Methylation of histone H4 lysine 20 by PR-Set7 ensures the integrity of late replicating sequence domains in *Drosophila*

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Abstract

The local chromatin environment is a key regulator of DNA-templated processes including transcription, DNA replication, and DNA repair. The methylation state of lysine 20 on histone H4 (H4K20) has been linked to cell cycle progression, origin recognition complex (ORC) binding, pre-Replication Complex (pre-RC) assembly, and activation of replication origins. Monomethylation of H4K20 (H4K20me1) is mediated by the cell cycle-regulated histone methyltransferase PR-Set7. PR-Set7 depletion in mammalian cells results in defective S-phase progression and the accumulation of DNA damage, which could be partially attributed to a defect in pre-RC formation and origin activity. However, these studies were limited to only a handful of mammalian origins, and it remains unclear how PR-Set7 and H4K20 methylation impact the replication program on a genomic scale. Using *Drosophila* Kc167 cells, we employed genetic, cytological, and genomic approaches to better understand the role of PR-Set7 and H4K20 methylation in regulating DNA replication and governing genome stability. We find that depletion of *Drosophila* PR-Set7 and loss of H4K20me1 result in the accumulation of DNA damage and an ATR-dependent cell cycle arrest. The cell cycle arrest occurs during the second S phase following loss of PR-Set7 activity, suggesting that accumulation of nascent H4K20 is recalcitrant to the DNA replication program. Deregulation of H4K20 methylation had no impact on origin activation throughout the genome; instead, we found that the DNA damage marker, phosphorylated H2A.v (y-H2A.v), accumulated specifically in late replicating domains in the absence of PR-Set7. We conclude that PR-Set7-mediated monomethylation of H4K20 is critical for maintaining the genomic integrity of late replicating domains.

Author Summary

The complete and faithful duplication of the genome is essential to ensure normal cell division and organismal development. Post-translational modifications (PTMs) of histones modulate the local chromatin environment and thus have the potential to regulate the transcription and DNA replication programs. A number of histone PTMs, including methylation of lysine 20 on histone H4 (H4K20), have been linked to the regulation of the replication program; however, the majority of these studies have been limited in scope, surveying only a few select loci for origin activity. Here we use genome-wide approaches to comprehensively survey the impact of deregulated H4K20 methylation on the *Drosophila* DNA replication program. Deregulation of H4K20 methylation had no impact on origin activation. In contrast, we found that loss of H4K20 monomethylation results in the accumulation of DNA damage at late replicating sequences in S phase and the activation of the ATR-dependent intra-S phase checkpoint. Together, our data suggest that H4K20 methylation is important for replication fork progression and stability to ensure the integrity of late replicating sequences.

Introduction

Histone post-translational modifications (PTMs) regulate almost all DNA-templated processes including DNA replication, transcription and DNA repair. Deregulation of these epigenetic histone modifications has the potential to lead to catastrophic consequences at both the cellular and organismal level. One such epigenetic mark, methylation of histone H4 lysine 20 (H4K20), is critical for maintaining genome stability, and its deregulation impacts transcription, chromatin compaction, DNA repair, cell cycle progression, and DNA replication [[reviewed in 1,2,3]].

Monomethylation of H4K20 (H4K20me1) is catalyzed by the histone methyltransferase PR-Set7/Set8, orthologues of which exist in all metazoans [4,5]. H4K20 can also be di- and tri-methylated by the Suv4-20 h1 and h2 homologs in mammalian cells and a single Suv4-20 in *Drosophila* [6,7]. The levels of mammalian PR-Set7 and H4K20me1 are cell cycle regulated. PR-Set7 harbors a conserved PIP-box motif and undergoes PCNA- and CRL4^{Cdt2}-mediated degradation during S phase [8–12]. This interaction between the PR-Set7 PIP-box and PCNA is conserved in *Drosophila* cell lines, where the levels of PR-Set7 and H4K20me1 display a similar cell cycle oscillation pattern as seen in mammalian systems [13].

Not only are PR-Set7 and H4K2ome1 levels coupled to DNA replication via the PIP-box dependent degradation of PR-Set7, but the DNA replication program is also regulated, in part, by the methylation status of H4K2o. Mammalian cells depleted of PR-Set7 are defective in S phase progression, accumulate DNA damage, and activate the DNA damage response [14–16]. Mechanistically, this may be partially attributed to the role of mammalian PR-Set7 in recruiting pre-Replication Complex (pre-RC) components onto chromatin [11], thus promoting the activity of select replication origins. Stabilization of PR-Set7 resulting from the expression of a

degradation resistant PIP-box mutant version of PR-Set7 leads to re-replication [11]. Similarly, PR-Set7 mutant Drosophila neuroblasts show reduced mitotic and S-phase indices [17], and PR-Set7 RNAi treated S2 cells have an increased S-phase population [18]; however, re-replication resulting from PR-Set7 overexpression has not been observed in the fly.

The ability for mammalian PR-Set7 to regulate replication origin activity is dependent on its catalytic function [11] and the presence of Suv4-20h1/h2 which catalyze the di- and tri-methylation of H4K20 [19]. Consistent with this, H4K20me2 and H4K20me3 may function to stabilize ORC binding via the BAH domain of ORC1 or the WD40 domain of LRWD1/ORCA [19–21]. However, H4K20me2 constitutes more than 80% of total histone H4 [7], which implies that 96% of all nucleosomes will contain at least one histone H4 dimethylated at lysine 20. Similarly, trimethylated H4K20 is, for the most part, limited to heterochromatic regions [22,23]. Together, these results suggest that additional mechanisms must exist to specify origin selection in mammalian genomes. Moreover, it is estimated that mammalian cells have more than 40,000 origins of replication [24], while the influence of PR-Set7 on origin licensing has only been examined at a select few origins [11].

Here, we investigate the function of PR-Set7 and H4K20 methylation in regulating the DNA replication program in *Drosophila* Kc167 cells. Similar to mammalian systems, loss of PR-Set7 perturbs the normal cell cycle progression of Kc167 cells, presumably due to the accumulation of unmethylated H4K20 over the course of multiple cell cycles. We show that PR-Set7 depletion results in DNA damage and activation of an ATR-dependent cell cycle checkpoint, implying a defect in DNA replication. However, the genome-wide pattern of replication origin activation is not impaired; instead, we find that late replication domains are specifically sensitive to downregulation of PR-Set7 and accumulate DNA damage. Together, these data suggest that

PR-Set7 and H4K20me1 are critical for maintaining the integrity of late replicating domains in *Drosophila*.

Results

H4K20 monomethylation is critical for the completion of the subsequent cell cycle

The methylation state (none, mono-, di- or tri-methylation) of H4K20 has been linked to genome stability and cell cycle progression in a number of model systems [[reviewed in 1]. To assess the impact of H4K20 methylation states on cell cycle progression in *Drosophila* Kc167 cells, we manipulated the levels of the H4K20 methyltransferases, PR-Set7 and Suv4-20 (Fig 1A). Depletion of the monomethyltransferase PR-Set7 with double-stranded RNA (dsRNA) mediated RNAi for 72 hours resulted in a 50% reduction of H4K20me1 levels compared with control RNAi. Conversely, depletion of Suv4-20, which catalyzes di- and tri-methylation of H4K20, resulted in a dramatic increase in H4K20me1 levels. Overexpression of a stabilized PIP box mutant of PR-Set7 (F523A and F524A; PR-Set7^{PIPm}) under the control of a copper-inducible metallothionein promoter alone or in combination with depletion of Suv4-20 also resulted in an accumulation of H4K20me1. We also observed a slight increase in H4K20me1 levels relative to control RNAi when simultaneously depleting PR-Set7 and Suv4-20. This slight increase is likely due to differences between the rates of H4K20 monomethylation by PR-Set7 and subsequent dimethylation by Suv4-20.

Cell cycle progression was monitored in parallel by flow cytometry for each of the genetic manipulations described above (Fig 1B). Depletion of PR-Set7 for 72 hours in Kc167 cells resulted in an accumulation of cells with 4N or near 4N DNA content, similar to previous reports in *Drosophila* S2 cells [13]. Interestingly, we also observed an accumulation of cells with 4N

DNA content following the simultaneous depletion of PR-Set7 and Suv4-20 despite the slight increase in H4K20me1. Given that PR-Set7 activity is strictly coupled to the cell cycle and H4K20me1 is a transient epigenetic mark deposited in late S phase and converted to H4K20me2/3 by late mitosis [8,13], we hypothesize that the cell cycle arrest may not be due to the depletion of parental H4K20me1 levels, but rather the accumulation of nascent histone H4.

To test this hypothesis, we examined the kinetics of the cell cycle arrest following PR-Set7 downregulation in synchronized cells (Figs 1C and 1D). Cells were synchronized in G1 by overexpressing the *Drosophila* P27 homolog, Dacapo [25], in the presence of control or PR-Set7 dsRNA. The cells were then released from the Dacapo-induced arrest and allowed to re-enter the cell cycle. In contrast to control cells, cells depleted for PR-Set7 exhibited no increase in H4K20me1 following release from the Dacapo-induced G1 cell cycle arrest (Fig S1). Despite the absence of PR-Set7 and a failure to monomethylate H4K20 during S phase, the cell cycle profile for PR-Set7 RNAi treated cells was indistinguishable from control RNAi treated cells after 24 hours, implying a decoupling of cell cycle progression and H4K20me1 levels. Only after 48 hours did we observe a strong cell cycle arrest. We propose that the monomethylation of nascent H4K20 deposited during S phase is critical for the completion of the next cell cycle.

Global H4K20 monomethylation occurs in late S phase

Cell cycle regulated levels of PR-Set7 start to peak in late S phase with the accumulation of H4K20 monomethylation [8,13]. Recent reports indicate that *Drosophila* PR-Set7 is coupled to the replication fork via an interaction with DNA Pol alpha [13], which may suggest that the accumulation of H4K20 monomethylation in late S phase will be associated with recently replicated sequences. We used cytological and genomic approaches to characterize and monitor the accumulation of H4K20 monomethylation throughout the cell cycle in *Drosophila* Kc167

To more precisely ascertain the patterns of H4K20 monomethylation, we used chromatin immunoprecipitation coupled with high-throughput sequencing (ChIP-seq) to map H4K20 monomethylation at the beginning or end of S phase. Cells were synchronized at the G1/S transition (early S phase) by treatment with hydroxyurea (HU), an inhibitor of nucleotide reductase. To obtain a synchronous population of late S-phase cells, cells were released from the HU arrest and allowed to progress through S phase for six hours (Fig 2C).

H4K20me1 levels are cell cycle regulated and exhibit their lowest levels in early S phase. Despite only detecting background levels of H4K20me1 by western blot in cells arrested in early S phase at the G1/S transition (Fig 2C), we identified discrete peaks of H4K20me1 enrichment

throughout the genome (Fig 2D). These peaks coincided with transcription start sites and regulatory elements as previously noted (Fig 2E; [27]). In late S phase, in addition to enrichment of H4K20me1 at regulatory elements (Fig 2D), we also observed a global increase in H4K20me1 in both genic and intergenic sequences throughout the *Drosophila* genome (Fig 2F). As an alternative approach to isolate cell populations in early or late S phase, we used cell sorting. Chromatin was isolated from these distinct cell populations and the genome-wide distribution of H4K20me1 was determined for early and late S phase. The results were similar to the HU treated samples (Fig S2). Together, these results indicate that monomethylation of H4K20 by PR-Set7 is not limited to replication forks but occurs throughout the genome in late S phase.

Loss of *Drosophila* PR-Set7 induces DNA damage and activates the ATR checkpoint

Loss of PR-Set7 results in an altered cell cycle distribution with an accumulation of cells with greater than 2N DNA (Fig 1B; [13,15]), suggesting the activation of the DNA damage response. Prior experiments investigating the occurrence of PR-Set7-mediated DNA damage by immunostaining of the phosphorylated histone H2A.v (γ-H2A.v, the *Drosophila* homolog of H2AX) led to conflicting results [13,17,18]. The discrepancy in whether or not DNA damage accumulates in PR-Set7 depleted or mutant cells may, in part, result from the specificity of the antibodies used to detect γ-H2A.v. Using a recently developed monoclonal anti-γ-H2A.v antibody [28], we observed an increase in the number and intensity of γ-H2A.v foci when Kc167 cells were depleted of PR-Set7 by RNAi treatment for 72 hours (Figs 3A, S3A and S3B). The observed increase in γ-H2A.v resulting from PR-Set7 depletion was supported by the direct detection of double-strand DNA breaks (DSBs) using a neutral DNA comet assay (Figs 3B and 3C).

The accumulation of DNA damage in the absence of PR-Set7 activity led us to test the role of ATM and ATR in mediating the cell cycle arrest. As previously reported [17], we found that depletion of ATR abrogated the checkpoint and cell cycle arrest due to loss of PR-Set7. In contrast to the co-depletion of ATR and PR-Set7, loss of ATM and PR-Set7 had no impact on the cell cycle arrest (Fig 3D). To confirm the efficacy of ATM RNAi, we tested its ability to reduce the phosphorylation of H2A.v when cells were stressed with methyl methanesulfonate (MMS), a strong DNA damaging agent. While ATR or ATM dsRNA alone did not reduce the levels of γ -H2A.v detected by immunofluorescence, the downregulation of both significantly reduced the H2A.v phosphorylation that occurred in response to MMS (Fig S3C). We propose that PR-Set7 depletion in *Drosophila* Kc167 cells results in aberrant DNA replication which activates the ATR-dependent intra-S checkpoint.

Drosophila H4K20 methylation states do not affect replication origin activity

In mammalian systems, PR-Set7 is thought to regulate pre-RC assembly and origin activation [11], presumably dependent on Suv4-20h1/h2 mediated di- or trimethylation of H4K20 [19]. We first hypothesized that loss of PR-Set7 may activate the ATR-dependent intra-S phase checkpoint due to aberrant (reduced) origin activation. We used BrdU-seq to map the genome-wide origin activation profile [29]. Briefly, we incubated asynchronous cells with dsRNA targeting PR-Set7 or a control sequence for 36 hours before adding HU for 4 hours to inhibit active replication forks. We then added BrdU (in the presence of HU) for another 16 hours. This provided the cells sufficient time to enter the second cell cycle following RNAi treatment so that most cells would be arrested at the second G1/S transition, and only those sequences proximal to early activating origins would incorporate BrdU. As expected, the BrdU incorporation observed in both the control and PR-Set7 depleted cells peaked at ORC binding

We next considered the hypothesis that PR-Set7 depletion might impair late origin activation. To test this hypothesis, we co-depleted PR-Set7 and ATR in the presence of HU and BrdU. ATR depletion and abrogation of the intra-S phase checkpoint permitted the activation of late and/or dormant origins in addition to early origins. However, the origin activation profile upon co-depletion of ATR and PR-Set7 was indistinguishable from the depletion of ATR alone (Figs 4A and 4B). These results suggest that the cell cycle arrest following depletion of PR-Set7 is not due to a defect in origin activation.

Finally, deregulation of PR-Set7 by overexpression results in DNA re-replication and DNA damage in mammalian cells [11]. However, *Drosophila* PR-Set7^{PIPm} overexpression did not lead to re-replication, nor did Suv4-20 RNAi result in any cell cycle defects (Fig 1B). To test if any of these conditions might impact origin activation while keeping the cell cycle progression intact, we performed BrdU-seq in the presence of HU as described above. We detected no significant perturbations in the distribution or activity of replication origins throughout the genome (Fig 4B). We conclude that deregulation of H4K20 methylation does not directly impact replication origin activity in *Drosophila*.

Loss of PR-Set7 accumulates DNA damage in late replicating domains

Since replication origin activity was not impacted by PR-Set7 depletion, we hypothesized that the DNA damage lesions (Fig 3) may occur during replicative stress and result from collapsing replication forks. The presence of DNA breaks marked by γ -H2A. ν provided an opportunity to identify the genomic locations and features associated with DNA damage resulting from the loss

of PR-Set7. We used ChIP-Seq to identify the genome-wide distribution of y-H2A.v. The control and PR-Set7 depleted samples exhibited numerous y-H2A.v peaks (Fig S5), presumably reflecting endogenous replication- and transcription-mediated stress [30]. In order to identify the genomic locations specifically enriched for y-H2A.v by PR-Set7 depletion, we normalized the y-H2A.v signal by calculating the log2 difference in signal between PR-Set7 and control RNAi treated samples (see Materials and Methods). Following normalization, we observed discrete domains of elevated v-H₂A,v throughout the genome (Fig 5A). The vast majority of v-H₂A,v enriched domains were associated with late replicating sequences and were inversely correlated with replication timing data (Pearson correlation = -0.696; Figs 5A and 5B; [31]). We used a Hidden Markov model to define 102 y-H2A.v domains and compared them with the replication timing domains identified in a prior study [31]. We found that 88 of the y-H2A.v domains overlapped with a late replicating domain while 120 of the 166 late replicating domains coincided with a y-H2A.v domain (Table S1). We did not observe hotspots of y-H2A.v within the late domains, but rather found that y-H2A.v spanned the entire late domain region (Fig. 5C). Given that ChIP-seq is a population-based assay averaged over millions of cells, the lack of discrete y-H2A.v 'hotspot' peaks within the late replicating domains implies a spatial pattern of stochastic DNA damage likely resulting from replication fork collapse in the absence of H4K20me1.

Discussion

Depletion of PR-Set7 in *Drosophila* Kc167 cells results in a decrease in H4K20 monomethylation, activation of an ATR-dependent checkpoint, and a cell cycle arrest in late S phase. The activation of the ATR-dependent checkpoint was accompanied by the accumulation of DNA damage. We found that PR-Set7 and H4K20 monomethylation of nascent histones were not required to complete the first S phase, but rather cells entering their second cell cycle failed

H4K20 is the most common histone target for post-translational modification. The vast majority (>80%) of H4K20 exists in the dimethyl state [7]; therefore, every cell cycle the bulk of nascent histone H4 deposited during DNA replication needs to undergo monomethylation by PR-Set7 and subsequent di- or tri- methylation by Suv4-20. The conserved PIP-box mediating the PCNA and CRL4^{Cdt2} dependent degradation of PR-Set7 tightly couples PR-Set7 activity to the cell cycle. PR-Set7 globally monomethylates the nascent pool of H4K20 on lysine 20 in late S phase. Depletion of PR-Set7 in G1 and loss of late S phase specific H4K20 monomethylation had no impact on the progression through S phase in the first cell cycle, indicating that there is not an immediate "just in time" requirement for H4K20me1 to complete S phase or enter into mitosis. DNA damage and cell cycle arrest occur in the next cell cycle with the accumulation of nascent nucleosomes containing unmodified H4K20 throughout the genome (Fig. 6).

DNA damage resulting from loss of PR-Set7 and H4K20 methylation was restricted to late replicating regions of the genome, suggesting a defect in the DNA replication program at the end of S phase. A paucity of activated origins has been shown to lead to DNA damage, fragile sites, and increased mutations in human cells due to the increased distances replication forks must traverse [32–34]. However, unlike in mammalian cells, deregulation of H4K20 methylation did not impair origin activation in *Drosophila*. In mammalian cells, tethering of PR-Set7 to the DNA promotes the locus-specific recruitment of replication factors and pre-RC assembly (presumably

via increased levels of H4K20me2 and ORC stabilization) [11,19,20]. However, given that almost all nucleosomes contain at least one histone H4 methylated at lysine 20, it is difficult to envision how tethering could lead to a greater local concentration of H4K20me2. It remains to be seen if the locus-specific effects observed at select origins in mammalian cells will represent a genome-wide paradigm [11].

Loss of PR-Set7 leads to a cell cycle arrest with nearly 4N DNA content. Depletion of the putative DNA damage checkpoint ATM did not relieve the cell cycle arrest. This may be due to the dispensable role of Drosophila ATM in maintaining the premitotic-checkpoint in response to DNA damage [35,36]. Instead, depletion of ATR activity relieved the checkpoint and cell cycle arrest suggesting that loss of PR-Set7 and H4K20 methylation activate the ATR-dependent intra-S phase checkpoint. The checkpoint is likely activated due to the accumulation of ssDNA and/or DSBs resulting from replication fork stalling and collapse (Reviewed in [37]). We found that the DNA damage resulting from loss of PR-Set7 accumulates in late replicating regions of the Drosophila genome. We also found that the pattern of γ -H2A. ν marked DNA damage was broadly distributed across late replicating domains suggesting a model whereby the accumulation of unmodified H4K20 may render the chromatin recalcitrant to replication fork progression resulting in stochastic replication fork collapse.

Why is *Drosophila* PR-Set7 and H4K20 methylation critical for the integrity of late replicating domains? In contrast to early replicating domains which are enriched for activating chromatin PTMs, late replicating regions of the genome are generally gene-poor and depleted of most chromatin PTMs [29,38]. Thus, H4K20me1 may be a critical epigenetic mark to maintain proper chromatin architecture of late replication domains. Consistent with this hypothesis, H4K20me1 is necessary to ensure proper chromatin compaction [18,39]. Loss of PR-Set7 leads to altered

nuclear morphology; specifically, an enlarged nucleus due to abnormal chromatin decompaction (Fig 3A; [18]). Together, these results suggest that normal chromosome compaction is a determinant of replisome stability and fork progression.

Alternatively, H4K20 methylation may be required for repairing and restarting spontaneous stalled replication forks which are more likely to be enriched in late replication regions (Reviewed in [40]). H4K20me2 has been implicated in interacting with 53BP1 in mammalian cells, which promotes the non-homologous end-joining (NHEJ) pathway to repair DNA DSBs and antagonizes the homologous recombination (HR) pathway factor BRCA1 [41,42]. It is proposed that PR-Set7 deposits de novo monomethylation marks on H4K20 flanking DSBs, which provide substrates for sequential conversion to H4K20me2 by Suv4-20h1 [43,44]. In mammalian U2OS cells, depletion of either an HR factor, Rad51, or an essential replication factor, Cdc45, eliminated the accumulation of y-H2AX induced by PR-Set7 siRNA [14]. These results suggest that PR-Set7 activity may promote NHEJ over HR during S phase. However, co-depletion of PR-Set7 and the *Drosophila* Rad51 or BRCA2 homolog had no impact on reducing the intensity of y-H2A.v by immunofluorescence in our system (Fig S6). Also, loss of Drosophila Suv4-20 had little if any impact on the cell cycle progression of Kc167 cells (Fig. 1B) or animal survival [45]. This may reflect the variability of the replicative stress response pathways across species, or likely a less essential role of H4K20me2 in *Drosophila*, despite it being the most abundant form of H4K20 [7].

PR-Set7 is essential for development. Loss of PR-Set7 in mice leads to embryonic lethality prior to the eight-cell stage [39]; similarly, flies homozygous null for PR-Set7 die as third instar larvae [46]. While it has been assumed that the essential function of the PR-Set7 methyltransferase is to monomethylate H4K20, it remains possible that it could target other critical proteins for

methylation. Indeed, nonhistone targets of PR-Set7's catalytic activity have been identified in mammalian cells, including p53 [47] and PCNA [48], both of which are critical cell cycle regulators.

Recently, McKay and colleagues developed a novel histone replacement system to directly assess the function of specific histone lysine residues in *Drosophila* [49]. Strikingly, more than half of the flies harboring a mutation of histone H4 lysine 20 to alanine (H4K20A), although sick, were able to survive into adulthood. H4K2OA mutant flies also exhibited robust EdU incorporation in germline nurse and somatic follicle cells, which is consistent with our findings that origin activation is not impacted by loss of PR-Set7 activity. Despite the tight cell cycle-coupled regulation of PR-Set7, we found that depletion of PR-Set7 and loss of H4K20 monomethylation did not impair the first S phase but rather impacted the subsequent cell cycles. While we cannot completely rule out the possibility that it may take more than 24 hours for a PR-Set7 methylated non-histone protein to turn over, we suggest that cell cycle arrest in subsequent cell cycles is an epigenetic effect coupled to the accumulation of unmodified H4K20 on the chromatin. What is unclear, is precisely how much H4K20 methylation throughout the genome is required to maintain genomic stability? In the case of the H4K2oA mutant flies, it remains possible that the incorporation of minimal amounts of the replication independent histone H4, His4r, (which was not mutated) may be sufficient to prevent the accumulation of DNA damage during late S phase. Finally, the majority of tissues derived from differentiated cells in the fly are polyploid resulting from developmentally programmed endocycles [50]. These cells have under-replicated regions which frequently correspond with late replicating domains [51] and are de-sensitized to the DNA damage response pathway [52]. Thus, H4K20 methylation may have a minimal impact on cells that are predisposed to fail to replicate and/or activate the DNA damage response in late replicating regions of the genome.

Materials and Methods

Drosophila cell culture

Drosophila Kc167 cells were cultured at 25°C in Schneider's Insect Cell Medium (Invitrogen) supplemented with 10% heat-inactivated fetal bovine serum (Gemini) and 1% penicillin/streptomycin/glutamine (Invitrogen). Dacapo and PR-Set7^{PIPm} were cloned into the pMK33 plasmid under the control of a Cu²+ inducible metallothionein promoter and transfected into Kc167 cells with the Effectene Transfection Reagent (Qiagen). Two days after transfection, cells were selected and maintained with 0.125 μg/ml Hygromycin B (Sigma). To arrest cells in G1, Dacapo was overexpressed with 1 mM copper sulfate for 24 hours. Similarly, PR-Set7^{PIPm} was induced with 0.5 mM copper sulfate. To arrest cells at the G1/S transition, 1 mM hydroxyurea (Sigma) was added for 24 hours (unless otherwise noted).

Double-strand RNA synthesis and RNA interference

Double-strand RNAs (dsRNAs) were generated by *in vitro* transcription (Promega T7 RiboMax) from 400-800 bp PCR products representing the target gene. Target sequences were flanked by T7 promoters on both strands. Primer sequences used are found in Table S2. To perform RNAi, cells were washed and diluted in serum-free medium to a concentration of 2×10^6 cells/ml, and $15~\mu g$ dsRNA per 10^6 cells was added and incubated for one hour before diluting cells to 1×10^6 cells/ml with 2x medium. The cells were then incubated at $25^{\circ}C$ for 24-72 hours.

Western blot analysis

Primary antibodies used for western blot analysis: rabbit anti-PR-Set7 (gift from Danny Reinberg) at 1:500, rabbit anti-H4K20me1 (Abcam) at 1:1000, rabbit anti-histone H3 (Abcam)

at 1:2000. Alexa Fluor 680 goat anti-rabbit IgG (Invitrogen) was used at 1:10000 as a secondary antibody. Immunoblots were scanned using a LiCor infrared system and analyzed with Odyssey software.

Flow cytometry and cell sorting

For cell cycle analysis by flow cytometry, cells were harvested by centrifugation, washed in 1x cold PBS, resuspended in 500 μ l PBS and fixed with 5 ml ice-cold 100% ethanol at 4°C overnight. The cells were then washed in 1x cold PBS, resuspended in PBS with 0.3 mg/ml RNase A and 15 μ g/ml propidium iodide, and incubated for 30 minutes at room temperature. Flow cytometry was performed on BD FACscan analyzer machine and 30000 cells were recorded for each sample. Histograms of cell cycle distribution were generated with the R bioconductor flowCore package [53].

To sort cells into G1/S and late S/G2 populations, asynchronous cells were fixed with 1% formaldehyde for 10 min and quenched by 125 mM glycine for 10 min. Then 0.5 μ g/ml Hoechst 33342 (Invitrogen) was used to stain DNA before the cells were sorted into G1/S and late S/G2 populations based on DNA content using BD DiVa sorter.

Immunofluorescence

For EdU pulse labeling, cells were incubated with 10 μ M EdU for 30 minutes before harvest, fixed with 4% paraformaldehyde in PBS for 10 minutes, permeabilized with 0.5% Triton X-100 in PBS for 3 minutes, and labeled with the Click-iT EdU imaging kit (Invitrogen). Cells were then blocked with 3% BSA in PBS with 0.1% Triton X-100 (PBS-T), stained with rabbit anti-H4K20me1 (Abcam) at a dilution of 1:1000. Alexa Fluor 568 goat anti-rabbit (Invitrogen)

was used at 1:500 as the secondary antibody. For detecting γ-H2A.v, cells were fixed, permeabilized, and blocked as above. Cells were stained with mouse monoclonal anti-γ-H2A.v (DSHB UNC93-5.2.1-s) at 1:500 and Alexa Fluor 488 goat anti-mouse (Invitrogen) at 1:500. All slides were mounted with VECTASHIELD Mounting Medium with DAPI (Vector Laboratories) and observed with Zeiss Axio Imager widefield fluorescence microscope with 40x objective.

DNA comet assay

The neutral DNA comet assay was performed using the CometAssay Kit (Trevigen). DNA was stained with the SYTOX Green nucleic acid stain (Invitrogen, dilute to 1:10000 in TE) and images were acquired with an Zeiss Axio Imager widefield fluorescence microscope. The comet tail moment was quantified using the the ImageJ plugin OpenComet [54].

Chromatin immunoprecipitation

Chromatin immunoprecipitation were performed as previously described [55]. H4K20me1 and γ -H2A.v ChIP experiments were performed using rabbit anti-H4K20me1 (Abcam, validated for ChIP-seq [56]) and mouse anti- γ -H2A.v (DSHB UNC93-5.2.1-s) antibodies, respectively. The γ -H2A.v ChIP experiment was performed in duplicate. The H4K20me1 experiment was also performed in duplicate by using two independent approaches (HU exposure and cell sorting) to obtain synchronous cell populations.

Genome-wide analysis of early origin activation

A 10 ml culture of asynchronous cells at a concentration of 1×10^6 cells/ml was treated with dsRNA or copper for 36 hours and then 1 mM HU was added and the cells were incubated for 4 hours to stall any active replication forks. Subsequently, 10 μ M BrdU was added to the culture

and the cells were incubated for another 16 hours before harvesting. Genomic DNA extraction and BrdU immunoprecipitation were performed as previously described [57].

Sequencing library preparation

Illumina sequencing libraries were generated from the DNA enriched by ChIP or BrdU immunoprecipitation using Illumina's TruSeq protocols.

Sequencing read alignment

Sequenced reads were aligned to the dm3 release 5.12 of the *Drosophila melanogaster* genome assembly (downloaded from the UCSC Genome Browser, and available at ftp://hgdownload.cse.ucsc.edu/goldenPath/dm3/bigZips/chromFa.tar.gz), using the software package Bowtie (version 0.12.7) [58]. The following Bowtie parameters were used: -n 2 -l 30 -m 1 --best --strata -y.

Data accession and genomic data analysis

All genomic data are publicly available at the NCBI GEO data repository with the accession number GSE73668.

For H4K20me1 ChIP-seq and BrdU-seq experiments, RPKM (reads per kilobase, per million mappable reads) was calculated over bins of 10 kb stepping every 1 kb. For the γ -H2A.v datasets, RPKM was determined over nonoverlapping bins of 5 kb. The list of gene regions was downloaded from UCSC Table Browser (Assembly BDGP R5/dm3) using FlyBase Genes track.

The H4K20me1 ChIP signal was normalized between early and late S-phase samples based on the relative enrichment in the top 5% of detected peaks [59].

For γ -H2A.v ChIP-seq analysis, we calculated the log2 difference of ChIP signal: log2 difference = log2(PR-Set7 RNAi RPKM + 1) - log2(control RNAi RPKM + 1). A positive value indicates increased γ -H2A.v upon PR-Set7 RNAi treatment. To identify discrete γ -H2A.v domains, we trained a two-state hidden Markov model (HMM, representing positive or negative γ -H2A.v domains), with each state emitting a univariate normal distribution corresponding to the log2 difference values. To generate the metaplot of replication timing over late replicating domains, each late domain was divided into 50 bins. Twenty-five flanking upstream and downstream bins, equivalent in width to the late domain bin, were also included. The γ -H2A.v log2 differences and replication timing values within each bin were then calculated and the respective means across all domains were plotted to generate a metadomain.

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Figures

Fig 1. H4K20me1 is necessary for cell cycle progression. (A-B) PR-Set7 downregulation arrests cell cycle progression. Asynchronous cells were depleted for the histone methyltransferases, PR-Set7 or Suv4-20, by RNAi, or alternatively, a stabilized mutant of PR-Set7, PR-Set7^{PIPm}, was overexpressed alone or in combination with Suv4-20 depletion for 72 hours. Cells were harvested for (A) western blot analysis for relative levels of PR-Set7 and H4K20me1 (histone H3 as the loading control), and for (B) cell cycle analysis by flow cytometry. (C-D) Kinetics of PR-Set7 RNAi induced cell cycle arrest. Dacapo was overexpressed for 24 hours with control or PR-Set7 dsRNA. The cells were then released from Dacapo overexpression and collected at 24 hours and 48 hours (C) western blot analysis of H4K20me1 (histone H3 as the loading control) and (D) flow cytometry analysis of the cell cycle profile.

Fig 2. H4K20me1 levels are globally increased throughout the genome in late S phase. (A) H4K20me1 is restricted to cells in late S phase and G2. Asynchronous cells were
pulse labeled with EdU for 30 minutes to detect cells in early/mid and late S phase. EdU and
H4K20me1 were detected by immunofluorescence microscopy and the nuclei were stained with
DAPI. (B) Quantification of H4K20me1 immunofluorescence intensity in early and late S phase
(n>20). (C) H4K20me1 is established in late S-phase cells released (6 hours) from an HU arrest.
(D) Enrichment of H4K20me1 (RPKM) over a representative 4 MB region of chromosome 2L for
cells arrested at the G1/S transition with HU (blue) or cells in late S phase following release from
HU for 6 hours (orange). (E) H4K20me1 is enriched at transcription start sites and gene
bodies. Aggregate plots of H4K20me1 enrichment in HU arrested (blue) and HU released
(orange) cells relative to transcription start sites. (F) H4K20me1 levels increase in both genic

and intergenic regions during late S phase. Boxplots representing the distribution of H4K20me1 RPKM within 13428 genic and 11430 intergenic regions (p<2.2×10⁻¹⁶).

Fig 3. PR-Set7 maintains genome integrity. (A) Depletion of PR-Set7 by RNAi treatment for 72 hours leads to elevated γ-H2A.v staining (cyan). Nuclear DNA is stained with DAPI (magenta). (B) Depletion of PR-Set7 results in DNA double-strand breaks as determined by a neutral DNA comet assay. Treatment with 0.03% MMS serves as a positive control. (C) Quantification of tail moments (tail length × percentage of DNA in tail) from the neutral DNA comet assay (n>45 for each condition). (D) Flow cytometry analysis of the cell cycle after RNAi treatment against control, PR-Set7, ATM/PR-Set7 and ATR/PR-Set7 for 72 hours..

Fig 4. De-regulation of H4K20 methylation does not impact the activation of replication origins. (A) Genome-wide early origin activity was mapped by BrdU-Seq for control and PR-Set7 depleted cells. Late origin activity was mapped by BrdU-Seq for cells depleted of ATR and ATR/PR-Set7. BrdU enrichment (RPKM) is depicted for a representative 2.5 MB window of chromosome 2L. (B) Heatmap depicting the pairwise Pearson correlations for origin activity across the different experimental conditions.

Fig 5. Loss of PR-Set7 leads to the accumulation of DNA damage in late replicating regions of the *Drosophila* genome. (A) The DNA damage marker, γ-H2A.v, accumulates in late replicating regions of the genome. The genome-wide distribution of γ-H2A.v was determined for control and PR-Set7 depleted cells by ChIP-Seq (Fig S5). The log2 difference in γ-H2A.v enrichment between PR-Set7 and control RNAi treated cells (blue) is depicted for a 5 Mb window of chromosome 2L. A replication timing profile (red) for chromosome 2L [31] is overlaid on the plot. (B) γ-H2A.v is inversely correlated with the time of replication in S-phase.

Each data point represents a 5 kb bin and the Pearson correlation between replication timing and the log2 difference in γ -H2A.v between PR-Set7 depleted and control cells was calculated. (C) γ -H2A.v is enriched throughout late replicating domains. Meta-domain analysis of γ -H2A.v enrichment in PR-Set7 depleted cells relative to control cells (blue) and replication timing values (orange) over late replicating domains. Each late replicating domain was divided into 50 bins and the average γ -H2A.v log2 difference between PR-Set7 depleted and control cells as well as replication timing values were plotted.

Fig 6. Model for the maintenance of *Drosophila* genome integrity by PR-Set7 in late replicating domains. The majority of parental histone H4 (orange) exists in the dimethylated state in G1. *Left branch (wild type)*: PR-Set7 is absent in early S phase and nucleosomes containing nascent histone H4 unmethylated at H4K20 accumulate on newly synthesized DNA. PR-Set7 levels and activity are restored in late S phase and there is a global conversion of nascent nucleosomes from H4K20me0 to H4K20me1. Following S phase the monomethylated H4K20 is converted to H4K20me2. *Right branch (absence of PR-Set7)*: In the absence of PR-Set7, the first S phase is completed and unmethylated nascent histone H4 (H4K20me0) is accumulated throughout the genome. In the subsequent S phase, early replicating domains are able to be copied; however, during late S phase there is an accumulation of DNA damage, activation of an ATR-dependent intra-S phase dependent checkpoint and cell cycle arrest. Presumably the presence of H4K20me0 in late replicating but not early replicating domains leads to replication fork stalling and collapse suggesting a critical role for PR-Set7 in maintaining the integrity of late replicating domains.

Fig S1. PR-Set7 depletion is sufficient to prevent the accumulation of H4K20me1 upon release from a Dacapo-induced G1 cell cycle arrest. (A) Control and PR-Set7

depleted cells enter late S/G2 within 8 hours following release from a Dacapo-induced G1 arrest. (B) RNAi-mediated depletion of PR-Set7 is sufficient to block the monomethylation of H4K20 in late S/G2 following release from the Dacapo induced cell cycle arrest. Immunofluorescence of H4K20me1 (red) in control and PR-Set7 depleted cells 8 hours following release into S phase. DNA is stained with DAPI (blue). (C) Boxplots resulting from the quantification of H4K20me1 intensity for each condition (n>180).

Fig S2. H4K20me1 levels are globally increased throughout the genome in late S phase. (A) Cell sorting. Hoechst 33342 stained cells were sorted into G1/S and late S/G2 populations based on DNA content. (B) Enrichment of H4K20me1 (RPKM) over a representative 4 MB region of chromosome 2L for cells enriched in G1/early S (blue) or late S/G2 (orange) of the cell cycle by cell sorting. (C) H4K20me1 is enriched at transcription start sites and gene bodies. Aggregate plots of H4K20me1 enrichment in G1/early S (blue) and late S/G2 (orange) cells relative to transcription start sites. (D) H4K20me1 levels increase in both genic and intergenic regions during late S phase. Boxplots representing the distribution of H4K20me1 RPKM within 13428 genic and 11430 intergenic regions (p<2.2×10⁻¹⁶).

Fig S3. PR-Set7 depletion leads to increased H2A.v phosphorylation independent of ATM. (A-B) Quantification of γ-H2A.v signal upon PR-Set7 RNAi treatment. (A) Average number of γ-H2A.v foci per cell from three independent experiments (p = 0.039, t-test). Each error bar is two SDs. (B) Boxplot of the immunofluorescence intensity of γ-H2A.v per cell from one representative experiment. (C) Verification of ATM downregulation by ATM dsRNA treatment. The cells were incubated with control, ATR, or ATM dsRNA for 48 hours before 0.05% MMS was added and incubated for 2 hours (labeled as "MMS +"). The intensity of H2A.v

phosphorylation (γ -H2A.v) in response to MMS treatment was measured by immunofluorescence microscopy and depicted with boxplots (n > 250).

Fig S4. BrdU incorporation in HU arrested cells is enriched at ORC binding sites.

Heatmap of BrdU incorporation (measured as RPKM) in relation to 5159 ORC binding sites [29]. Below the dashed line are the 1677 ORC sites at early origins and above the dashed line are ORC sites at late origins. The center of ORC peak is labeled as position o.

Fig S5. Genome-wide distribution of γ -H2A.v for control and PR-Set7 depleted cells. ChIP-seq analysis of the genome-wide distribution of γ -H2A.v for control and PR-Set7 depleted cells (RPKM).

Fig S6. Disruption of the homologous recombination (HR) pathway does not reduce DNA damage resulting from loss of PR-Set7. Cells were treated with control or PR-Set7 RNAi for 72 hours before harvest and immunofluorescence staining of γ -H2A.v. The immunofluorescence intensity of γ -H2A.v under each condition was depicted with boxplots (n > 250).

Table S1. List of the chromosomal positions of late replicating domains and γ-H2A.v domains identified by a hidden Markov model.

Table S2. List of primer sequences used for generating dsRNAs.

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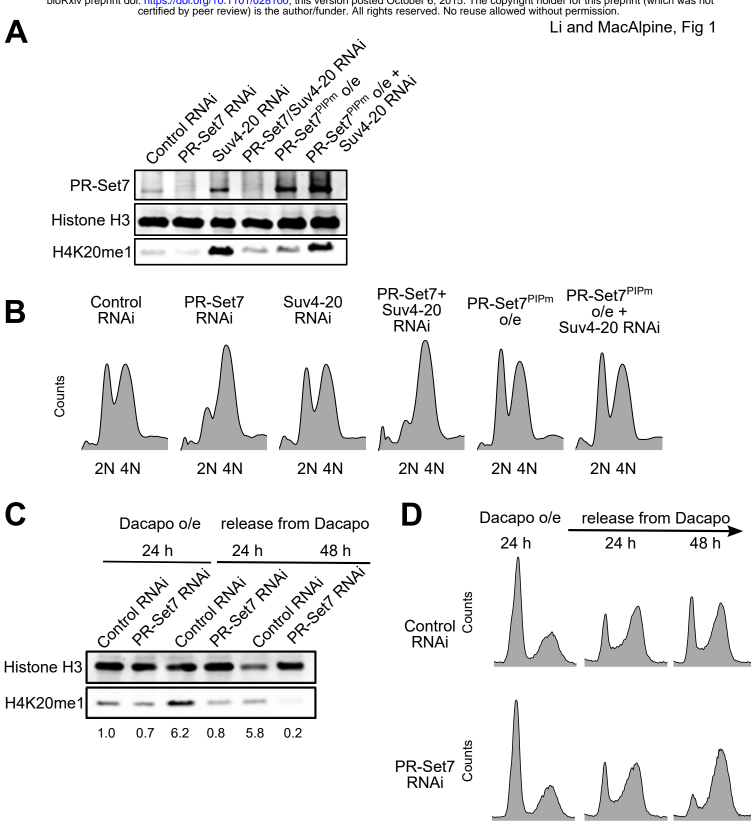
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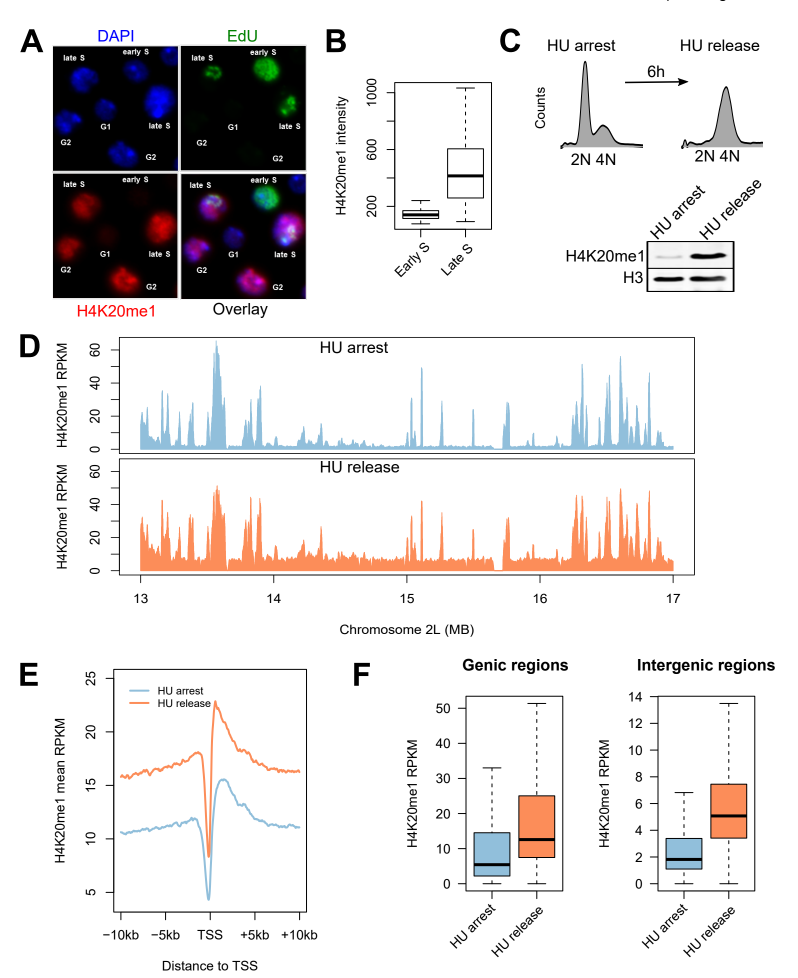
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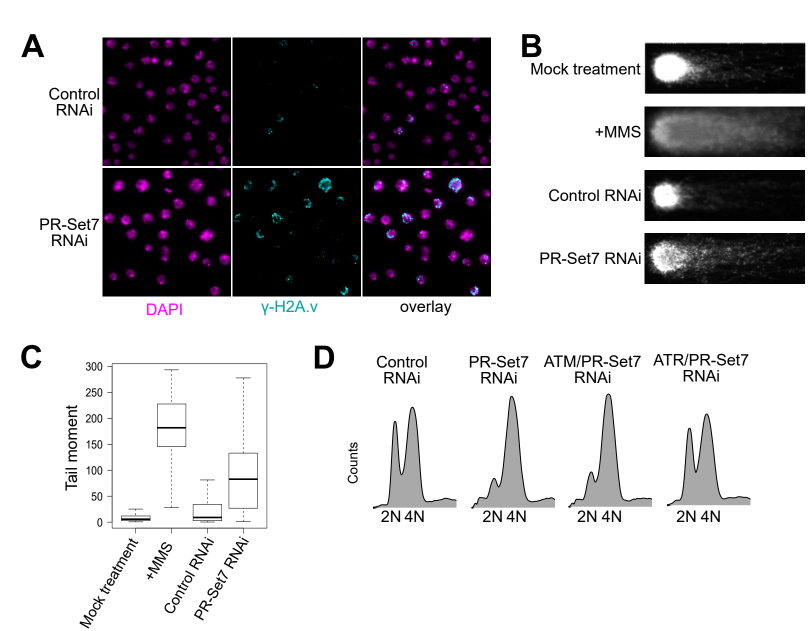
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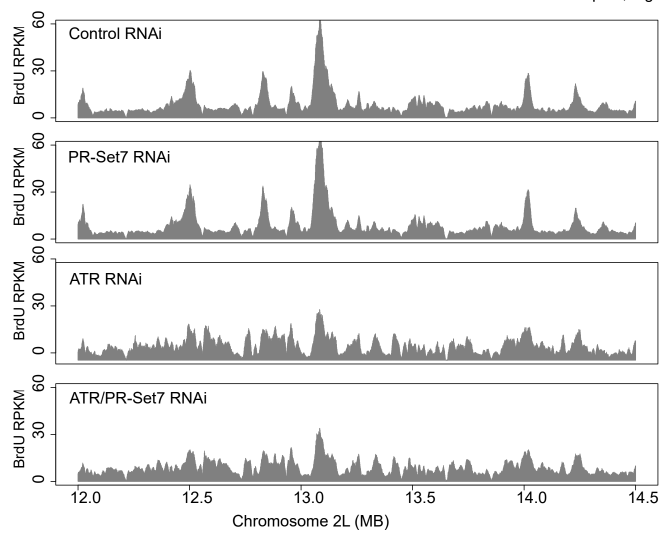
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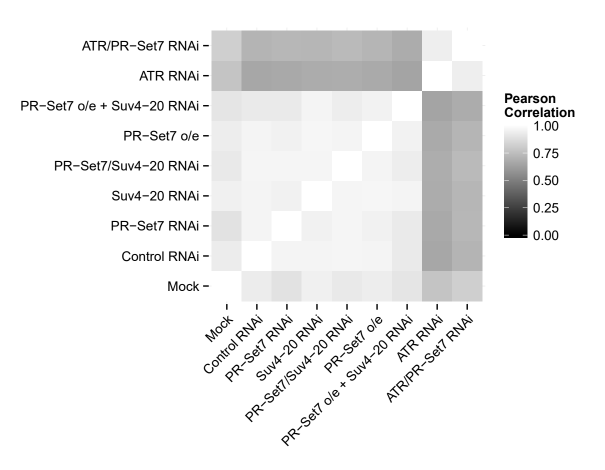


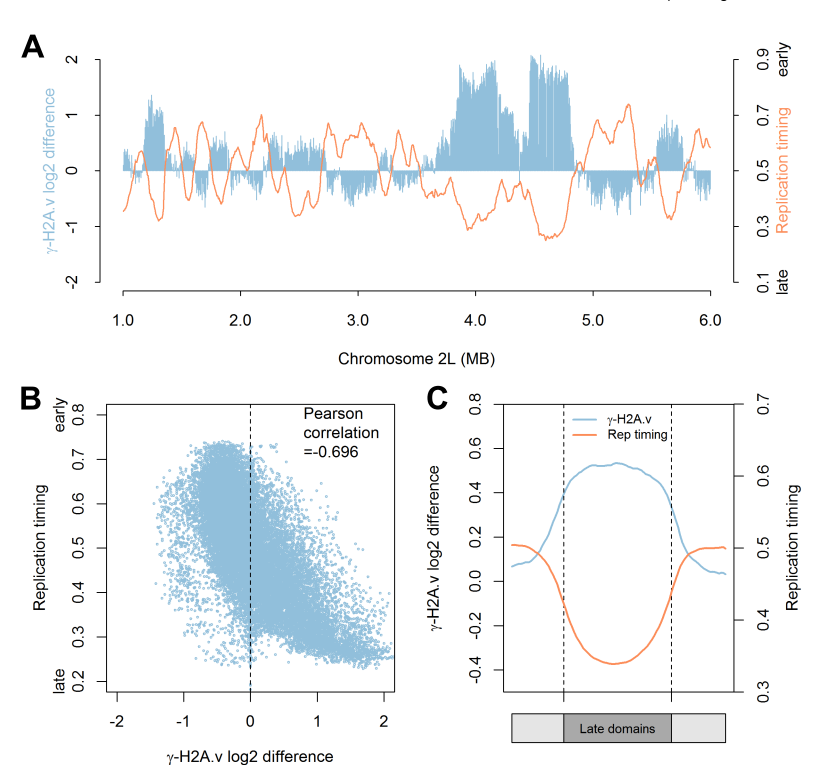


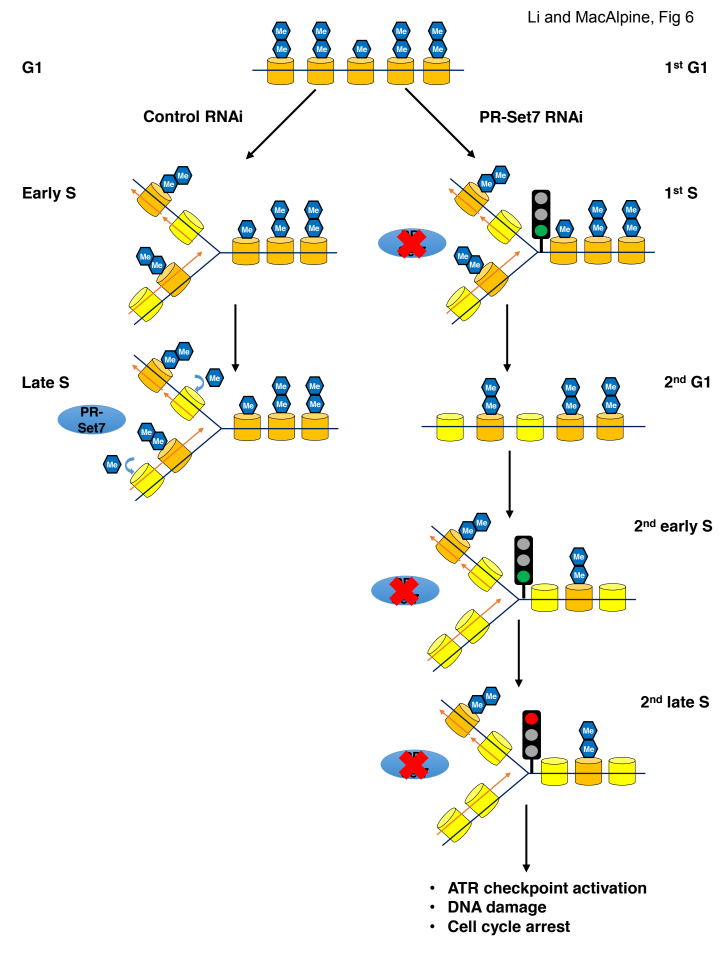




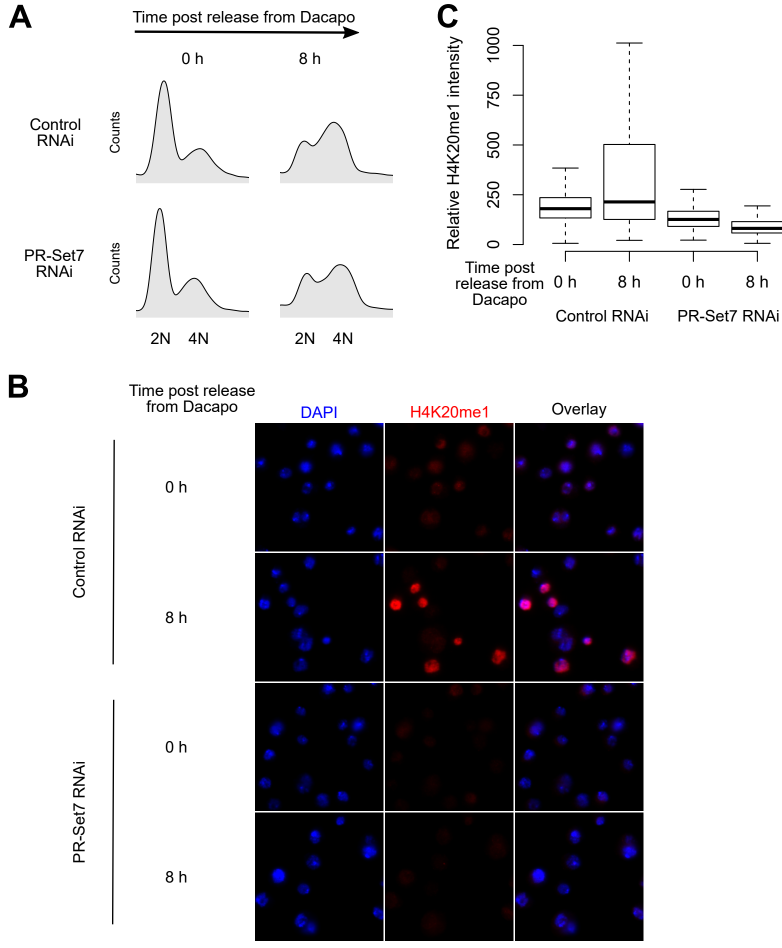


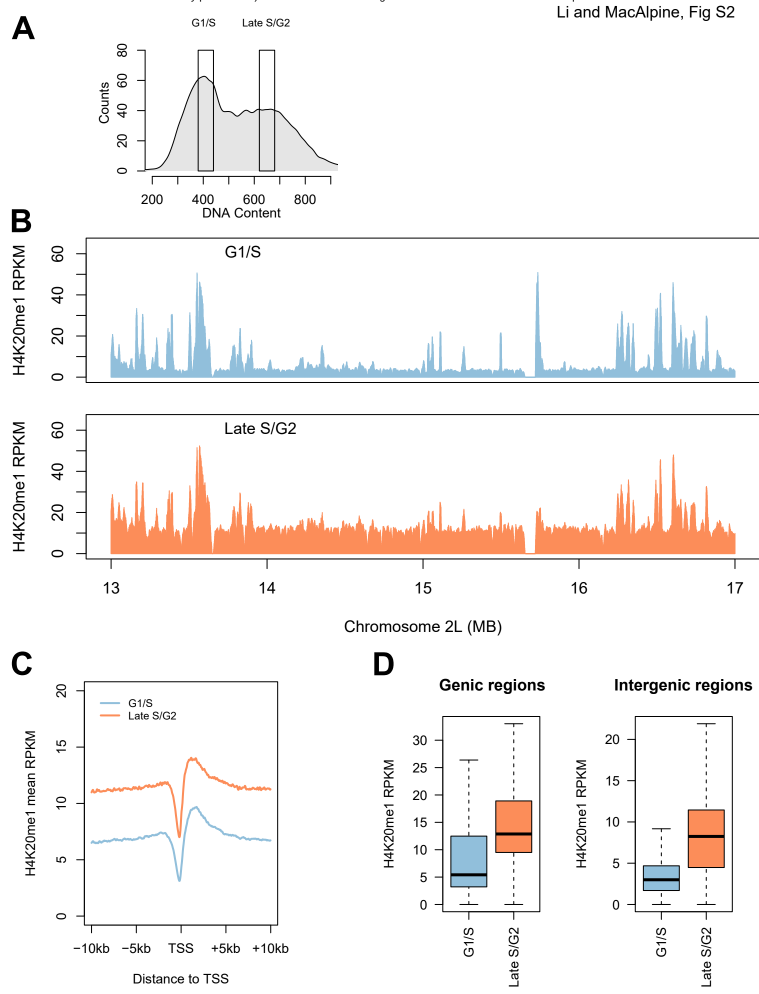




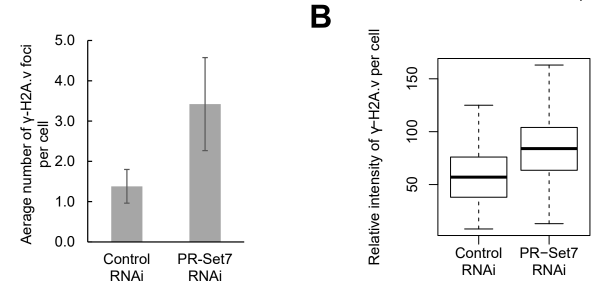


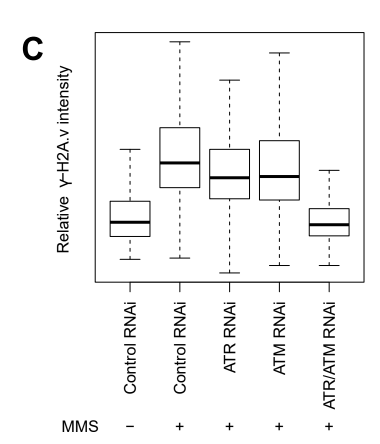


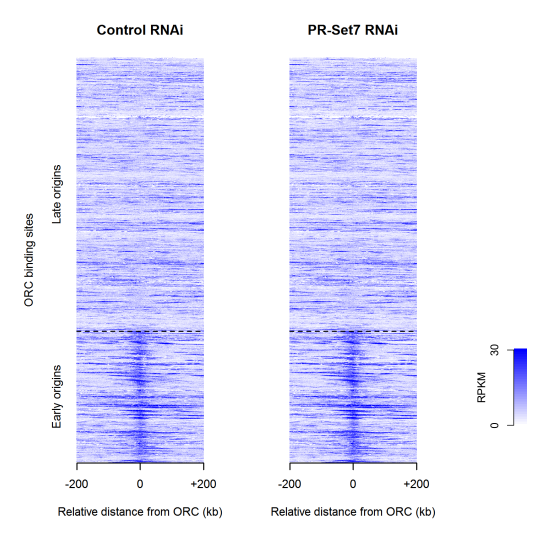




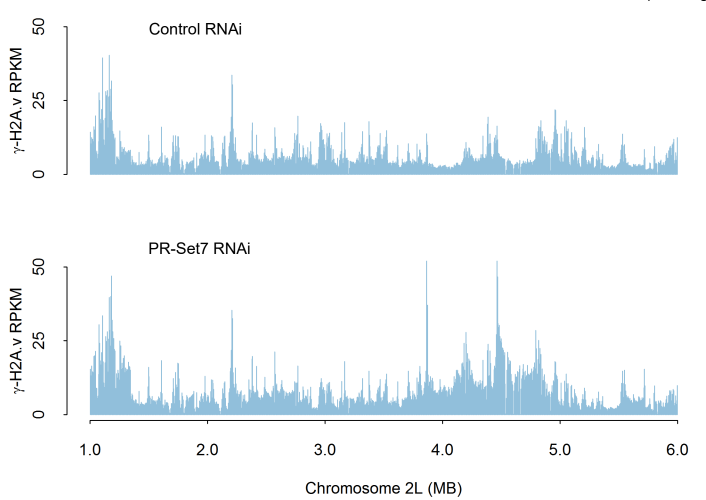
Li and MacAlpine, Fig S3







Li and MacAlpine, Fig S5



Li and MacAlpine, Fig S6

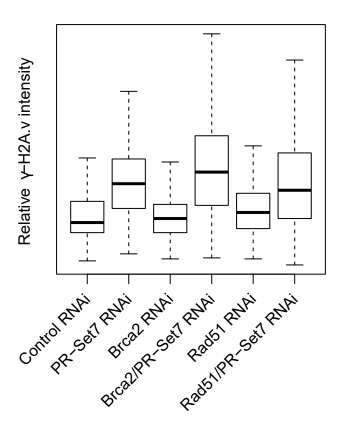


Table S1. Chromosomal positions of late replicating domains and γ H2A.v domains identified by HMM.

Late replicating domains			γ-H2A.v domains			
	Chr	Start	End	Chr	Start	End
1	2L	627002	813000	2L	580000	810000
2	2L	1005002	1048000	2L	900000	1080000
3	2L	1258002	1350000	2L	1170000	1590000
4	2L	2406002	2648000	2L	1730000	1910000
5	2L	3839002	4200000	2L	2220000	2730000
6	2L	4268002	4326000	2L	3620000	4880000
7	2L	4457002	4793000	2L	5520000	5850000
8	2L	6148002	6294000	2L	6100000	6340000
9	2L	8806002	8864000	2L	7500000	7800000
10	2L	10069002	10138000	2L	10510000	10950000
11	2L	10601002	10842000	2L	11240000	12000000
12	2L	11318002	11787000	2L	12100000	12440000
13	2L	11831002	12002000	2L	13900000	16270000
14	2L	12135002	12362000	2L	16850000	18540000
15	2L	14087002	14178000	2L	19190000	19360000
16	2L	14419002	14486000	2L	19570000	19750000
17	2L	14540002	15011000	2L	19820000	20070000
18	2L	15090002	15239000	2L	20440000	20650000
19	2L	15281002	15654000	2L	21750000	22070000
20	2L	15779002	16221000	2R	2940000	3130000
21	2L	16962002	17339000	2R	4610000	4780000
22	2L	17517002	18105000	2R	9450000	9710000
23	2L	18181002	18409000	2R	11270000	11410000
24	2L	19835002	20110000	2R	12230000	12430000
25	2L	20223002	20372000	2R	13030000	13280000
26	2L	20488002	20573000	2R	14730000	15040000
27	2L	21795002	22112000	2R	15360000	16150000
28	2R	1683002	1791000	2R	16210000	16790000
29	2R	2396002	2454000	2R	17630000	17970000
30	2R	2957002	3086000	2R	18570000	18700000
31	2R	4185002	4322000	2R	18820000	19260000
32	2R	4622002	4799000	3L	1870000	2390000
33	2R	4837002	4972000	3L	2840000	3050000
34	2R	5533002	5724000	3L	3600000	3820000
35	2R	6187002	6296000	3L	4410000	5750000
36	2R	6844002	6899000	3L	5880000	6120000
37	2R	7357002	7423000	3L	6210000	6920000
38	2R	7605002	7723000	3L	7010000	7130000
39	2R	7953002	8008000	3L	7390000	7730000
40	2R	9524002	9679000	3L	9130000	9340000
41	2R	10534002	10641000	3L	9850000	10640000
42	2R	10910002	10965000	3L	11790000	12040000

43	2R	11280002	11369000	3L	13030000	13260000
44	2R	11457002	11670000	3L	13360000	13920000
45	2R	12245002	12425000	3L	14180000	14480000
46	2R	13828002	13889000	3L	14780000	14980000
47	2R	14057002	14157000	3L	15300000	15490000
48	2R	14750002	15022000	3L	15690000	15950000
49	2R	15414002	15580000	3L	16230000	16350000
50	2R	15667002	16094000	3L	17020000	17430000
51	2R	16244002	16491000	3L	18100000	18670000
52	2R	16552002	16869000	3L	19080000	19390000
53	2R	17696002	17930000	3L	20530000	20710000
54	2R	18120002	18181000	3L	21670000	21820000
55	2R	18710002	18761000	3L	21940000	22260000
56	2R	18851002	18929000	3L	22400000	22720000
	2R	18972002		3R	1610000	2200000
57 58	2R		19237000			
<u>58</u>	2R 2R	19322002	19435000	3R	2270000	2580000
59		20278002	20386000	3R	3030000	3740000
60	3L	1981002	2147000	3R	4170000	4360000
61	3L	2203002	2350000	3R	6700000	7030000
62	3L	2686002	2731000	3R	7830000	8070000
63	3L	2862002	3064000	3R	8540000	8820000
64	3L	3640002	3760000	3R	8860000	9150000
65	3L	3955002	4008000	3R	9210000	9400000
66	3L	4720002	5094000	3R	10710000	10920000
67	3L	5381002	5569000	3R	12940000	13440000
68	3L	5838002	5887000	3R	13550000	13880000
69	3L	5947002	6068000	3R	14500000	15590000
70	3L	6293002	6878000	3R	15700000	16110000
71	3L	7017002	7113000	3R	16140000	16370000
72	3L	7156002	7234000	3R	17890000	18200000
73	3L	7446002	7716000	3R	18610000	18870000
74	3L	9204002	9293000	3R	19170000	19550000
75	3L	9921002	10203000	3R	21120000	21340000
76	3L	10508002	10582000	3R	21970000	22630000
77	3L	11835002	12045000	3R	23100000	24630000
78	3L	12586002	12663000	3R	24750000	24860000
79	3L	13060002	13187000	3R	25100000	25480000
80	3L	13573002	13799000	3R	26390000	26570000
81	3L	14310002	14370000	3R	26850000	27030000
82	3L	14846002	14962000	3R	27270000	27410000
83	3L	15358002	15418000	3R	27630000	27770000
84	3L	15740002	15786000	X	2660000	2910000
85	3L	15851002	15908000	X	3070000	3180000
86	3L	16247002	16330000	X	4590000	5310000
87	3L	17087002	17437000	X	6250000	6430000
88	3L	18201002	18584000	X	7210000	7410000
89	3L	19166002	19226000	X	9240000	9390000
			-		•	

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90	3L	19294002	19580000	X	9580000	10130000
91	3L	20584002	20758000	X	10770000	11130000
92	3L	20862002	20970000	X	11910000	12400000
93	3L	21056002	21125000	X	12670000	13020000
94	3L	21644002	21805000	X	13310000	13470000
95	3L	21977002	22246000	X	13880000	14920000
96	3L	22453002	22702000	X	15020000	15190000
97	3L	24280002	24398000	X	17860000	18140000
98	3R	1667002	2164000	X	19800000	20280000
99	3R	2332002	2586000	X	20370000	20910000
100	3R	2650002	2839000	X	21320000	21430000
101	3R	3059002	3300000	X	21620000	21700000
102	3R	3400002	3594000	X	21910000	22440000
103	3R	3635002	3710000			
104	3R	6260002	6576000			
105	3R	6732002	6982000			
106	3R	7107002	7346000			
107	3R	7923002	8153000			
108	3R	8361002	8424000			
109	3R	8600002	8800000			
110	3R	8918002	9068000			
111	3R	9207002	9408000			
112	3R	9667002	9767000			
113	3R	10755002	10937000			
114	3R	11566002	11609000			
115	3R	12317002	12449000			
116	3R	12491002	12785000			
117	3R	12983002	13191000			
118	3R	13271002	13337000			
119	3R	13648002	13966000			
120	3R	15055002	15413000			
121	3R	15867002	16066000			
122	3R	16176002	16367000			
123	3R	17910002	18226000			
124	3R	18634002	18819000			
125	3R	19224002	19471000			
126	3R	20204002	20304000			
127	3R	20776002	20826000			
128	3R	20971002	21039000			
129	3R	21187002	21263000			
130	3R	22132002	22244000			
131	3R	22390002	22522000			
132	3R	23177002	23375000			
133	3R	23428002	23735000			
134	3R	23900002	24048000			
135	3R	24195002	24338000			
136	3R	24481002	24610000			
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137	3R	24720002	24828000		
138	3R	25210002	25529000		
139	3R	26316002	26505000		
140	3R	26910002	27013000		
141	X	2747002	2791000		
142	X	3910002	3981000		
143	X	4301002	4409000		
144	X	4652002	4817000		
145	X	4892002	5170000		
146	X	5900002	6067000		
147	X	7284002	7375000		
148	X	8190002	8255000		
149	X	9273002	9387000		
150	X	9630002	10122000		
151	X	10768002	10955000		
152	X	11994002	12289000		
153	X	13375002	13448000		
154	X	13954002	14078000		
155	X	14204002	14532000		
156	X	14588002	14654000		
157	X	15058002	15157000		
158	X	17172002	17370000		
159	X	17811002	17949000		
160	X	18028002	18144000		
161	X	18859002	18915000		
162	X	19852002	20037000		
163	X	20101002	20249000		
164	X	20423002	20877000		
165	X	21335002	21456000		
166	X	22028002	22211000	 	

Table S2. Primer sequences used for generating dsRNAs.

	1 0	
dsRNA Target	Forward Primer	Reverse Primer
PR-Set7	CACAATGATGGCACACAACA	TTCGAAGCTCCGATTGATCT
Suv4-20	CTACGGCGAGGACTTCTTTG	CTTTTCGAGCAAAGAGACGG
ATR	CGACATGGACACAAAACTGG	TTTGGGTAAGCACTTGGACC
ATM	GCGTTCTGCTGGAAGATG	GCTCATCCAAACTAGCGTAA
Rad51	CTCGCACCTTCTATCAAATG	CATCGGCCAGGCGTTG
Brca2	CGACGAAATGGCCAAAATAT	CCTGCCTTGCTTCGGTT
Control (pUC119)	GCGCGGAACCCCTATTTGTTTA	GGAAGCTAGAGTAAGTAGTTCGC