The Histone H4 Lysine 20 Monomethyl Mark, Set by PR-Set7 and Stabilized by L(3)mbt, Is Necessary for Proper Interphase Chromatin Organization

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The Histone H4 Lysine 20 Monomethyl Mark, Set by PR-Set7 and Stabilized by L(3)mbt, Is Necessary for Proper Interphase Chromatin Organization

Ayako Sakaguchi1, Eric Joyce4, Tsutomu Aoki3, Paul Schedl3, Ruth Steward1,2

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Abstract

Drosophila PR-Set7 or SET8 is a histone methyltransferase that specifically monomethylates histone H4 lysine 20 (H4K20). L(3)MBT has been identified as a reader of methylated H4K20. It contains several conserved domains including three MBT repeats binding mono- and dimethylated H4K20 peptides. We find that the depletion of PR-Set7 blocks de novo H4K20me1 resulting in the immediate activation of the DNA damage checkpoint; an increase in the size of interphase nuclei, and drastic reduction of cell viability. L(3)mbt on the other hand stabilizes the monomethyl mark, as L(3)mbt-depleted S2 cells show a reduction of more than 60% of bulk monomethylated H4K20 (H4K20me1) while viability is barely affected. Ploidy and basic chromatin structure show only small changes in PR-Set7-depleted cells, but higher order interphase chromatin organization is significantly affected presumably resulting in the activation of the DNA damage checkpoint. In the absence of any other known functions of PR-Set7, the setting of the de novo monomethyl mark appears essential for cell viability in the presence or absence of the DNA damage checkpoint, but once newly assembled chromatin is established the monomethyl mark, protected by L(3)mbt, is dispensable.

Introduction

Dynamic changes in chromatin structure are directly influenced by the post-translational modifications of the N-terminal histone tails [1,2]. Specific amino acids within the tails are modified by phosphorylation, ubiquitination, ADP ribosylation, acetylation, and methylation [3]. It has been proposed that distinct histone modifications, on one or more tails, act sequentially or in combination to form a “histone code” that is read by other proteins to bring about distinct events [4,5].

H4K20 can be mono-, di-, or trimethylated. Drosophila PR-Set7 (also known as SET8) is a histone methyltransferase (HMT) that specifically monomethylates H4K20 [6,7,8,9], while di- and trimethylation of the same lysine is controlled by Hmt4-20, also known as Suv4-20 [10,11]. In vertebrate tissue culture cells, expression of PR-Set7 increases during S phase and peaks at mitosis [12]. The changing levels of PR-Set7 are controlled by its interaction with PCNA which stimulate PR-Set7 ubiquitylation and proteasome-dependent degradation [13,14,15].

Detailed analysis of the PR-Set7 complete loss-of-function phenotype in flies revealed that in neuroblasts the number of both mitotic and S phase cells is reduced, indicating that the cells stop dividing. This analysis also showed that the progression through early mitosis is delayed and that cyclin B protein is strongly reduced [9]. The abnormalities of mitotic progression and the cyclin B protein level were rescued when the DNA damage checkpoint was abolished, indicating that the DNA damage checkpoint is activated when PR-Set7 activity is compromised. However, though the DNA damage checkpoint is activated, there is no evidence of DNA breakage. On the other hand, PR-Set7 does show defects in chromosome condensation.

Therefore, we proposed that H4K20me1 is involved in the maintenance of proper higher order structure of chromatin. Our results were confirmed in human tissue culture cells; when PR-Set7 was depleted in these cells, the DNA damage checkpoint is also activated. However, in contrast to Drosophila, double strand breaks are detected [16,17]. In these cells, the depletion of PR-Set7 causes S phase delay and alterations in replication fork velocity and origin firing, suggesting that PR-Set7 is involved in S phase progression [16,17].

A vertebrate PR-Set7 interactor, L3MBTL1, has been identified and PR-Set7 is proposed to enhance its transcriptional repression [18]. L(3)MBT proteins contain several conserved domains, including three MBT repeats [19] that have been shown to bind to mono- and dimethylated histone H4K20 peptides in vitro [20,21]. L(3)MBT is thought to be a “reader” protein of H4K20me1 (for review see [22]) and to promote methylation-dependent chromatin compaction [23].

l(3)malignant brain tumor (l(3)mbt) was originally identified on the basis of a temperature-sensitive (ts) larval brain cell overgrowth [24]. l(3)mbt also has a strong maternal effect phenotype. Embryos
from l(3)mbt mutant females show abnormalities in cell proliferation [19]. Biochemically L(3)mbt was identified as part of the Drosophila Myb complex involved in regulating transcription [25,26]. Its human ortholog, L3MBTL1, is proposed to function as a negative regulator of E2F target genes such as c-myc and cyclin E7 [29].

To further elucidate the respective functions of PR-Set7 and l(3)mbt we used an RNAi approach in S2 cells. We find that depletion of PR-Set7 inhibits de novo H4K20 monomethylation and results in the activation of the DNA damage checkpoint and a drastic reduction of cell viability. We also detected a low level of abnormal nucleosome compaction in PR-Set7-depleted cells when the ATR checkpoint is abolished. Importantly, these PR-Set7 depleted cells consistently showed an increase in their nuclear diameter, apparently caused by abnormal organization of interphase chromatin. In contrast, the depletion of L(3) mbt reduced the stability of the epigenetic H4K20 monomethyl mark significantly, but it had surprisingly little if any effect on cell viability.

**Materials and Methods**

**RNA Interference and Preparation of S2 Cells Extracts**

Double-stranded RNAs for PR-Set7, l(3)mbt, mei-41, gp and lacZ were prepared as previously described [27]. The templates for synthesizing dsRNAs were produced by PCR.

(for PR-Set7, 5’-ctaatagcactagtgagaatgtatagttta-3’; and
5’-ctaatagcactagtgagaatgtatagttta-3’)

for l(3)mbt, 5’-ctaatagcactagtgagaatgtatagttta-3’
and 5’-ctaatagcactagtgagaatgtatagttta-3’; and

for mei-41, 5’-ctaatagcactagtgagaatgtatagttta-3’
and 5’-ctaatagcactagtgagaatgtatagttta-3’; and

for gp, 5’-ctaatagcactagtgagaatgtatagttta-3’
and 5’-ctaatagcactagtgagaatgtatagttta-3’; and

for lacZ, 5’-ctaatagcactagtgagaatgtatagttta-3’
and 5’-ctaatagcactagtgagaatgtatagttta-3’; and

Double-stranded RNAs for PR-Set7, L(3)MBT, Histone H4K20 me1

polysomal anti-PR-Set7 antibody was diluted 1:1000. Rabbit polysomal anti-monomethylated H4K20 (Upstate Biotechnology) was used at 1:1000 [29]. Rabbit polysomal anti-histone H3 (Abcam) was used at 1:15000. Mouse monoclonal anti-lamin Dm0, ADL67.10 was obtained from the Developmental Studies Hybridoma Bank and used at 1:2000. Rabbit polysomal anti-Chk1 (Phospho-Ser345; GenScript Corp.) and anti-actin (Sigma) were used at 1:500 and 1:1000.

For over expression of L(3)mbt, full length L(3)mbt cDNA was cloned into pMT-FLAG-HA expression vector. 1.2×10⁶ cells were cultured in 4 ml of the medium in 6-well plates for 12 h and then the plasmid was transfectioned using Effectene (Qiagen). 18 h after transfection, the expression was induced by addition of 50 μM CuSO₄. For immunoblotting, mouse monoclonal anti-HA.11 (COVANCE) was used at 1:1000.

**In vivo Histone Methyltransferase Assay**

2.0×10⁶ cells were seeded in 4 ml of medium in 6-well plates. After 12 h, 50 μg of dsRNA were added to each well. After 28 h, 0.5 ml of the cell culture was replated in 24-well plates and 8 h later, 15 μl (15 μCi) tritium-labeled SAM (GE Healthcare) was added to each well. After 2 days, cells were replated in 2 ml of medium in 6-well plates and incubated for another 3 days. Histone extracts were prepared from 2 ml of cell culture. The histones were separated by 14% SDS-PAGE and the gel was fixed in fixation buffer (isopropanol: water: acetic acid = 2:6:1) for 30 min, shaken slowly in Amplify solution (GE Healthcare) for 30 min and dried at 80°C for 2.5 h. The autoradiograph was developed after 10-day incubation at −80°C.

**Cell Viability and DNA Double Strand Break Assays**

1.6–2.8×10⁶ cells were placed in 4 ml of the medium in 6-well plates and dsRNA was added as described above (day 0). The number of cells was counted from day 0 to day 10 and the average ratio of each value to the values of lacZ knockdown cells was calculated. Each time the number of the cells reached 1×10⁵, the cell culture was diluted by 25% to avoid saturation density. For double strand break assay the cells were harvested 4 days after addition of dsRNA and stained with Rabbit polysomal anti–PH2Av (Ser139) antibody (Upstate Biotechnology) diluted 1:500. Double strand breaks were induced by exposing the cells to 10 Gy for 10 minutes.

**FACS Analysis**

To perform FACS analysis 1 ml of cell culture was centrifuged at 800×g for 5 min. The pelleted cells were washed in 10 ml of phosphate-buffered saline (PBS) and resuspended in 500 μl of PBS with 0.1% Triton-X containing 25 μg/ml propidium iodide. FACS was performed with Cytomix FC500 Flow Cytometer (Coulter).

**Micrococcal Nuclease Digestion**

RNAi treatment was performed as described above. S2 cells were harvested after 6 days of treatment. Nuclei were isolated and digested with micrococcal nuclease as described earlier [30], except that 0.1 unit of micrococcal nuclease (Sigma) was added per 1 O.D. (260 nm). Digested DNAs were separated on 1% agarose gels and transferred to nitrocellulose filters. The 5S RNA plasmid 22A8 [31] was labeled using Ready-To-Go DNA Labeling Beads (GE Healthcare Life Sciences) and hybridized.
FISH Analysis

Our FISH protocol has been previously described [32]. S2 cells were treated with dsRNA for 3 days. Oligo probes for the 359 and dodeca heterochromatic repeats were synthesized with a 5′ Cy5 and Alexa488 fluorescent dye, respectively, from Integrated DNA Technologies (IDT). The sequences are as follows: Cy5-359: Cy5-ggggtatgtagctgaattagctgc and Alexa488-dodeca: Alexa488-ACGGGACCGATGCAG. DNA probes to 28B and 69C were synthesized according to standard protocols. PI plasmids (Berkeley Drosophila Genome Project) containing cloned Drosophila genomic DNA corresponding to chromosomal regions 28B1–28B2 (abbreviated as 28B; DS01529) and 69C2–69C8 (abbreviated as 69C; DS02752; [33]) were synthesized and labeled by nick translation/direct labeling (Nysis) following the manufacturer’s protocol. [32,34].

FISH images were collected using an Olympus IX81 microscope with a 60X, N.A. 1.35 lens. Nuclei were imaged by collecting optical sections through the entire nucleus and the analysis of the images was performed by examining one section at a time. The images are from a single section. The number of FISH signals per nucleus was counted manually and the nuclear analysis of the images was performed by examining one section at a time. The images are from a single section. The number of FISH signals per nucleus was counted manually and the nuclear diameters were measured using Olympus cellSens software.

Results

We examined the importance of both PR-Set7 and l(3)mbt in monomethylation of H4K20 using an RNAi approach in S2 cells. To establish that l(3)mbt RNAi treatment reduces the level of the protein we expressed HA tagged full length L(3)mbt and monitored HA-L(3)mbt by western blots. We observed significant reduction of HA-L(3)mbt from day one onward (Figure 1a), indicating that the level of endogenous L(3)mbt is also likely to be strongly diminished. We next monitored PR-Set7 levels for ten days in PR-Set7 knockdown cells and found that the protein is strongly reduced within one day of RNAi treatment (Figure 1b). In l(3)mbt knockdown cells PR-Set7 levels also noticeably decreased on day 4 and stayed at this low level for the duration of our experiment. In contrast to what would be expected based on published results [18], other transcripts are also reduced, indicating that reducing L(3)mbt levels may affect transcription of many genes (Figure S1 and see [35]).

We next determined the effect of lowering PR-Set7, l(3)mbt, PR-Set7 l(3)mbt and, as control, $\beta\zeta$ on the level of H4K20me1 five days after RNAi (Figure 1c). In PR-Set7 no change in H4K20me1 was observed on western blots even though PR-Set7 is strongly reduced. Likely several factors contribute to this finding. First, when PR-Set7 is depleted, activation of the DNA damage checkpoint arrests cell cycle progression and presumably also limits the synthesis of new histones. Second, a pool of unincorporated H4K20me1 protein may exist in S2 cells as our limits the synthesis of new histones. Second, a pool of unincorporated H4K20me1 protein may exist in S2 cells as our

To better assess cell-cycle defects in knockdown cells, we used FACS analysis to profile the cell cycle (Figure 2b, c). In PR-Set7, a change in the number of G1 and G2 cells is already observed on day 1 (data not shown). By day 4 the number of S phase cells is significantly increased (p = 0.0022, t-Test, Fig. 2b), similar to what is observed in PR-Set7-depleted human cells [16,17]. On day 4 the histogram of DNA content of PR-Set7 cells shows that the two peaks at 1c (2N) and 2c (4N) are lower and broader (Figure 2c). The number of cells categorized as sub-G1 or polypliod is also increased, suggesting that in many cells the DNA content is abnormal. As would be expected from the minimal effect of the l(3)mbt knockdown on cell growth, the number of S phase cells and the histogram of these cells are not significantly different from those of control cells.

We previously proposed that H4K20me1 is involved in the maintenance of proper chromatin structure because in PR-Set7 mutant brains the DNA damage checkpoint is activated but DNA double-strand breaks are not increased [9]. To confirm that the depletion of de novo H4K20me1 causes the checkpoint activation, we looked for phosphorylated Chk1, a marker of checkpoint activation [37] and detected the phosphorylated form only in PR-Set7 (Figure 2d). We next asked whether the cell growth defects induced by knocking down PR-Set7 is rescued by simultaneously

PR-Set7, L(3)MBT, Histone H4K20 me1
knocking down the DNA damage checkpoint gene mei-41. Whereas knocking down PR-Set7 alone arrests cell growth and division after about 4 or 5 days, cells simultaneously knocked down for PR-Set7 and either mei-41 or the Drosophila checkpoint kinase grp continue to grow for 6 or 7 days before growth shows evidence of slowing (Figure 3A). We previously found that loss of PR-Set7 in Drosophila larvae did not result in DNA double strand breaks as judged by staining for anti–phosphorylated histone H2Av [9]. We did the same experiments on control and PR-Set7 depleted S2 cells and found that even though the DNA damage checkpoint is activated, we could not observe double strand breaks by anti–phosphorylated histone H2Av (Table S1). Taken together, these data suggest that PR-Set7 functions independently of L(3)MBT in controlling the H4K20 monomethyl mark.

Figure 1. PR-Set7 functions independently of L(3)MBT in controlling the H4K20 monomethyl mark. a, HA tagged full length L(3)MBT (HA-MBT) was expressed in S2 cells and monitored after RNAi against l(3)mbt. b, PR-Set7 levels in PR-Set7 and l(3)mbt knockdown S2 cells. c, Blots of extracts from S2 cells treated for five days with RNAi, probed with anti-mono Histone H4K20 (mono), anti-histone H3 (H3) and anti-Set7 antibodies (Set7). The graph shows the average ratio of each value to the values of lacZ cells. The value of monomethylated H4K20 was normalized to the value of histone H3. d, De novo methylation was assessed by supplying tritium-labeled SAM, to S2 cells treated with different RNAis. The graph shows the average ratio of de novo histone H4 methylation to total de novo histone methylation of H2A, H2B and H3. Error bars for both graphs show two SDs (n = 3). The knockdown of l(3)mbt does not affect the level of tritium labeling on histone H4, showing that newly monomethylated H4K20 is stable for duration of the experiment even in the absence of L(3)MBT.

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together these results suggest that the growth arrest in the single PR-Set7 knockdown cells is due to the activation of the DNA damage checkpoint, while another unknown mechanism must be responsible for the reduction in growth rate seen later on in the PR-Set7 mei-41 or PR-Set7 grp double knockdown cells.

In the Pr-Set7 knockdown experiments in vertebrates, cells tend to accumulate in S phase [16,17]. It is possible that abnormalities in chromatin organization could occur because de novo H4K20 methylation is tightly coupled to the process of nucleosome assembly, and therefore in the absence of this methylation mark DNA synthesis and de novo nucleosome assembly could become uncoupled. In this scenario, when Pr-Set7 activity is compromised, DNA replication would continue without properly assembling chromatin on the newly replicated DNA. The presence of stretches of abnormally packaged DNA would then activate the DNA damage checkpoint.

We therefore examined the effects of reducing PR-Set7 on chromatin architecture in double knockdown (PR-Set7;mei-41). Figures 3b and c show the nucleosomal ladders produced by digestion of nuclei prepared from either mei-41 (b) or PR-Set7;mei-41 (c) knockdown cells. The most striking finding is that the nucleosomal ladder remains largely intact in the PR-Set7;mei-41 knockdown cells, and that there is no evidence of extensive regions of DNA that lack nucleosomes. Since the double knockdown cells have divided once or twice under conditions that would have induced a cell cycle arrest in the single knockdown cells, it would appear that PR-Set7 is not required to sustain the process of de novo nucleosome assembly. While no gross disruptions in global nucleosome organization are observed, some differences between the double knockdown and the single knockdown control were evident in this and other experiments suggestive of differences in nucleosome packing. The double knockdown appears to be more sensitive to micrococcal nuclease digestion (Fig. 3, compare corresponding lanes in panels b and c). Larger polynucleosomes (4–8 mers) are present in lower yield and/or appear less distinct (e.g., 5 & 6) than in the control knockdown. This increased sensitivity could indicate that nucleosome arrays/bulk chromatin may not be as tightly compacted on average in the double knockdown compared to the control cells.

To confirm these findings we examined the chromatin organization of the ~100 tandemly repeated 5 s rRNA genes. Previous studies show that the two nucleosomes associated with the 375 bp 5 s RNA gene + spacer are positioned relative to the underlying DNA sequence [31], [38]. As a consequence, MNase preferentially generates multiples of two nucleosomes, while fragments corresponding to an odd number of nucleosomes are considerably less frequent (see Fig. 4a). The 5 s repeat also contains sequences that are preferentially cleaved by micrococcal nuclease in naked DNA digests. Normally these sequences are incorporated into nucleosomal DNA and are comparatively resistant to micrococcal cleavage; however, if nucleosomes were depleted in some of the 5 s rDNA repeats, these sites would be exposed and generate fragments of non-nucleosomal lengths. Figure 4b shows the characteristic di-nucleosomal ladder of 5 s...
rDNA repeats generated by micrococcal nuclease digests of chromatin from PR-Set7; mei-41 double knockdown cells, and there is no evidence of enhanced cleavage at naked DNA sites. This result supports the main conclusion drawn from the MNase cleavage pattern in bulk chromatin, namely that de novo nucleosome assembly can proceed in the absence of PR-Set7 activity under conditions in which activation of the DNA damage checkpoint is blocked. Additionally, as was evident in the bulk chromatin digest, there appear to be some abnormalities in the 5 s rDNA digestion pattern that are suggestive of differences in nucleosome compaction. The 5 s chromatin in the double knockdown cells is more sensitive to MNase digestion and large oligonucleosomes appear earlier in the digestion and are lost more rapidly. The yield of odd numbered bands (3, 5 and 7

Figure 3. Cell viability and nucleosome analysis in normal and PR-Set7 mei-41 cells. The graph shows the average ratio of each cell count to that of lacZ RNAi cells (n≥3). The knockdown of either mei-41 (Drosophila ATR) or grp (Chk1) disrupts the DNA damage checkpoint and therefore rescues the lethality of the PR-Set7 cells partially. Nucleosomal ladders from mei-41 (b), and PR-Set7 mei-41 (c) knockdown cells. Nuclei were subjected to micrococcal nuclease digestion for the times indicated, separated electrophoretically. doi:10.1371/journal.pone.0045321.g003
nucleosomes) also is visibly elevated compared to the control. Both
the increased sensitivity of 5 s chromatin and the increase in the
yield of odd number bands (particularly the trinucleosome) was
seen in several experiments.

Though the experiments reported above clearly demonstrate
that PR-Set7 is not needed for de novo nucleosome assembly, the
fact that overall micrococcal nuclease sensitivity appeared to be
elevated, albeit only slightly, was intriguing when considered in the
context of previous studies on larval brains in PR-Set7 mutants [9].
In these studies, we found that interphase nuclei in the PR-Set7
mutants are enlarged compared to nuclei in wild type brains.
Indeed, as was observed in the fly, about 20% of PR-Set7-depleted
tissue culture cells had nuclear volumes at or above the 95\textsuperscript{th}
percentile of control knockdown cells. The average nuclear
diameter in PR-Set7-depleted cells was 6.6 μm compared to
4.5 μm in controls (Fig. 5). We considered the possibility that this
phenotype could reflect polyploid cells that arise due to the role of
PR-Set7 in mitosis. However, the modest increase in polyploidy
observed in our FACS analysis cannot explain the significant
proportion of cells with this phenotype.

An alternative explanation is that the enlarged nuclei are caused
by an alteration in high order chromatin compaction or
organization. We reasoned that if higher-order chromatin folding
was perturbed in PR-Set7-depleted cells, the spatial integrity of

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<td>1.12</td>
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<td>0.97</td>
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<tr>
<td>nucleosome 8/7</td>
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Figure 4. PR-Set7-depleted cells show abnormal nucleosome spacing. Nuclei isolated from control mei-41 cells (a), and PR-Set7 mei-41 (b),
were subjected to micrococcal nuclease digestion for the indicated times, and separated electrophoretically. The DNA was transferred to
nitrocellulose and hybridized with a \textsuperscript{32}P-labeled 5S probe [31]. The intensity of the radioactivity was measured and calculated as a ratio of the even
numbered nucleosome divided by the odd one; e.g. ratio of dinucleosome/trinucleosome, tetrinucleosome/trinucleosome etc.
doi:10.1371/journal.pone.0045321.g004
large chromosomal regions might be altered relative to control cells. To test this possibility, we performed DNA-FISH in male-derived S2 cultured cells [39] that contain two X chromosomes due to tetraploidy of the cell line [40–41]. Following PR-Set7 knockdown, we observed an increase in the number of FISH signals targeting the 359-bp repeat, located in the pericentromeric heterochromatin of the X chromosome; only 65% of PR-Set7-depleted cells exhibited a single FISH signal as compared to 74% in control cells (Fig. 5; p < 0.0244). The increase in the number of FISH signals was not unique to the X chromosome and was also observed when FISH targeted the dodeca satellite repeat on the 3rd chromosome as well as two euchromatic regions, 28B and 69C, on the 2nd and 3rd chromosomes, respectively (Fig. 5). Similar results were obtained in a recent screen that showed that dsRNAs targeting PR-Set7 caused a disruption in MSL staining, a complex important for dosage compensation, and an increase in the number of FISH signals in Drosophila cultured cells, which the authors suggests is due to a disruption in somatic homolog pairing [42].

We propose that increased FISH signals could be explained by defects in chromatin condensation, chromosome breakage, or separation of sister chromatids or homologs, each of which would

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**Figure 5. PR-Set7-depleted cells show disrupted nuclear organization.** (a) Representative images of PR-Set7-depleted S2 cells that exhibit larger nuclei and increased number of FISH signals per nucleus. Scale bars are 5 μm. (b) The average nuclear diameter is increased as compared to control cells (lacZ RNAi). (c) dsRNA directed against PR-Set7 decreases the number of nuclei with a single FISH signal when targeting peri-centromeric heterochromatin regions (359 on X, red; dodeca on 3rd, green) or euchromatic regions (28B on 2nd; 69C on 3rd). A minimum number of 100 nuclei were counted for each of three replicate tests. P-values were calculated using an unpaired t test.

doi:10.1371/journal.pone.0045321.g005
be consistent with a role for PR-Set7 in higher-order chromatin organization during interphase in Drosophila cells. This would suggest that H4K20me1 is essential for chromatin packaging both during interphase and mitosis [9,16,43]. The failure to fully condense chromatin could also explain our nucleosome results. It is likely that the differences between normal and PR-Set7 nucleosome positioning are lost when chromatin that is not properly condensed is digested.

Discussion

In our studies we established that PR-Set7 sets the H4K20 monomethyl mark in vivo and that at least in S2 cells K20 is the only amino acid that is methylated. We further found that depleting PR-Set7 in Drosophila S2 cells leads to the activation of the DNA damage checkpoint and within about 10 days to cell death. When the DNA damage checkpoint is abrogated by double knock-down of PR-Set7 and the checkpoint genes met-41 or grp, the half-life of the cells is increased by 1 to 2 days, but ultimately the cells still die, suggesting that whatever is perturbed in the absence of PR-Set7 cannot be repaired. We did not observe double strand breaks when staining for anti–phosphorylated histone H2A. This does not agree with results observed in vertebrate cells [17] [16] and may be because, in Drosophila, H2Av is not phosphorylated in the absence of H4K20me1 or the specific epitope is obscured. Alternatively, double strand breaks may not exist and the checkpoint is activated because of abnormal chromatin organization or because protein complexes are not removed in a timely manner as observed in S. cerevisiae [44].

In this context it is interesting to note that in vertebrates H4K20 me2 is implicated in double strand break repair [45]. Because H4K20me1 is the likely substrate for Suv4-20H1 and H2, the di- and trimethyltransferases, an additional link between H4K20 methylation and double strand breaks seems to exist. However, besides potentially setting the monomethyl mark at double strand breaks, PR-Set7 would have to have additional functions, because in both flies and vertebrates PR-Set7 mutants have a substantially stronger phenotype than the loss of the Suv4-20 enzymes [46] [11].

The increase in nuclear volume, together with the changes in the number of FISH signals per nucleus observed in interphase cells following PR-Set7 RNAi would be consistent with a role for PR-Set7 in chromosome compaction and higher-order chromatin organization (9,16,43). Bateman, 2012 #179). Interestingly, mass spectrometry experiments show that the H4K20 monomethyl mark is set at the G2/M transition well after newly synthesized histone H4 is incorporated into chromatin in S phase [47]. These findings suggest that the abnormalities in chromosome compaction and organization evident in interphase nuclei might be due to defects arising during the G2/M transition. Consistent with this possibility, cells depleted for only Pr-Set7 appear to arrest mostly in early mitosis. But unlike what is observed in larval brains, there is also a subset of cells that arrest in S phase [9]; these may represent cells that despite the abnormalities in higher order chromatin organization are able to continue through the cell cycle until a checkpoint is activated during S. The discrepancy between the brain and tissue culture cells may be a reflection of differences in their cell cycle and developmental potential.

Results from several laboratories suggest that PR-Set7 function is coupled to DNA replication based on its targeting to the dividing fork via its interaction with PCNA [17] [15]. Our findings indicate that while abnormalities in chromatin organization and compaction appear to accumulate after growth without PR-Set7 activity, these defects are inconsistent with massive disruptions in de novo nucleosome assembly during replication [48]. Instead, the DNA damage checkpoint activation must arise from more subtle abnormalities in chromatin or DNA structure.

As for the l(3)mbt, its functional requirement does not appear to overlap with that of PR-Set7, neither in tissue culture as shown here, nor in flies. In larva the loss of l(2)mbt results in an expansion of the neuroblast pool and subsequent tumorous overgrowth of the optic lobe [49] while in PR-Set7 mutants the cell cycle of neuroblasts arrests in early mitosis resulting in fewer cells [9]. PR-Set7 is essential for de novo methylation of H4K20. While the loss of H4K20me1 could occur either because in the absence of L(3)mbt protection the H4K20me1 is lost, or it could be a secondary effect. Consistent with the latter explanation, recent results show that L(3)mbt binds to DNA boundary elements and affects the level of transcription of Salvador-Wart-Hippo pathway genes both positively and negatively [42]. That L(3)mbt possibly controls expression of many genes is also supported by our observation that the transcription level of all genes we tested was reduced compared to wild type (Fig. 1B).

Supporting Information

Figure S1 Transcriptional levels of several genes measured by quantitative real-time PCR five days after RNAi treatment against PR-Set7 and l(3)mbt. Each value is normalized to the level of rp–49. The graph shows the average ratio of each value of PR-Set7 and l(3)mbt knockdown cells to the values of lacZ knockdown cells (control). Error bars show two SDs (n ≥ 3). The asterisk means that the value was not detected. (TIFF)

Figure S2 Time-course experiment to detect the reduction of monomethylated H4K20 in PR-Set7 and l(3)mbt knockdown cells. The number of cells was counted at each point and the number of cell divisions was calculated. The intensity of the bands was quantified by ImageJ, and the value of monomethylated H4K20 (mono) was normalized to the values of both histone H3 and lacZ knockdown cells (control). Cells were plated at two densities (1.6×10^6 and 2.8×10^6). Two examples of western blots are shown (a and b). In the l(3)mbt cells H4K20 monomethylation is reduced to around 40% of controls after four divisions. In the PR-Set7 cells monomethylation is reduced by at most 21%, suggesting that the slow growth of PR-Set7 cells (see Figure 2a) is not the reason why we failed to detect reduction of the monomethylation. (TIF)

Table S1 The number of double strand breaks is not increased in PR-Set7 depleted cells. (>750 cells were counted for each sample). (PDF)

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Author Contributions

Conceived and designed the experiments: AS EJ PS RS. Performed the experiments: AS EJ TA. Analyzed the data: AS EJ PS RS. Contributed reagents/materials/analysis tools: AS EJ TA PS RS. Wrote the paper: AS EJ PS RS.
References