concerns regarding the extent through which defects in mutant strains (shifted to an elevated and non-permissive temperature) are predicated on comparisons to a hypercondensed rDNA locus. It thus becomes crucial to investigate the mechanism through which condensation is regulated by temperature, improve experimental strategies to better accommodate for those effects and then revisit results from prior studies predicated on conditional alleles and analysis of the rDNA locus. Moreover, our observations are likely to be of clinical interest, given the use of hyperthermia as a cancer treatment.\textsuperscript{68} For instance, the mechanism by which tumor cells become heat sensitive, compared with wildtype cells, is not fully understood. Our study provides a starting point through which defects in mutant strains (shifted to an elevated temperature) are predicated on comparisons to a hypercondensed rDNA locus. Moreover, our observations are likely to be of clinical interest, given the use of hyperthermia as a cancer treatment.\textsuperscript{68}

Materials and methods

Yeast strains and strain construction: Saccharomyces cerevisiae strains used in this study are listed in strain table (Table 1). GFP-tagging and deletion of genes was performed following published protocol.\textsuperscript{69}

rDNA condensation assay: Condensation assays were modified based on published FISH protocol.\textsuperscript{17} Briefly, synchronized cells were fixed by paraformaldehyde (100 \( \mu l \) 36% formaldehyde per 1 ml culture) for 2 hr at 23 °C. Cells were washed with distilled water 3 times and resuspended in spheroplast buffer (1M sorbitol, 20 mM KPO4, pH7.4), then spheroplasted by adding β-mercaptoethanol (1/50 volume) and Zymolyase T100 (1/100 volume) and incubating for 1 hour at 23°C. Resulting cells were pelleted and resuspended in 1.5 pellet volume of spheroplast buffer with 0.5% Triton X-100. 10 \( \mu l \) of the cell suspension were added to each well on poly-L-lysine coated slides, set at room temperature for 10 min, then removed the liquid gently by pipette. 20 \( \mu l \) of 0.5% SDS were added to each well, and set for 10 min, room temperature, then remove the liquid gently by pipette. Air dry the slides. Cells were then dehydrated by immersing the slides in fresh 3:1 methanol:acetic acid for 5 min at room temperature. Slides were stored at 4°C until completely dry, then cells were treated with RNase (100 \( \mu g/ml \)) in 2XSSC buffer (0.3M NaCl, 30 mM Sodium Citrate, pH7.0), incubate 1 hour at 37°C. Slides were washed 4 times in fresh 2XSSC (2 min/per wash), then went through a series of cold (−20°C) ethanol washes (start with 70%, followed by 80%, 95% ethanol washes, 2 min/per wash), then air dry. Slides were prewarmed to 37°C, then put into denaturing solution (70% formamide, 2X SSC) at 72°C for 2 min. Slides were immediately washed through a series of cold (−20°C) ethanol washes (start with 70%, followed by 80%, 90%, 100% ethanol washes, 1 min/per wash), then air dry. DNA mass were detected by DAPI (0.05ug/ml) staining and assayed under microscope. Cell cycle progression were confirmed by detection of DNA content using flow cytometry as described.\textsuperscript{27}

Net1-GFP condensation assay: rDNA condensation assays were performed following similar strategy as previous publications.\textsuperscript{24,27} Briefly, cells were arrested at preanaphase and fixed by paraformaldehyde for 10 min at 30°C. GFP signal were then assayed under microscope. Cell cycle progression were confirmed by detection of DNA content using flow cytometry as described.\textsuperscript{27}

Chromosome arm condensation assay: Chromosome arm condensation assays were performed as previous described.\textsuperscript{24} Briefly, cells were arrested at preanaphase and fixed by paraformaldehyde for 10 min at 30 °C. Distances between GFP dots were measured by microscopy. Cell cycle progression were confirmed by detection of DNA content using flow cytometry as described.\textsuperscript{27} To generate unbiased detection criteria of the GFP distance, cells contain a single GFP dot may reflect sister chromatids positioned vertical to the z-axial focal plane were thus excluded from analysis. In cases where GFP foci were planar to the field of view (eliminated z-axial contributions), we also excluded cells that contained 3 or 4 GFP loci, since the cohesion defect at one or both loci made it impossible to determine which GFP dot represented intra- or inter- sister chromatid loci. Thus, we focused our analysis on cells that contained 2 GFP dots resolvable within a single focal plane.

Statistical Analyses: F-Tests were used to assess the equality of 2 variances in chosen experimental groups, followed by Student’s T-Tests to assess the statistical significance (\( P < 0.05 \)).

Abbreviation

(FISH) Fluorescence in situ hybridization

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

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