Rearrangement of macronucleus chromosomes correspond to TAD-like structures of micronucleus chromosomes in *Tetrahymena thermophila*

Zhengyu Luo,¹,³ Tengfei Hu,¹,³ Hong Jiang,¹ Ruoyu Wang,¹ Qianlan Xu,¹ Simo Zhang,² Jun Cao,¹ and Xiaoyuan Song¹

¹Hefei National Laboratory for Physical Sciences at the Microscale, CAS Key Laboratory of Brain Function and Disease, School of Life Sciences, Division of Life Sciences and Medicine, University of Science and Technology of China, Hefei, Anhui, 230026, China; ²Department of Biology, Indiana University, Bloomington, Indiana 47405, USA

The somatic macronucleus (MAC) and germline micronucleus (MIC) of *Tetrahymena thermophila* differ in chromosome numbers, sizes, functions, transcriptional activities, and cohesin complex location. However, the higher-order chromatin organization in *T. thermophila* is still largely unknown. Here, we explored the higher-order chromatin organization in the two distinct nuclei of *T. thermophila* using the Hi-C and HiChIP methods. We found that the meiotic crescent MIC has a specific chromosome interaction pattern, with all the telomeres or centromeres on the five MIC chromosomes clustering together, respectively, which is also helpful to identify the midpoints of centromeres in the MIC. We revealed that the MAC chromosomes lack A/B compartments, topologically associating domains (TADs), and chromatin loops. The MIC chromosomes have TAD-like structures but not A/B compartments and chromatin loops. The boundaries of the TAD-like structures in the MIC are highly consistent with the chromatin breakage sequence (CBS) sites, suggesting that each TAD-like structure of the MIC chromosomes develops into one MAC chromosome during MAC development, which provides a mechanism for the formation of MAC chromosomes during conjugation. Overall, we demonstrated the distinct higher-order chromatin organization in the two nuclei of the *T. thermophila* and suggest that the higher-order chromatin structures may play important roles during the development of the MAC chromosomes.

[Supplemental material is available for this article.]

Chromatin conformation capture (3C) (Dekker et al. 2002)-related approaches, such as Hi-C and its further developed version, in situ Hi-C, have revealed that mammalian cells have hierarchical chromatin organizations, from larger scale A/B compartments to smaller scale topologically associating domains (TADs) and even smaller scale chromatin loops (Lieberman-Aiden et al. 2009; Dixon et al. 2012; Rao et al. 2014). The binding of the insulator protein CTCF and the cohesin complex on chromatin are enriched at TAD boundaries in mammals, playing important roles in higher-order chromatin organization (Dixon et al. 2012; Rao et al. 2014, 2017). Removal of cohesin or cohesin-loading protein NIPBL eliminated chromatin loops and TADs while enhancing compartmentalization in mammalian cells (Rao et al. 2017; Schwarzer et al. 2017). However, recently a superresolution chromatin tracing technique revealed that the TAD-like structure is present in single cells after cohesin depletion, which indicates that cohesin is not required for maintaining the TAD-like structure in single cells (Bintu et al. 2018). Thus, the mechanism of TADs formation and maintenance is still largely unknown.

*Tetrahymena thermophila*, a ciliated protozoan, contains two nuclei in the same cell: the transcriptionally active polyploid macronucleus (MAC) and the transcriptionally inert diploid micronucleus (MIC) (Yao 1996; Chalker 2008; Mochizuki 2012). Although both the MAC and the MIC develop from the same zygotic nucleus during the sexual reproduction (conjugation) induced by starvation and share 70% of their genome sequences (Hamilton et al. 2016), their genomes differ in chromosome numbers, sizes, and functions. During conjugation, the new MAC develops from the zygotic nucleus and undergoes a programmed genome rearrangement which involves chromosome breakage at chromosome breakage sequence (CBS) sites and DNA elimination of internal eliminated sequences (IESs) (Noto and Mochizuki 2017, 2018). Ultimately, the MAC harbors about 181 stably maintained chromosomes derived from the five chromosomes in the zygote which remain unchanged in the MIC, and the MAC is ≈45–50 N (polyploid) while the MIC is 2 N (diploid) (Hamilton et al. 2016; Lin et al. 2016). However, the higher-order chromatin organization in *T. thermophila* is still unclear.

Recent work in mammals suggested that higher-order chromatin organization is associated with DNA double-strand breaks (DSBs) and chromosomal rearrangements (Canela et al. 2017). Since the development of the *T. thermophila* MAC during conjugation is associated with chromosome breakage at CBS sites and DNA rearrangement, we speculate that the higher-order chromatin organization in *T. thermophila* might be related to chromosome breakage and genome rearrangement in the MAC during conjugation. In addition, there is no CTCF in *T. thermophila* and the cohesin complex is only located in the MIC but not in the MAC.

³These authors contributed equally to this work.

Corresponding author: songxy5@ustc.edu.cn

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Thus, *T. thermophila* provides a naturally controlled system in the same cell for both cohesin depletion (in the MAC) and presence (in the MIC). Therefore, we investigated the higher-order chromatin organization in the two different nuclei of *T. thermophila*.

**Results**

**Determining Hi-C data for analyzing MAC or MIC higher-order chromatin structures**

To examine the higher-order chromatin structures in the two distinct nuclei of *T. thermophila*, we planned to perform in situ Hi-C (Rao et al. 2014) with *T. thermophila* whole cells, isolated MICs, or isolated MACs, respectively, at different stages of conjugation, the sexual life cycle of *T. thermophila*. After starving the cells for 24 h, we mixed two different mating types of *T. thermophila* strains to induce conjugation and collected the *T. thermophila* cells at three different conjugation stages: 1.5 h post-mixing (hpm), when the cells were in premeiosis stage; 3 hpm, when the cells were in meiotic recombination stage, and the MIC underwent transcription and elongation, where the elongated meiotic MIC was also called crescent MIC; and 24 hpm, at the end of conjugation when the new MIC and the new MAC were formed and progenies would be produced upon feeding (Fig. 1A). We then performed in situ Hi-C with *T. thermophila* whole cells, isolated MACs, and isolated MICs at these three different conjugation stages (see Methods).

We first analyzed the reads covered on IESs among our Hi-C data to check the purity of the isolated MICs, as the IESs are only located in the MIC (Yao and Gorovsky 1974; Coyne et al. 1996; Hamilton et al. 2016; Noto and Mochizuki 2018). There are a total of ∼7500 IESs that are precisely mapped on the MIC genome with a total length of about 28.5 Mb (Hamilton et al. 2016; Noto and Mochizuki 2018). We divided the 7500 IESs into three classes with the same combined length (∼9.5 Mb) based on their sizes, and class 1 contained the largest IESs, while class 3 contained the shortest IESs. The results showed that the Hi-C data in *T. thermophila* whole cells had a similar low coverage on IESs compared to that in the isolated MACs (Supplemental Fig. S1A), which suggested that the Hi-C reads generated in the *T. thermophila* whole cells mainly came from the MAC DNA. We thus used the whole *T. thermophila* Hi-C data for the following MAC higher-order chromatin structures analysis with KR normalization (Knight and Ruiz 2012), as the nuclei isolation processes might influence the higher-order chromatin structures in the MAC.

For the Hi-C data generated in the isolated MICs, the coverage on IESs increased compared to that in the isolated MACs or the whole cells but was still lower than that of the other genomic regions (Supplemental Fig. S1), indicating that some of the Hi-C data from isolated MICs were contaminated with MAC genome. Since cohesin component protein Smc1 is only located in the MIC of *T. thermophila* (Howard-Till et al. 2013), to explore the higher-order chromatin organization in the MIC and rule out the effects from the MAC genome, we used HiChIP (Mumbach et al. 2016) with anti-HA antibodies in strains expressing tagged fusion protein Smc1-HA (Howard-Till et al. 2013). The reads coverage on IESs of Smc1-HA HiChIP was higher than that of the isolated MICs Hi-C (Supplemental Fig. S1). For the class 1 and class 2 IESs, the reads coverage of HiChIP data was comparable (or even higher for class 1) to that of the other genomic regions (Supplemental Fig. S1), which indicates that the Hi-ChIP data is more accurate than the Hi-C data for studying the higher-order chromatin structures in the MIC.
suggested that the cohesin HiChIP data mainly came from the MIC with little MAC contamination. We reasoned that IESs in class 3 were too small so that they were lost during the process of Hi-C or HiChIP experiments. We then used Smc1-HA HiChIP data for the following analysis of the MIC higher-order chromatin structures with KR normalization.

**Crescent-specific interactions in the meiotic MIC of T. thermophila**

Our Smc1-HA HiChIP data showed that there were genome-wide crescent-specific chromosome interactions in the MIC only at 3 hpm (Fig. 1B; Supplemental Fig. S2). The crescent-specific inter-arm chromosome interactions existed from telomeres to centromeres on all five chromosomes in the elongated crescent MIC (Fig. 1B). In addition, the crescent-specific interactions also existed between different chromosomes, which formed an X-shaped interaction pattern on the inter-chromosome heatmaps (Fig. 1B, upper right; Supplemental Fig. S2B). These results suggested that the telomeres of all five MIC chromosomes interacted and clustered together, as did all of the five centromeres. Therefore, in the crescent MIC, the two telomeres of one chromosome gathered together, making the chromosome fold at the centromere in the middle, so that centromeres and telomeres were clustered at opposite ends of the nucleus. The X-shaped crescent-specific interactions revealed by our HiChIP data perfectly explained the existence of a meiotic chromosome bouquet in T. thermophila, which was organized by centromeres and promoted inter-homolog recombination (Loidl et al. 2012). We therefore demonstrated this phenomenon at the molecular level as a supplement to the imaging data previously supported by DNA FISH and IF (Mochizuki et al. 2008).

**Optimizing identification of the centromere midpoint based on crescent-specific interactions**

In addition to the above effects, the X-shaped crescent-specific chromosome interactions had other functions in the MIC. Currently, the midpoint of the centromere in T. thermophila is defined as follows: The two CBS sites that are closest to the centromere are designated as R-CBS (on the right chromosome arm) and L-CBS (on the left chromosome arm), respectively; the region between L-CBS and R-CBS is titled the putative centromere region, and the midpoint of this putative centromere region is considered as the midpoint of the centromere (Hamilton et al. 2016). Our cohesin HiChIP data showed that the centromeres were clustering together in the crescent MIC, and there were intra- and inter-chromosome interactions around the centromere region. Thus, the crossover points of the X-shaped crescent-specific chromosome interactions provided another, more accurate way to define the midpoints of centromeres (Fig. 1D,E; Supplemental Table S1). Our cohesin HiChIP data at the crescent stage therefore optimized the identification of the accurate midpoints of centromeres in the MIC.

In addition, we found that there were some square-shaped chromosome interaction boxes that were located far away from the diagonal of the heatmaps (Supplemental Fig. S3), suggesting that these regions were very likely located near each other on the same chromosome but were mistakenly annotated to different chromosome regions. This result indicated that there might be some biases in the current MIC genome, although it did not affect our analysis in this work. Together, these results clearly showed that the higher-order chromatin organization information in the MIC of T. thermophila will be useful in assembling the MIC genome more accurately.

**The MAC chromosomes lack A/B compartments, TADs, and chromatin loops**

As there are two nuclei in the same T. thermophila cell, we analyzed our T. thermophila whole cells Hi-C data using both the MIC genome (Hamilton et al. 2016) and the MAC scaffolds (Coyne et al. 2008). Because the MAC genome assembly remains incomplete, we chose 568 relatively large (longer than 1 kb) and annotated
MAC scaffolds, out of the total 1158, which occupy the majority of the genome (101.92 Mb out of 103.01 Mb). We reordered these 568 MAC scaffolds in the genome according to their relative location on the MIC chromosomes, based on the homology of MAC and MIC, for the following analysis. The results showed that there were square-shaped chromatin interaction patterns along the diagonal of the heatmaps, which were similar to the TADs in mammals, when we mapped the whole cells Hi-C data to the MAC scaffolds (Fig. 2A), but there were no square-shaped chromatin interaction patterns inside the MAC scaffolds, even in the largest ones (Fig. 2A). In addition, we found that every one of the MAC chromosomes, defined by the occupancy of double-end telomeres on the MAC scaffolds, was consistent with one of the TAD-like structures on the heatmaps when mapping the whole cells Hi-C data to the MIC genome (Fig. 2B,C). Thus, although these square-shaped interaction patterns along the diagonal of the heatmaps were similar to TAD structures in mammals, they were consistent with the MAC (whole) chromosomes but did not exist inside the MAC chromosomes, indicating that they were very similar to the chromosome territories in mammals. Additionally, there were no TADs identified by HiCExplorer (Ramírez et al. 2018) with the whole cells Hi-C data when we mapped them to the MAC scaffolds (Supplemental Fig. S4A), and the number of the TADs identified when mapping the whole cells Hi-C data to the MIC genome was close to the number of chromosomes in the MAC (Supplemental Fig. S4A). Thus, our data showed that the TAD-like structures might not exist in the MAC of *T. thermophila*, and the square-shaped interaction patterns along the diagonal of the heatmaps are the MAC chromosome territories.

In addition to TADs, in mammalian cells, Hi-C and related studies revealed that the chromatin is also organized into A/B compartments and chromatin loops (Lieberman-Aiden et al. 2009; Dixon et al. 2012; Rao et al. 2014). PCA analysis of the *T. thermophila* whole cells Hi-C data did not show clear segmentation, whether mapping to the MIC genome or to the MAC scaffolds (Supplemental Fig. S4B), indicating that there were no A/B compartments in the MAC. Although we could identify chromatin loops with the *T. thermophila* whole cells Hi-C data by HiCCUPS (Durand et al. 2016), these chromatin loops could not be confirmed when visually compared to the chromatin loops identified in mammalian cells (Supplemental Fig. S4C,D), suggesting that chromatin loops might not exist in the MAC either. This result was consistent with the fact that there are no reports of distal enhancers or long-range gene regulation in *T. thermophila*. Together, our results showed that the gene-dense, fragmented MAC genome shows no obvious A/B compartments, TADs, and chromatin loops, but each MAC chromosome is a single isolated unit that is similar to the chromosome territory in mammals.

**TAD-like structures are present in the MIC of *T. thermophila***

Next, we examined the A/B compartments, TADs, and chromatin loops in the MIC of *T. thermophila* using Smc1-HA HiChIP data. The results showed that there were no A/B compartments (Fig. 3A) or visible chromatin loops on the heatmaps either, similar to that in the MAC. The heatmaps of Smc1-HA HiChIP, however, clearly showed that there were square-shaped chromatin interaction patterns along the diagonal inside the chromosome (Fig. 3B), which we named TAD-like structures as they were similar to the TADs structure in mammals. There were 166, 190, and 159 TAD-like structures identified by HiCExplorer in the MIC at 1.5 hpm, 3 hpm, and 24 hpm, respectively (Fig. 3C). The number of TAD-like structures in the MIC was close to the number of chromosome territories in the MAC, indicating that TAD-like structures in the MIC might be related to the chromosomes in the MAC. However, although Smc1 is only localized in the MIC, there might still be some MAC contamination in the Smc1-HA HiChIP data; since we could not guarantee 100% ChIP efficiency. To rule out the possibility that the highly similar feature between TAD-like structures in the MIC and the chromosome territories in the MAC might be due to the MAC contamination in our HiChIP data, the ideal way is to isolate 100% pure MICs to perform Hi-C, but it is technically not possible currently.

We then took another way to solve this potential MAC contamination problem. Since IESs are MIC-specific sequences (Hamilton et al. 2016; Noto and Mochizuki 2018), we selected...
the IES-associated interactions, which could only come from the MIC, from the whole cells Hi-C data. Although the IES-associated Hi-C interactions were only about 1.4% of the total paired reads, the IES-associated interaction heatmaps clearly showed that there were square-shaped chromosome interaction patterns along the diagonal of the heatmaps in the MIC (Fig. 4A), which were highly consistent with that on the heatmaps of HiChIP data (Fig. 4B,C). Thus, our data showed that there are no A/B compartments and chromatin loops in the MIC, while there are TAD-like structures in the MIC of *T. thermophila*.

**Boundaries of the TAD-like structures in the MIC coincide with CBS sites**

During conjugation, both the MIC and the MAC are developed from the same zygotic nucleus, and the MIC possesses the five chromosomes as the zygotic nucleus, while the MAC undergoes programmed genome rearrangement with, finally, 181 stably maintained chromosomes (Hamilton et al. 2016; Lin et al. 2016). The MAC chromosomes are formed from the five zygotic chromosomes after chromosome breakage at CBS sites and elimination of IESs (Fig. 5A; Hamilton et al. 2016; Lin et al. 2016). As the number of TAD-like structures in the MIC was similar to the number of chromosomes in the MAC, we asked whether the TAD-like structures on the MIC chromosomes are associated with the formation of the MAC chromosomes. We found that the TAD-like structures in the MIC were all aligned to the chromatin regions between the CBS sites, and the boundaries of the TAD-like structures in the MIC were highly coincident with CBS sites (Fig. 5B; Supplemental Fig. S5). About 57% of CBS sites were exactly localized to the boundaries of TAD-like structures, which was increased to 77.3% when we included the 20-kb surrounding regions of the identified boundaries (Fig. 5C). The averaged chromatin contact frequency centered on boundaries of TAD-like structures or CBS sites were also highly consistent (Fig. 5D,E), suggesting that CBS sites on the MIC chromosomes are the boundaries of the TAD-like structures in the MIC.

**Chromosome territories in the MAC are consistent with TAD-like structures in the MIC**

A comparison of the TAD-like structures in the MIC and the chromosome territories in the MAC revealed clear similarity (Fig. 6A). For some MAC scaffolds that have telomere sequences at both ends, which means that they are MAC chromosomes, the pattern of the whole cells Hi-C heatmaps mapping to the MAC scaffolds (Fig. 6B) or to the MIC genome (Fig. 6C) was highly consistent with the TAD-like pattern on the MIC genome (Fig. 6D). These results together suggested that the TAD-like structures in the MIC might play an important role in the formation of the MAC chromosomes during conjugation. Thus, we proposed a model for the genome organization in the two distinct nuclei of *T. thermophila* during conjugation: The higher-order genome structure (TAD-like structure) may form in the zygotic nucleus by cohesin and/or other factors before the new MAC and the new MIC differentiation; the MIC then retains cohesin and keeps its higher-order chromatin organization, while the cohesin complex disappears in the developing MAC; the higher-order genome structure of MAC chromosomes is directly formed from the higher-order genome structure of MIC chromosomes after chromosome breakage at CBS sites (and elimination at IESs) during the MAC development. Therefore, the chromosome territories in the MAC are similar to the TAD-like structures in the MIC, and these TAD-like structures may function in the development of the new MAC during conjugation (Fig. 6E).

**Discussion**

*T. thermophila* has two distinct nuclei in the same cell, with a different genome and function, while these two nuclei both develop from the same zygotic nucleus during conjugation. However, little is known about the higher-order chromatin organization and its function in this process. Here, we explored the higher-order chromatin organization in *T. thermophila* and found that there are no A/B compartments and chromatin loops in both of the MAC and the MIC. There are also no TADs in the MAC, while the MIC has TAD-like
Boundaries of TAD-like structures in the MIC are consistent with CBS sites. (A) The schematic diagram shows the development of MAC and MIC from the same zygotic nuclei during conjugation. (B) The location of CBS sites, boundaries, and TAD-like structures compared to the heatmaps on the Chromosome 4 (2.5 Mb to 10.5 Mb) at 3 hpm. (C) The bar plot shows percentage of CBS sites localized to the boundaries of the TAD-like structures (10 kb) or 20 kb around the boundaries (total 50 kb). (D) Heatmaps showing averaged Hi-C interaction counts (Smc1-HA HiChIP, 1.5 hpm) around the boundaries of TAD-like structures. (E) Heatmaps showing averaged Hi-C interaction counts (Smc1-HA HiChIP, 1.5 hpm) around the CBS sites.

In addition, we noticed that the TAD-like structures in the MIC of *T. thermophila* have clearer boundaries compared to the TADs identified in mammals. Considering recent advances about TADs in mammals, these clearer boundaries might be the nature of TAD-like structures in the MIC of *T. thermophila*. Recently, single-cell 3D genome studies showed that there are TAD-like structures in the single cell and the TADs detected by Hi-C with a pool of cells are the average of TAD-like structures in individual mammalian cells (Flyamer et al. 2017; Bintu et al. 2018). In a single mammalian cell, any genomic position may be the boundary of a TAD-like structure, while the boundaries are likely to be binding sites of CTCF and cohesin (Bintu et al. 2018). The binding of CTCF and cohesin is highly dynamic in individual mammalian cells with active transcription. Therefore, the TADs obtained in the Hi-C data of a population of cells (the average result of the superposition of TAD-like structures in each single cell) have the enrichment of CTCF and cohesion at the TAD boundaries, which will be, in turn, a region rather than a point. In contrast, the MIC of *T. thermophila* is transcriptionally inert (except for the transient transcription activity in crescent stage), and its genome may be much more stable compared to the transcriptionally active genome in mammals. Thus, the TAD-like structures in the MIC have clearer boundaries (boundaries are limited to individual points of the CBS sites, not to relative wide region as in mammals).

In mammals, TADs are reorganized during spermatogenesis and are almost lost during meiosis (Vara et al. 2019; Wang et al. 2019). However, we observed that TAD-like structures were still clear in the MIC at 3 hpm during conjugation when *T. thermophila* cells enter meiosis. There were, however, crescent-specific X-shaped chromosome interactions in the MIC at this stage which was similar to the X-shaped chromosome interactions in meiotic prophase chromosomes during mouse spermatogenesis (Patel et al. 2019). The TADs are lost from mouse meiotic prophase chromosomes when the X-shaped chromosome interactions formed (Patel et al. 2019), and we speculated that the TAD-like structures might also be lost or attenuated from the MIC chromosomes at 3 hpm when the crescent-specific X-shaped chromosome interactions formed. The reason for the apparent TAD-like structures at 3 hpm is very likely the fact that the percentage of the crescent MICs in the conjugation is <40% (Loidl et al. 2012). Thus, we could not obtain pure crescent MICs, and the crescent-specific interactions were generated from the crescent MICs while the TAD-like structures might come from the MICs at other stages during conjugation. Hi-C or HiChIP with 100% pure crescent MICs or single-cell Hi-C methods (Nagano et al. 2013, 2017; Ramani et al. 2017; Bintu et al. 2018) may help us to answer these questions in the future.
CTCF is found to play important roles in TADs organization in mammals (Dixon et al. 2012; Phillips-Cremins et al. 2013; Rao et al. 2014), and TADs are almost lost after depletion of CTCF in mammalian cells (Nora et al. 2017). However, there are also some species that have TAD-like structures but do not have a CTCF ortholog (Le et al. 2013; Hsieh et al. 2015; Marbouty et al. 2015), similar to T. thermophila. However, in budding yeast or bacteria, which have no CTCF ortholog, the boundaries of the TAD-like structures are enriched in genes with high transcription activities (Le et al. 2013; Hsieh et al. 2015, Marbouty et al. 2015). Thus, there is a CTCF-dependent pathway for TADs formation in mammalian cells and also a CTCF-independent but transcription-dependent pathway for TAD-like structures formation in some species without a CTCF ortholog. The MIC of T. thermophila does not even have transcription activity at most times, suggesting that the formation of the TAD-like structures in the MIC may be independent of CTCF or transcription. We showed that the boundaries of TAD-like structures in the MIC were consistent with CBS sites, indicating that the CBS sites-related proteins or other factors, such as chromatin modifiers, may function in the formation of TAD-like structures in the MIC of T. thermophila. Future works to identify binding proteins and chromatin features at CBS sites and their nearby regions will disclose the underlying mechanism of TAD-like structures formation in the MIC of T. thermophila, which will also help to understand the mechanism of the formation of MAC chromosomes during conjugation. Taken together, our work provided the first genome-wide view of the higher-order chromatin organization in T. thermophila, which carries two distinct nuclei in the same cell, and revealed the relationship between the TAD-like structures in the MIC and the chromosomes in the MAC. These findings have the potential to reveal unique insights into how higher-order chromatin organization is established and maintained.

Methods

T. thermophila culture, starvation, and conjugation

T. thermophila strains B2086, Cu428, B2086 with Smc1-HA (gift from Dr. Josef Loidl), and Cu428 with Rec8-GFP (gift from Dr. Josef Loidl) were used for Hi-C and HiChIP experiments. Cells were cultured at 30°C in 1x SPP medium (10 g protease peptone, 2 g dextrose, 1 g yeast extract, 0.03 g Sequestrene (Fe-EDTA), adding water to final 1 L and autoclave prior to use). For conjugation, cells were first starved in 10 mM Tris-HCl pH 7.4 to a final density of ~2.5 x 10^7 cells/mL for 24 h at 30°C without shaking. Equal numbers of cells representing two different mating types were mixed and incubated at 30°C without disturbing, until cells were collected at 1.5 h, 3 h, or 24 h for the subsequent experiments.

Collection of mating T. thermophila cells for Hi-C

We added 37% formaldehyde (Sigma-Aldrich) to mating cells in 10 mM Tris-HCl pH 7.4 to make final a 1% concentration, mixed well, and incubated for 10 min at room temperature, then quenched the formaldehyde with final 125 mM glycine (Sigma-Aldrich), mixed well, and incubated for 5 min at room temperature, followed by 15 min on ice. We centrifuged the cells at 800 g at 4°C and discarded the solution, then washed the cell pellet with ice-cold 10 mM Tris-HCl pH 7.4. We then centrifuged again and discarded the solution, transferred cells to 1.5-mL tubes and removed as much of the residual solution as possible. The cell samples were stored at -80°C for future experiments or immediately used.

T. thermophila MAC and MIC isolation

We resuspended 2 x 10^7 fixed cells in 1 mL of Hi-C lysis buffer (20 mM Tris-HCl pH 8.0, 100 mM NaCl, 20 mM KCl, 1.5 mM MgCl2, 1% NP-40, 1x PIC), and incubated at 4°C for 30 min with slow rotation. Then we centrifuged at 800g 4°C for 2 min and washed once with ice-cold lysis buffer. Next, we resuspended cells

Figure 6. The MAC chromosomes are related to the MIC TAD-like structures during conjugation. (A) The heatmap regions (Chromosome 1: 3.2 Mb to 7.2 Mb) of the MIC (Smc1-HA HiChIP at 3 hpm, upper) and the MAC (whole cells Hi-C at 3 hpm, below) compared to the CBS sites on the MIC and the MAC scaffolds are plotted. (B) The heatmap showing the Hi-C interactions on six of the MAC scaffolds at 3 hpm (whole cells Hi-C data mapped to the MAC scaffolds). (C) The heatmap showing the Hi-C interactions on the MIC Chromosome 4 (4.11 Mb to 8.14 Mb) at 3 hpm (whole cells Hi-C data mapped to the MIC genome). (D) The heatmap showing the Hi-C interactions on the MIC Chromosome 4 (4.11 Mb to 8.14 Mb) at 3 hpm (Smc1-HA HiChIP data mapped to the MIC genome). (E) A model for the MAC chromosomes development during the conjugation of T. thermophila. The TAD-like structures established in the zygotic nuclei were followed by chromosome breakage at CBS sites.
in 400 μL of 0.5% SDS (Sigma-Aldrich) and incubated at 62°C for 6–12 min with shaking (checked by DAPI stain) until cell membranes were broken (not waiting for all cell membranes to be broken; we stopped when 30%–50% were broken). We put the tubes on ice and quenched the SDS by adding 400 μL of 5% Triton X-100 (Sigma-Aldrich). Nuclei were checked by DAPI stain, using dounce (Sigma-Aldrich) if necessary. Then, we centrifuged at 780 g for 10 min at 4°C, transferred the solution which contains MIC to a new tube (the pellet which contains MAC was also collected and washed twice with 1× NEBuffer 2.1), centrifuged the tubes with MIC again at 780 g for 10 min at 4°C, repeated this step 3–4 times and checked by DAPI stain to see if the MIC was pure. Then, we spun at 8000 g at 4°C for 10 min to pellet the MIC for the following Hi-C experiments.

**Hi-C and HiChIP**

The detailed Hi-C method was described previously (Rao et al. 2014). In brief, we resuspended *T. thermophila* whole cells (after Hi-C lysis), isolated MIC, or isolated MAC in 1× NEBuffer 2.1, and digested overnight with HindIII or MboI (NEB) at 37°C. After inactivating HindIII or MboI by incubating at 65°C for 10 min, we filtered in chromosome ends with dATP (Thermo Fisher Scientific), dTTP (Thermo Fisher Scientific), dGTP (Thermo Fisher Scientific), and biotin-14-dCTP (Thermo Fisher Scientific) at 37°C for 45–60 min with shaking. The DNA fragments were ligated with T4 DNA ligase (NEB) at room temperature for 4 h with slow rotation. After ligation, we incubated at 65°C overnight in the presence of Proteinase K (Thermo Fisher Scientific) to crosslink and then purified the ligated DNA. The Hi-C DNA was then sheared to 200–500 bp and selected by Dynabeads MyOne Streptavidin C1 beads (Thermo Fisher Scientific). Then, we prepared the sequencing libraries on beads, with end-repair, dA-tailing, and adapter ligation. After PCR, we purified the DNA between 300 and 500 bp with AMPure XP beads (Beckman Coulter). The Hi-C libraries were sequenced using the Illumina HiSeq platform. For HiChIP, we generated Smc1-HA HiChIP using standard methods with HindIII or MboI digestion (Mumbach et al. 2016). The antibody used in HiChIP is: anti-GFP antibody-ChIP grade (Abcam ab290).

**Hi-C and HiChIP data processing**

The reads coverage on IES regions was calculated by deepTools (Ramírez et al. 2014) and normalized by reads per kilobase per million mapped reads (RPKM). The paired-end Hi-C and HiChIP sequencing data were mapped and processed through Hi-C Pro 2.10.0 (Servant et al. 2015) to build the contact matrix. The paired reads were mapped to the MIC genome (Hamilton et al. 2016) or the MAC scaffolds (Coyne et al. 2008) reordered by their localizations on the MIC chromosome by Bowtie 2 (Langmead and Salzberg 2012). The Hi-C matrices were then transferred to Juicebox .hic format for visualization or further analysis with KR normalization. The IES-associated interactions were selected from the whole cells Hi-C data by HiC-Pro 2.10.0 while setting KR normalization. The IES-associated interactions were selected on the MIC chromosome by BWA (Li and Durbin 2009). The Hi-C matrix was then generated by hicBuildMatrix tool in HiCExplorer with 10-kb bin size and then KR balanced by hicCorrectMatrix tool. The A/B compartments analysis was performed with the hicPCA tool, while the TADs and boundaries were found by the hicFindTADs tool. For chromatin loops, we identified chromatin loops at 5-kb, 10-kb, and 25-kb resolution and merged in our Hi-C and HiChIP data of *T. thermophila* together with mES Hi-C data (Borev et al. 2017) using CPU version of HiCCUPS (Durand et al. 2016).

**Data access**

The raw and processed sequencing data generated in this study have been submitted to the NCBI Gene Expression Omnibus (GEO; https://www.ncbi.nlm.nih.gov/geo/) under accession number GSE140429.

**Competing interest statement**

The authors declare no competing interests.

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**Author contributions:** X.S. conceived and designed the project. Z.L., T.H., and Q.X. performed the experiments. Z.L., T.H., H.J., R.W., and S.Z. analyzed the data. All authors discussed and interpreted results. X.S., Z.L., and T.H. wrote the manuscript. All authors read and approved the final manuscript.

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