TALE-light imaging reveals maternally guided, H3K9me2/3-independent emergence of functional heterochromatin in Drosophila embryos

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Metazoans start embryogenesis with a relatively naïve genome. The transcriptionally inert, late-replicating heterochromatic regions, including the constitutive heterochromatin on repetitive sequences near centromeres and telomeres, need to be re-established during development. To explore the events initiating heterochromatin formation and examine their temporal control, sequence specificity, and immediate regulatory consequence, we established a live imaging approach that enabled visualization of steps in heterochromatin emergence on specific satellite sequences during the mid-blastula transition (MBT) in Drosophila. Unexpectedly, only a subset of satellite sequences, including the 359-base-pair (bp) repeat sequence, recruited HP1a at the MBT. The recruitment of HP1a to the 359-bp repeat was dependent on HP1a’s chromoshadow domain but not its chromodomain and was guided by maternally provided signals. HP1a recruitment to the 359-bp repeat was required for its programmed shift to later replication, and ectopic recruitment of HP1a was sufficient to delay replication timing of a different repeat. Our results reveal that emergence of constitutive heterochromatin follows a stereotyped developmental program in which different repetitive sequences use distinct interactions and independent pathways to arrive at a heterochromatic state. This differential emergence of heterochromatin on various repetitive sequences changes their replication order and remodels the DNA replication schedule during embryonic development.

[Keywords: late replication; heterochromatin formation; satellite sequence; mid-blastula transition; HP1a; Drosophila]

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Very early in metazoan embryogenesis, epigenetic reprogramming remodels nucleosomes, changes histone modification patterns, and, in many organisms, refashions DNA methylation. Active mechanisms, which are most heavily investigated in mammals, along with the passive dilution effect caused by the rapid cell divisions characteristic of many early embryos, are thought to reset the newly formed zygotic genome to a relatively naïve state to restore totipotency (Fadloun et al. 2013; Farrell and O’Farrell 2014). As the embryo continues developing, the epigenetic constraints are progressively restored and thus canalize the cells toward distinct cell fates. Although it lies at the heart of developmental programing, we have only rudimentary ideas about how embryonic restoration of epigenetic marks is controlled.

Selective silencing of the genome by heterochromatin formation is one of the most important epigenetic events (Beisel and Paro 2011). In multicellular organisms, two types of heterochromatin exist. Constitutive heterochromatin, characterized by enrichment of dimethylated or trimethylated H3K9 [H3K9me2/3] and HP1a, silences genomic regions enriched with tandem repeats of DNA motifs (also known as satellite sequences) and remnants of transposable elements (TEs) (Elgin and Reuter 2013), whereas facultative heterochromatin represses selective domains of euchromatin in particular cells, thereby participating in cell fate determination during development. In early embryos, most signatures of both types of heterochromatin are erased and have to be re-established de novo. How to deploy these powerful silencing systems accurately is a developmental challenge. Here, by focusing on the formation of the constitutive heterochromatin on different satellite sequences in Drosophila embryos, we start to unveil the molecular process of transmisson and re-establishment of epigenetic regulation during embryogenesis.
Like many other organisms, *Drosophila* begins embryonic development with rapid cell divisions (Farrell and O’Farrell 2014). Following the first gononomic division, during which the male and female pronuclei meet, the zygotic nuclei undergo seven mitotic divisions (cycles 2–8) that are perhaps the shortest cell cycles ever documented (8.6 min for every division), resulting in a shell of nuclei inside a shared cytoplasm—the syncytium. At cycle 9, this shell of nuclei migrates to the surface of the embryo and forms the blastoderm. The nuclei then continue dividing rapidly another four times (cycles 10–13), with progressively increased interphase durations from ~9 min in cycle 10 to 14 min in cycle 13. This commitment to the cell cycle quickly expands the number of zygotic nuclei. Many other cellular processes and developmental events are deferred until the slowing of the cell cycle at cycle 14. The interphase of cycle 14 is abruptly lengthened to ≥70 min depending on the spatial position of the cell in the embryo. In this lengthened interphase 14, cortical nuclei are cellularized, many maternal mRNAs and proteins are degraded, and zygotic transcription is fully activated. The embryonic development thus switches from maternal to zygotic programming (Lee et al. 2014). This major embryonic transformation is called the mid-blastula transition (MBT).

Various satellite sequences in somatic cells are packaged into constitutive heterochromatin, which has unique attributes, including high compaction, enrichment of H3K9me2/3 and HP1a, transcriptional quiescence, and late replication. Most of these attributes are absent in the preblastoderm embryos, and the satellite sequences seem to take on these features successively as the embryo develops. Analysis of the C-banding patterns of the mitotic chromosomes (Vlassova et al. 1991) as well as assessment of the volume of a particular satellite sequence (Shermoen et al. 2010) suggest that satellite DNA becomes compacted prior to the blastoderm stage. However, neither the establishment of H3K9 methylation in heterochromatic regions nor the H3K4 methylation in euchromatic regions was observed until interphase 14 (Rudolph et al. 2007). Using a heat-shock-driven *lacZ* transgene inserted near satellite sequences, it was reported that the silencing activity of constitutive heterochromatin was first detectable in the gastrulating embryo, which is approximately in G2 of interphase 14 (Lu et al. 1998). In addition, BrdU or dUTP incorporation experiments revealed that satellite sequences became late-replicating in interphase 14 (Edgar and O’Farrell 1990, Shermoen et al. 2010). These observations imply that constitutive heterochromatin is matured around the time of MBT. The early embryonic formation of constitutive heterochromatin seems to play a unique role in long-term programming of gene expression, as selectively compromising HP1 function in early embryos by genetic manipulations influenced transcriptional silencing activity of heterochromatin in adults (Gu and Elgin 2013).

One caveat of the studies described above is the lack of spatial resolution. Constitutive heterochromatin contains many compositionally different satellite sequences located at distinct genomic positions. Without analyzing the behavior of each individual satellite sequence, our understanding of constitutive heterochromatin formation would be oversimplified. Here, by using the recently developed transcription activator-like effector (TALE)-light method (Yuan et al. 2014), we characterized the formation of heterochromatin on satellite sequences in *Drosophila* early embryos at unprecedented spatial–temporal resolution. We observed, at the time of MBT, a stereotyped program of introduction of molecular markers of heterochromatin on some repetitive sequences [359 base pairs (bp)] but not on the others [1,686]. The resultant change of local chromatin landscape altered the order of replication of those repeats. Moreover, we interrogated the molecular mechanism of heterochromatinization of the 359-bp repeat and uncovered that signals from the mother guided the initial accumulation of HP1a at the 359-bp loci independently of the histone mark H3K9me2/3. This study highlights the sophistication of the programs that introduce heterochromatic features to different regions of the genome within a nucleus and generates new insight into the inheritance of constitutive heterochromatin.

**Results**

**Establishment of repressive histone modifications at the MBT**

Previously, we observed that the replication of one satellite sequence, the 359-bp repeat, was only moderately delayed at the onset of late replication in interphase 14 but shifted to a much later time in interphase 15 (Fig. 1A; Shermoen et al. 2010; Yuan et al. 2014). We wanted to probe the mechanism of this selective sudden delay in replication and reasoned that local epigenetic changes might be involved. DNA methylation is absent in *Drosophila melanogaster* early embryos (Zemach et al. 2010), and histone modifications play a key role in early embryonic epigenetic reprogramming. We thus analyzed several representative histone modifications during the MBT.

While there was almost no detectable immunostaining with antibodies targeting H3K9me2 or H3K9me3 prior to interphase 13, staining for H3K9me2 and H3K9me3 increased dramatically in interphase 14, with increases in H3K9me2 slightly preceding H3K9me3. This staining localized to the apical pole of nuclei where the pericentric sequences surround the clustered centromeres (Fig. 1B). Acetylation of H3K9 [H3K9ac], which is exclusive of H3K9me2/3 and marks euchromatin on chromosome arms, was also undetected in the early embryo but appeared in cycle 12 (Supplemental Fig. S1A). Interestingly, staining for H3K27ac as well as H4ac was readily detectable as early as in the preblastoderm embryos (Supplemental Fig. S1C,D), but its level decreased slightly in mid–late interphase 14 when H3K27me3 started to appear (Supplemental Fig. S1B).

In summary (Fig. 1C), during the early embryonic development, histone acetylation appears early, accompanying the rampant DNA replication and the gradual activation of the zygotic genome (Li et al. 2014). On the other hand, the repressive histone modifications H3K9me2/3...
and H3K27me3 do not significantly accumulate until the much prolonged interphase 14, suggesting that the slowing of cell cycle at the MBT provides sufficient time for the establishment of repressive histone modifications.

Different repetitive sequences acquire distinct chromatin features at the MBT

Various highly repetitive sequences that occupy >20% of the *D. melanogaster* genome are almost always heterochromatic and comprise “constitutive heterochromatin” (Wei et al. 2014). The appearance of H3K9me2/3 in interphase 14 marks a step in embryonic formation of constitutive heterochromatin. To probe whether each type of repeat sequence responds similarly to the developmental cues, we analyzed the emergence of histone modifications at the subheterochromatic level on different satellite sequences.

We previously developed protein probes called TALE-light that can be programmed to recognize a given DNA sequence (Fig. 2A). Injection of a TALE-light at an optimal concentration highlighted particular satellite sequences live without disrupting embryogenesis (Supplemental Movie S1). Similarly, staining of fixed embryos with the TALE-light lit up the target sequences (Supplemental Fig. S2A). The dynamics of the mitotic chromosome seemed to affect the TALE-light’s recognition of its target, as the TALE-light signal went down during mitosis in live embryos but not in fixed samples (Supplemental Fig. S2B).

We made TALE-lights that recognize two major satellite sequences in the *D. melanogaster* genome: 359-bp and 1.686 (Yuan et al. 2014). Costaining of the TALE-light and anti-histone 3 antibody suggested that these
repetitive regions contained nucleosomes in the early embryo (Supplemental Fig. S2C,D), although the AT-rich 1.686 region seemed to either be less compact or have lower nucleosome occupancy (Krassovsky and Henikoff 2014) when compared with adjacent regions. Early embryos showed little or no staining for most of the histone modifications tested in these two repetitive regions, except for trace staining of acetylated H3K27 and H4 at the 359-bp loci early on (Supplemental Fig. S2F). In interphase 14, both H3K9me2 or H3K9me3 marks on the 359-bp repetitive sequences by combining the TALE-light imaging with the Fab-based live endogenous modification [FabLEM] technique [see the Materials and Methods]. Time-lapse images of nuclei in the injected interphase 14 embryos are shown. The reformation of the interphase nucleus is set to be 00:00 (minutes:seconds). The Fab fragment recognizing the H3K9me2 mark is shown in green, and TALE-lights are in red. The dotted circles in the top panels outline the TALE-light-labeled regions. Bars, 5 µm.

Figure 2. Differential accumulation of methylated H3K9 on the 1.686 and 359-bp repeats. (A) Schematic of the in vivo and in vitro applications of the TALE-lights. (B) The appearance of H3K9me2 and H3K9me3 marks on the 359-bp repetitive sequence during embryogenesis. Fixed embryos at different developmental stages were stained with antibodies and TALE-light targeting the 359-bp repeat. H3K9me2 is shown in green, H3K9me3 is shown in red, TALE-light staining of the 359-bp repeat is shown in blue, and DNA is shown in magenta. Arrows point to the 359-bp locus. All of the images were captured and processed using the same microscopic settings so that the fluorescent intensities faithfully reflect the relative quantities of the indicated modifications. Bar, 5 µm. (C,D) Both H3K9me2 (C) and H3K9me3 (D) accumulate on the 359-bp repeat (red dotted circles) but not on the 1.686 repeat (blue dotted circles) in cycle 14. Fixed interphase 14 embryos were stained with antibodies and TALE-lights. H3K9me2 or H3K9me3 is shown in green, and TALE-light-stained 359-bp and 1.686 repeats are shown in red and blue, respectively. Bars, 5 µm. (E,F) In vivo live imaging of accumulation of the H3K9me2 mark on particular repetitive sequences by combining the TALE-light imaging with the Fab-based live endogenous modification [FabLEM] technique [see the Materials and Methods]. Time-lapse images of nuclei in the injected interphase 14 embryos are shown. The reformation of the interphase nucleus is set to be 00:00 (minutes:seconds). The Fab fragment recognizing the H3K9me2 mark is shown in green, and TALE-lights are in red. The dotted circles in the top panels outline the TALE-light-labeled regions. Bars, 5 µm.
Developmental regulation of heterochromatin formation on the 359-bp repeat

The differential accumulation of H3K9me2/3 during cycle 14 suggests heterochromatin formation on the 359-bp repeat but not on the 1.686. It is thought that H3K9me2/3 binds and recruits HP1a, and the establishment of a stable HP1a-bound state underlies or at least marks the molecular process of heterochromatin formation.

To characterize how the 359-bp repeat is heterochromatinized, we made recombinant HP1a protein with a GFP tag fused to its N termini (Supplemental Fig. S3A). Localization of the GFP-tagged HP1a along with the TALE-light probe to visualize the HP1a recruitment step in heterochromatin formation on particular satellite sequences in developing embryos. Western blot analysis showed that the injected exogenous GFP-HP1a was at a concentration comparable with that of the endogenous HP1a (Supplemental Fig. S3B).

At each mitosis, the TALE-light probe targeting the 359-bp repeat was displaced from mitotic chromosomes but soon reaccumulated to the 359-bp loci as the nuclei exited mitosis. We followed the accumulation of GFP-HP1a within the TALE-light-labeled region in each interphase during early embryogenesis (e.g., Fig. 3B, bottom panels, dotted circle) and quantified the enrichment of GFP-HP1a at the 359-bp loci by calculating the fold enrichment over a control area in the same nucleus (Materials and Methods; Supplemental Fig. S3C). From interphase 11 to interphase 13, GFP-HP1a was enriched at several discrete loci in the nucleus, but little was found in the 359-bp region (Fig. 3A, Supplemental Movie S2), which was consistent with the observation that H3K9me2/3 was absent at this stage. Dramatic accumulation of GFP-HP1a at the 359-bp loci was observed in interphase 14 (Fig. 3B, Supplemental Movie S3). After a rapid accumulation phase from 15 min to 30 min into interphase 14, the amount of GFP-HP1 within the 359-bp region reached a plateau (Fig. 3E, blue curve). This HP1a-bound state of the 359-bp repeat was stably maintained thereafter (Fig. 3F, blue curve), except for transient mitotic displacement. As shown in Figure 3C, in interphase 15, the 359-bp loci was decorated by GFP-HP1a right after mitotic exit and became part of the chromocenter.

It is noteworthy that we observed a small amount of GFP-HP1a within the 359-bp region at the end of interphase 13 and even interphase 12 (Fig. 3A, Supplemental Movie S2), suggesting that the rapid cell divisions interrupted recruitment to the 359-bp repeat. To test this, we knocked down all three mitotic cyclins by RNAi to arrest the embryos in interphase 13 (Farrell et al. 2012) and analyzed the accumulation of GFP-HP1a at the 359-bp loci. Indeed, stopping the rapid cell cycle allowed early recruitment of GFP-HP1a to the 359-bp loci (Fig. 3G, orange curve). This result suggests that interphase extension at the MBT is sufficient for the formation of heterochromatin at the 359-bp loci.

The 1.686 repeat, on the contrary, did not recruit GFP-HP1a during interphase 14 (Fig. 3D–F). These observations, based on GFP-HP1a, were confirmed by antibody staining of endogenous HP1a (Supplemental Fig. S3D). Together, our results reveal clear distinctions in the process of heterochromatin formation between different repetitive sequences. As building blocks of the constitutive heterochromatin, it is apparent that different repetitive sequences take different routes to reach the heterochromatic state during embryogenesis.

Ordering the events during heterochromatinization

The heterochromatinization of the 359-bp repeat in interphase 14 provides an opportunity to dissect the underlying molecular mechanism. It is known that H3K9me2/3 helps recruit HP1a, and HP1a promotes the spreading of the H3K9me2/3 mark by recruiting histone methyltransferase (Canzio et al. 2014). We thus tested what initiates the heterochromatinization process on the 359-bp repeat.

The N-terminal chromodomain (CD) of HP1a specifically recognizes the H3K9me2/3 mark, whereas the C-terminal chromoshadow domain (CSD) dimerizes and forms an interface that recruits proteins containing the PxVxL motif, where x is any amino acid (Supplemental Fig. S4B,C). A single amino acid substitution in the CD (V26M) abolishes the recognition of H3K9me2/3, and a substitution (W200A) in the C-terminal extension sequence disrupts the binding of the PxVxL motif without affecting HP1a’s dimerization (Mendez et al. 2011, 2013; Canzio et al. 2014). The V26M and W200A double mutant lost all of the specific localization in interphase nuclei (Fig. 4A; Supplemental Fig. S4A), suggesting that one or the other targeted site is required for HP1a binding at this developmental stage. Interestingly, restoration of a functional CD and hence the H3K9me2/3 binding activity (HP1a-W200A) brought back a subset of the HP1a foci but did not rescue the timely accumulation at the 359-bp locus (Supplemental Movie S4). It was only faintly enriched within the 359-bp region toward the end of interphase 14 (Fig. 4B), and quantification analysis showed that it lacked the rapid accumulation phase seen with the wild type (Fig. 4E). In contrast, HP1a-V26M, which is capable of binding to the PxVxL motif but not the H3K9me2/3 mark, was efficiently recruited to the 359-bp loci (Fig. 4C; Supplemental Movie S5), although the quantification analysis indicated that its accumulation curve plateaued at a reduced height (Fig. 4E), which could simply be due to the increased background or might imply that a deficiency in H3K9me2/3 recognition compromises the late accumulation phase of HP1a. Together, these results suggested that the initial recruitment of HP1a to the 359-bp repeat did not depend on the capability to bind the H3K9me2/3
mark but required an ability to interact with proteins that contain the PxVxL motif. Consistent with this, the HP1α-I191E mutant that cannot dimerize and hence loses the ability to bind the PxVxL ligand (Brasher et al. 2000; Menendez et al. 2011, 2013; Canzio et al. 2014) also failed to accumulate on the 359-bp repeat (Supplemental Fig. S4D). A small amount of GFP-HP1α could be seen in the 359-bp region at the end of interphase 13 (Supplemental Fig. S4A,C, first two columns). We further quantified and compared this premature recruitment among different HP1α mutants. As shown in Figure 4D, no significant reduction was observed for HP1α-V26M when compared with the wild type; however, mutants without the ability to bind the PxVxL motif failed to be recruited at this stage. Thus, we conclude that the initial recruitment of HP1α to the 359-bp region does not depend on binding to H3K9me2/3. This is consistent with the late and gradual accumulation of these methyl marks described above.

### Altering the position of the 359-bp repeat within the nucleus does not alter the timing of HP1α recruitment

We wanted to know what signal guided the HP1α accumulation on the 359-bp repeat. Constitutive heterochromatin is spatially separated from euchromatin in the nucleus of most somatic cells and is often positioned...
adjacent to the nuclear lamina and the periphery of the nucleoli (Padeken and Heun 2014). The unique nuclear position has been suggested to play a role during the establishment of heterochromatin (Jachowicz et al. 2013). We assessed the functional input of nuclear position during the heterochromatinization of the 359-bp repeat. The majority of the 359-bp repeat localizes on the proximal end of the X chromosome. In the scute 8 mutant, which is associated with a major inversion on the X chromosome with an end point within the proximal part of the 359-bp repeat, most of the 359-bp repetitive sequences are translocated to the distal end of the X chromosome, with a small block of the 359-bp repeat remaining at its original centromere-proximal position [Supplemental Fig. S5C,D].

We characterized the GFP-HP1a accumulation at the large, more telomeric block of the 359-bp repeat, which is positioned at the basal pole of the nuclei (Supplemental Fig. S5F), as well as the smaller, more centromeric block of the repeat, which is positioned apically (Supplemental Fig. S5E). GFP-HP1a was recruited to both loci of the 359-bp repeat simultaneously in interphase 14, suggesting that nuclear position at this stage is not a major contributor to the formation of heterochromatin. Quantification indicated that the accumulation of GFP-HP1a to the basal 359-bp locus plateaued at a reduced height (Supplemental Fig. S5G), which might be partially due to the measurement inaccuracies during live embryo imaging caused by the elongating nuclei in interphase 14 (Waters 2009).
which pushed the basal 359-bp locus up to 14 µm away from the coverslip.

Maternal cues guide heterochromatin formation

A widespread feature of early embryogenesis is that the mother preloads the egg with material that directs most of the early developmental programs [Farrell and O’Farrell 2014]. We reasoned that maternal signals might guide the heterochromatin formation on the 359-bp repeat. To test this, we removed the 359-bp repeat from the mother’s genome and evaluated the heterochromatinization of the 359-bp repeat in the offspring embryos.

Repetitive sequences evolve rapidly. Another closely related species, Drosophila simulans, has no 359-bp or 359-bp-like repeat in its genome and can be crossed to D. melanogaster [Ferree and Barbash 2009]. As illustrated in Figure 5A, when a female D. simulans was husbanded with a male D. melanogaster, the F1 hybrid female embryos would have the maternal supply from the D. simulans mother that lacks the 359-bp repeat and would meanwhile obtain a copy of the 359-bp repeat from the D. melanogaster father. Most of these hybrid female embryos exhibited an early embryonic lethality due to chromosome missegregation in the syncytial cycles [Supplemental Fig. S5A,B]. However, a small fraction developed to a later

Figure 5. Maternal cues guide heterochromatinization of the 359-bp repeat. [A] Schematic of the hybrid cross between D. simulans (D. sim) and D. melanogaster (D. mel). Only sex chromosomes are shown. Note that the 359-bp repeat located on the X chromosomes of D. melanogaster is absent in D. simulans. [B, C] TALE-light and H3K9me3 antibody stainings in the control and hybrid female embryos produced from the D. simulans and D. melanogaster cross. The H3K9me3 histone mark [arrows] exists within the 359-bp region in the control [B] but not the hybrid [C] female embryos. H3K9me3 is shown in green, the 359-bp repeat is shown in red, and DNA is shown in blue. Arrows point to the corresponding 359-bp loci. Arrows point to the corresponding 359-bp loci. Bars, 5 µm. [D] Frames from videos at the indicated times [minutes:seconds] show a lack of GFP-HP1a accumulation on the 359-bp repeat (arrowheads) in the hybrid female embryos. Note that the reappearance of the nucleus in each interphase is set to be 00:00. Arrows point to the corresponding 359-bp loci. Bars, 5 µm. [E] Enrichment of GFP-HP1a within the 359-bp region at the end of interphase 13 in 359-bp-positive embryos produced from the indicated crosses. No significant difference was observed in embryos laid by D. melanogaster mothers [unpaired t-test, \( P = 0.0653 \)]; however, embryos from the D. simulans mothers have less GFP-HP1a at the 359-bp loci when compared with control [unpaired t-test, \( P = 0.0002 \)]. Error bars represent the SD. [F] Quantification of GFP-HP1a accumulation in interphase 14 at the 359-bp loci in the 359-bp-positive embryos produced from the indicated crosses. Note that the Zhr\(^{-}\) mutant of D. melanogaster [D. mel:Zhr\(^{-}\)] lacks the 359-bp repeat on the X chromosome but still has many tiny 359-bp derivatives on the autosomes. \( n > 5 \). Error bars represent the SD.
embryonic stage beyond the MBT. We analyzed the H3K9 methylation in these embryos. While H3K9me3 was found on the 359-bp repeat in control interphase 14 embryos [Fig. 5B, arrow], it was absent in the hybrid female embryos [Fig. 5C, arrowheads]. Consistently, in these embryos, the HP1a accumulation at the 359-bp locus was greatly reduced when compared with that in the control or the embryos from the reciprocal cross [Fig. 5D,E; Supplemental Movie S6]. The initial recruitment of HP1a at the end of interphase 13 was also compromised in these embryos. In the reciprocal cross in which D. melanogaster was the mother, embryos exhibited early recruitment of HP1a, whereas embryos from the D. simulans mother failed to do so [Fig. 5E].

The cross-species mating suggests that the maternal signal might depend on the maternal presence of the 359-bp repeat. We wanted to test this with D. melanogaster strains. Besides a major block of 359-bp repeat on the X chromosome, D. melanogaster has many tiny 359-bp derivatives on its autosomes [Wei et al. 2014]. The Zhr1 mutant lacks the majority of the 359-bp repeat due to compound X chromosomes but still has the autosomal 359-bp derivatives [Ferreau and Barbash 2009]. We mated the Zhr1 females with control males and analyzed HP1a accumulation to the paternally derived 359-bp locus in the offspring female embryos. The removal of most of the 359-bp repeat from the maternal genome had a quantitative effect, as the accumulation of HP1a in interphase 14 at the 359-bp locus was significantly reduced in the offspring embryos when compared with that in the control [Fig. 5F; Supplemental Movie S7]. In contrast, in the reciprocal cross in which the normal 359-bp repeat was present in the maternal parent, progeny embryos showed normal accumulation of HP1a, indicating that the effect of the deletion on HP1a recruitment is a maternal effect phenotype [Fig. 5F].

We conclude that maternally provided factors contribute to the heterochromatinization of the 359-bp repeat. This maternal contribution appears to depend on the maternal presence of the 359-bp sequences. Deficits of maternal 359-bp sequences specifically affect the 359-bp locus in the progeny embryos.

Establishment of stable HP1a binding delays the timing of replication in S phase

The observed selective heterochromatinization of the 359-bp repeat in interphase 14 could potentially explain the previously reported sudden delay in replication timing of this repeat in interphase 15 [Yuan et al. 2014]. To directly test whether the establishment of the HP1a-bound state delays replication, we induced HP1a accumulation on the 1.686 repeat and measured its timing of replication. Artificial tethering HP1a to DNA has been shown to induce local heterochromatin formation [Hathaway et al. 2012]. We fused HP1a to the C terminus of the TALE-light probe recognizing the 1.686 repeat and injected a low level of this 1.686-HP1a fusion protein into the embryo to induce ectopic heterochromatin formation [Fig. 6A,B]. The targeted HP1a fusion protein induced robust accumulation of untargeted GFP-HP1a at the 1.686 loci in interphase 15 [Fig. 6D]. Moreover, the replication-coupled decompaction of the 1.686 loci seen in the control embryos was postponed [Fig. 6C,D], indicating a delay in replication. We injected GFP-PCNA to directly visualize the bulk replication of the 1.686 repeat and compared its timing in embryos with or without the 1.686-HP1a. Indeed, the presence of the 1.686-HP1a fusion protein delayed the replication of the 1.686 in interphase 15 [Fig. 6E, Supplemental Fig. S6A].

To examine the consequences of loss of HP1a recruitment, we used the GFP-PCNA reporter to follow the replication timing of the 359-bp repeat in the D. simulans and D. melanogaster hybrid embryos in which the 359-bp repeat failed to recruit HP1a. The replication of the 359-bp locus was advanced when compared with that in the control embryos [Fig. 6F; Supplemental Fig. S6B]. These two sets of experiments provided direct evidence for the idea that heterochromatin formation delays the local timing of replication in S phase. We thus conclude that the sudden delay of replication timing of the 359-bp repeat in interphase 15 is due to the developmentally regulated heterochromatinization of this repeat.

As summarized in Figure 6G, during the syncytial blastoderm stage, high Cdk1 activity and/or the rapid cell cycle inhibit the accumulation of HP1a at the 359-bp loci. At the MBT in interphase 14, down-regulation of Cdk1 [Farrell and O’Farrell 2014] and other developmental inputs [Blythe and Wieschaus 2015] slows DNA replication and extends interphase, allowing the HP1a accumulation and hence heterochromatin formation on the 359-bp repeat. This local change in chromatin landscape impacts the DNA replication schedule in the following cell cycles, as the heterochromatic state of the 359-bp repeat selectively delays its timing of replication relative to the other repeats.

Discussion

TALE-lights: visualizing specific DNA sequences in vivo and in vitro

New tools are needed to probe the complex and dynamic nature of the organization of the eukaryotic genome. For >30 years, FISH has been the dominant method to label a given DNA sequence [Levsky and Singer 2003], but the requirement of denaturation of DNA often compromises the integrity of the sample and hinders its applications in living organisms. The integration of the lacO or TetO repeats into a particular genomic locus and the use of fluorescently labeled LacI or TetR protein provide strategies to visualize DNA targets live [Robinett et al. 1996]. However, these methods lack the flexibility in target selection. Recent breakthroughs have allowed the systematic engineering of DNA sequence recognition, which stimulated the development of several new DNA visualization methods, including techniques based on the zinc fingers [ZFs] [Lindhout et al. 2007], the CRISPR/dCas9 [Chen et al. 2013; Anton et al. 2014; Deng et al. 2015], and the TALEs [Ma et al. 2013; Miyanari et al. 2013; Thanisch...
et al. 2014; Yuan et al. 2014). Our TALE-lights belong to the last. The sequence-specific DNA recognition by a TALE-light comes from the one-to-one binding of the TALE modules to each of the DNA bases. Therefore, its programmability is as flexible as the CRISPR/dCas9 and better than the ZFs, as one ZF domain recognizes a 3-bp 5′-GNN-3′ DNA sequence (Segal et al. 1999). Moreover, our results demonstrate that the TALE-lights work well in both fixed and live conditions. Because of this feature, we believe that the TALE-lights will be a valuable tool for the studies of genome organization. At this stage, we visualized only repetitive DNA elements. By adapting proper signal amplification strategies, single-copy genes might become visible.

The use of exogenous proteins or RNPs to label endogenous DNA elements in live cells could potentially interfere with normal cellular functions. Indeed, it has been reported that the binding of LacI to the lacO repeats blocks DNA replication (Duxin et al. 2014). The use of TALE-lights in our experiments, however, was seemingly compatible with DNA replication, affecting neither cell cycle length nor embryogenesis (Yuan et al. 2014). Since TALE-lights can be used in vitro as a “sequence-specific DNA antibody,” we examined fixed embryos and found that the appearance of histone marks on repetitive sequences was consistent with the dynamics scored in live embryos. This again suggests that TALE-lights binding in vivo did not cause major disruptions. In addition, we noticed that the majority of the TALE-light was displaced from the...
Constitutive heterochromatin formation in development

The ability of the TALE-lights to discriminate individual satellite sequences has given us the capacity to reveal their differences in the establishment of constitutive heterochromatin during early embryogenesis. The 359-bp repeat recruits HP1a and becomes heterochromatinized in interphase 14, and it appears to typify one group of repeat sequences to which the initial recruitment of HP1a does not depend on the presence of H3K9me2/3. This group of repeat sequences was visualized as distinct loci of accumulation of the mutant HP1a (V26M), which is incapable of binding to H3K9me2/3 (Fig. 4C). In contrast, the HP1a mutant with a deficient CSD (W200A) failed to localize to the 359-bp repeat but did localize to several foci in interphase 14 that appear to represent a second group of repeat sequences that recruits HP1a using an alternative mode of interaction, perhaps by CD binding of H3K9me2/3 (Fig. 4B). However, we have yet to identify a TALE-light marking this group of foci, and additional work is needed to characterize the mode of HP1a recruitment to these sites. The third group of repetitive sequences includes 1.686. In terms of the classical hallmarks of heterochromatin, these repeats remain largely naive at the time of the MBT. We do not know the molecular process of heterochromatin formation on these repeats.

While we use the term heterochromatinization to describe the ensemble of changes associated with recruitment of HP1a to the 359-bp satellite, our findings suggest complexity in the processes that establish heterochromatin and some resulting ambiguity in terminology. HP1a and the histone modification with which it associates, H3K9me2/3, are often taken as markers that define heterochromatin. Even though this study provides direct evidence for the functional impact of a transition in chromatin structure marked by HP1a recruitment, the results also highlight the diversity in forms of heterochromatin. The satellite sequences have distinctive features prior to the recruitment of HP1a, including compaction and late replication in cycle 14 (Shermoen et al. 2010). Although the delay in replication of the 359-bp repeat prior to HP1a binding is slight, it is nonetheless delayed in cycle 14. Furthermore, the 1.686 satellite fails to mature to an HP1a-bound form until later in development yet maintains a late replication program. These observations reinforce a perspective that many factors influence the formation of “heterochromatin,” and the degree of uniformity of the resulting chromatin structures remains to be established.

Controlling heterochromatinization in time and space at the MBT

The whole block of the 359-bp repeat undergoes heterochromatinization precisely in interphase 14. How does the embryo know when to form this heterochromatin? As mentioned earlier, fly embryos begin development with extremely fast nuclear divisions characterized by unusually high mitotic Cdk activity (Farrell and O’Farrell 2014). Given that the binding of HP1a and its silencing activity are both regulated by dynamic phosphorylation/dephosphorylation events (Zhao and Eisenberg 1999; Zhao et al. 2001), the high kinase activity and the rapid cell divisions in the early embryo might curb the establishment of the stable heterochromatin state. Indeed, our results showed early accumulation of HP1a on the 359-bp repeat when Cdk1 activity was down-regulated and the cell cycle was arrested, suggesting that factors needed for heterochromatinization of the 359-bp repeat are already present in the early embryos and that developmentally regulated cell cycle slowing provides the first opportunity for their action (Fig. 6G). However, recruitment of HP1a to foci of repeated sequences does not occur immediately in cycle 14, and the precise timing of this recruitment suggests sophisticated and as yet unknown regulatory circuitry.

Notably, constitutive heterochromatin is not always heterochromatic, at least according to the molecular hallmarks of this state. What marks the 359-bp sequences for selective formation of heterochromatin? It is notable that some repeat sequences, such as those of the 359-bp locus, are selectively compacted during early mitotic cycles despite the absence of H3K9me2/3 (Shermoen et al. 2010), suggesting that some intrinsic feature of the satellite sequence might specify its special behavior. However, because only some of the constitutively heterochromatic repeats recruit HP1a and because there have been numerous findings suggesting that maternal signals might act transgenerationally to direct aspects of gene activity, we sought to distinguish between a transacting maternal signal and sequence-autonomous features of the input.

Genetic manipulation of the amount of the 359-bp repeat in the mothers’ genome (using either D. simulans or D. melanogaster Zhr1 mutants as mothers) (see Fig. 5) specifically influenced the heterochromatin formation on the 359-bp repeat in the offspring, whereas changes in the fathers’ genome (the reciprocal crosses) had no such effect. These results suggest that maternal 359-bp sequences contribute to the zygotic establishment of heterochromatin at the 359-bp locus and encourage us to think of the type of signal that might be conveyed from mother to progeny. It has been established that particular RNAs expressed in the maternal germline are processed into a special class of small RNAs called piRNAs that act to silence gene expression. RNAs homologous to the 359-bp repeat were found in the germinal tissues, and the abundance of these transcripts was increased in mutants defective in components of the piRNA pathway, suggesting that transcription of these sequences occurs and that the piRNA pathway targets this expression (Usakin et al. 2007). Current
understanding of piRNA suppression the expression of repeated sequences suggests that copies of the sequence in a specialized piRNA cluster in heterochromatic regions are expressed, processed, and loaded onto an Argonaut protein called Piwi. Piwi in complex with a specific piRNA sequence appears capable of doing three things: directly targeting homologous DNA sequences to promote heterochromatin formation and suppress transcription, contributing to slicing activity that destroys the RNA products of homologous loci, and working in conjunction with other piRNA pathway components to amplify the piRNA signal by processing complementary and homologous RNA (Guzzardo et al. 2013). Piwi has been found to be associated with small RNAs derived from the 359-bp repeat region (Saito et al. 2006), consistent with a potential role of this pathway in regulating heterochromatin formation on these sequences. Our finding that HP1α is recruited to the 359-bp sequence at a specific time in cycle 14 and in a process that depends on the maternal presence of 359-bp sequences could be explained by suggestions that the 359-bp-derived piRNAs are required for timely heterochromatinization of these sequences, although the details of the mechanism remain to be addressed.

**Heterochromatin formation and the timing of replication**

We began with an interest in the developmental onset of late replication. We found that this onset is triggered by the down-regulation of mitotic cyclin:Cdk1, which, if active during interphase, can trigger early replication of otherwise late-replicating sequences [Farrell et al. 2012]. While this finding indicates that a change in a “trans factor” acting on pre-existing heterochromatin might be responsible for this onset, we found that heterochromatic marks were introduced at about the time of onset of late replication [Shermoen et al. 2010]. Additionally, we noted that the onset of late replication is modulated differently at different repeat sequences [Yuan et al. 2014]. These later observations suggest that changes of the chromatin structure of the repeat regions (cis changes) also occur and might modulate the replication timing.

Widespread coupling between heterochromatin and late replication has fostered the idea that heterochromatin is late-replicating. Additionally, the genomic context has been shown to influence the timing of firing of origins of replication, a finding that is generally interpreted as showing that local chromatin structure specifies replication timing. Nonetheless, despite strong connections, experimental support for a causal connection between heterochromatin and replication timing is indirect. Global knockdown of HP1α in *Drosophila* Kc cells by RNAi advanced the replication of centromeric repeats but delayed replication of many other genomic regions (Schwaiger et al. 2010). Such complex effects might be due to indirect effects of chromatin structure on overall genome arrangement [Sexton and Yaffe 2015].

By manipulating HP1α levels locally at specific repetitive loci, our results demonstrated that failure to establish a HP1α-bound state at the 359-bp repeat locus advanced its replication timing in cycle 15 and that experimentally induced TALE-mediated HP1α recruitment to the 1.686 locus delayed its replication in cycle 15. This, together with the natural developmental program of HP1α recruitment to the different satellite sequences, strongly supports a causal connection between HP1α recruitment and replication timing. It should be noted, however, that recruitment of HP1α can engage a number of reinforcing interactions among factors promoting heterochromatin formation. Hence, the results argue for a causal connection, but not necessarily a direct or even simple connection, between HP1α recruitment and replication timing.

HP1α-positive regions unfold in conjunction with PCNA recruitment and replication and with recruitment of repair proteins and repair of DNA damage [Shermoen et al. 2010, Chiolo et al. 2011]. It is presently unclear whether the events of repair and replication unfold the compacted heterochromatic structure or whether a separate decompaction process avails the compacted sequences of these reactions. However, in either case, compaction could impede these processes, and disassembly of the compacted structure might be regulatory. The activation and firing of replication origins involves complex assemblies regulated by several kinases and phosphatases. The dispersal of compacted heterochromatin might also be regulated by the dynamic phosphorylation/dephosphorylation events, and interplay between the structure and the activating processes might govern timing. More future studies are needed to fully uncover the underlying mechanisms of late replication, and we believe that the TALE-light technique will play an important role during this process.

**Materials and methods**

**TALE-light assembly**

Customized TALE arrays recognizing a given repetitive sequence were designed using TALE Nucleotide Targeter 2.0 [http://tale-nt.cac.cornell.edu]. The following target sequences were chosen: 5′-AGACTGTAATAGCTCAGTCTAAAACAGATATT-3′ and 5′-AGCTGCTAAAAACAGATATT-3′ to target the 359-bp repeat and 5′-AGATAACATAGAATAACAT-3′ to target the 1.686 repeat. The TALE arrays were assembled using Golden Gate TALEN and TALE kit 2.0 [Addgene]. Backbone plasmid MR015 was used in the final reaction. The full-length TALE array was then subcloned into the customized PET-28 bacterial expression vector carrying a C-terminal GFP or mCherry tag. These constructs were subsequently used to produce the recombinant TALE-light proteins.

**Fly stocks**

*Drosophila* strains were maintained on standard cornmeal–yeast medium. *D. melanogaster* strains used in this study were as follows: the Sevelen line as the wild type, *scute 8* [full genotype is In[1]sc8,y31Df(1)Zhr, provided by Dr. Daniel Barbash]. In the hybrid cross, 40–50 *D. simulans* virgins were
mated to 50–60 *Sevelen* males in a confined vial for 2 d. Flies from several such vials were pooled into a population cage for embryo collection.

**Protein production and microinjection**

TALE-lights as well as GFP-HP1a and GFP-PCNA were produced in BL21 (DE3)-competent *Escherichia coli* cells (Bioline). Briefly, BL21 (DE3) transformants cultured in Luria-Bertani medium were treated with 0.5 mM IPTG for 12 h at room temperature to induce protein expression. Bacteria were pelleted by centrifugation and resuspended in lysis buffer (20 mM Tris at pH 7.9, 500 mM NaCl, 10 mM imidazole). Next, the bacteria were incubated for 1 h on ice in the presence of 0.1 mg/mL lysozyme and 10 µM PMSF and further lysed by sonication (ultrasonic liquid processors, Misonix). The recombinant protein in the cleared bacterial lysate was purified using Ni-NTA agarose (Macherey-Nagel). After thorough washes with wash buffer (20 mM Tris at pH 7.9, 500 mM NaCl, 20 mM imidazole), the protein was eluted in elution buffer (20 mM Tris at pH 7.9, 500 mM NaCl, 300 mM imidazole) and then dialyzed into 40 mM Hepes (pH 7.4) and 150 mM KCl.

The microinjection was performed as previously described [Farrell et al. 2012]. The TALE-lights were used at 1 µg/µL, GFP-HP1a was used at 6 µg/µL, and GFP-PCNA was used at 2 µg/µL. The fusion protein 1.686-HP1a was used at 0.1 µg/µL to induce ectopic HP1a recruitment.

**Immunofluorescence and TALE-light staining**

Embryos were collected on grape agar plates, dechorionated for 2 min in 50% bleach, and fixed in methanol-heptane (1:1) for 5 min. The fixed embryos were stored in methanol at −20°C. Before immunostaining, the embryos were first rehydrated gradually [5 min each in 1:3, 1:1, and 3:1 PTA:methanol and then 10 min in PTA]. PTA consisted of PBS supplemented with 0.1% Triton X-100, and 0.02% azide. The embryos were then blocked in PBT [PTA plus 1% BSA] for 30 min and incubated with primary antibodies [1:100 in PBTA] for 1 h at room temperature or overnight at 4°C. The following primary antibodies were used: anti-Histone 3 [Abcam, ab17971], anti-H3K9me1 [Active Motif, 39249], anti-H3K9me2 [Active Motif, 39683 and 39735], anti-H3K9me3 [Active Motif, 39765; Millipore, clone CMA308], anti-H3K9ac [Abcam, ab10812], anti-H3K27ac [Abcam, ab4729], anti-H3K27me3 [Cell Signaling, 9733], and anti-H4ac [Millipore, 06-598]. The embryos were washed three times for 5 min each in PBTA and incubated with the appropriate fluorescently labeled secondary antibodies (Molecular Probes) for 1 h in the dark at room temperature. They were then washed four times for 5 min each in PBTA and mounted in VectaShield mounting medium with DAPI (Vector Laboratories). For TALE-light stainings, GFP- or mCherry-tagged purified TALE-light protein (1:500) was included during the incubation with secondary antibodies.

**FabLEM imaging**

All the FabLEM reagents were kindly provided by Dr. Hiroshi Kimura. Briefly, histone modification–specific monoclonal antibodies were treated by protease digestion to generate Fab fragments, and the Fab fragments were then labeled with amine-reactive fluorescent dyes to make the FabLEM probes [Hayashi-Takanaka et al. 2011]. Under many experimental conditions, the FabLEM did not disturb normal cellular functions, including the HP1a binding in our experiment, probably because the binding between the Fabs and their targets is very transient and dynamic [Hayashi-Takanaka et al. 2011; Sato et al. 2013; Staveich et al. 2014; Kimura et al. 2015].

**Imaging, data quantification, and interpretation**

Embryos were imaged on a spinning-disk confocal microscope as previously described [Farrell et al. 2012], and the images were analyzed using Volocity 6 (Perkin Elmer). When an individual repetitive sequence was imaged, the Z-scale was selected based on the TALE-light signals [usually 3–4 µm], and the step size in the Z-axis was 0.5 µm for fixed samples and 1 µm for live embryos. All of the images in one experiment were acquired and processed using identical microscopic settings. In the TALE-light staining experiments, a single optic section across the indicated repetitive loci was shown.

For the quantification of HP1a recruitment, optic sections containing the specific genomic loci were projected and corrected for photobleaching in Volocity. In each experiment, three to seven embryos and five to seven nuclei in each embryo were selected for quantification. In each nucleus, the mean intensity of GFP-HP1a at the TALE-light-positive loci was divided by that at a control locus in that same nucleus to calculate the fold enrichment [Supplemental Fig. S3C]. The mean and SD of the calculated fold enrichment in nuclei from different embryos were plotted against time, and then nonlinear regression analysis was performed using Prism (GraphPad).

The results shown in this study were obtained with injected recombinant GFP-HP1a protein. We also performed the experiments with GFP-HP1a expressed from a transgene, and the results were identical. HP1a functions as a dimer, and thus a fraction of the injected GFP-HP1a mutant proteins could form heterodimers with the endogenous wild-type HP1a. The possible existence of the heterodimer pool complicated the experimental scenario but did not influence the interpretation of the contributions of different domains to the initial recruitment of HP1a.

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