Direct interaction between DNMT1 and G9a coordinates DNA and histone methylation during replication

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Chromatin methylation is necessary for stable repression of gene expression during mammalian development. During cell division, DNMT1 maintains the DNA methylation pattern of the newly synthesized daughter strand, while G9a methylates H3K9. Here, DNMT1 is shown to directly bind G9a both in vivo and in vitro and to colocalize in the nucleus during DNA replication. The complex of DNMT1 and G9a colocalizes with dimethylated H3K9 (H3K9me2) at replication foci. Similarly, another H3K9 histone methyltransferase, SUV39H1, colocalizes with DNMT1 on heterochromatic regions of the nucleoli exclusively before cell division. Both DNMT1 and G9a are loaded onto the chromatin simultaneously in a ternary complex with loading factor PCNA during chromatin replication. Small interfering RNA (siRNA) knockdown of DNMT1 impairs DNA methylation, G9a loading, and H3K9 methylation on chromatin and rDNA repeats, confirming DNMT1 as the primary loading factor. Additionally, the complex of DNMT1 and G9a led to enhanced DNA and histone methylation of in vitro assembled chromatin substrates. Thus, direct cooperation between DNMT1 and G9a provides a mechanism of coordinated DNA and H3K9 methylation during cell division.

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Epigenetic information, encoded into the chromatin by virtue of covalently modified DNA and histone molecules (Holliday 1994; Turner 2000) provides a heritable mechanism to regulate and maintain gene expression patterns through DNA replication. The basic unit of chromatin, the nucleosomes, consists of 147 base pairs (bp) of double-stranded DNA wrapped around two copies of core histones H2A, H2B, H3, and H4 (Luger et al. 1997). Adjacent nucleosomes can fold into higher order structures that can restrict the accessibility of various basal and specific transcription factors or enzymes. During cell division and DNA replication, maintenance of the DNA methylation pattern onto the daughter strand is essential for stable repression of genes. In the G1 and G2 phases of the cell cycle, DNMT1 remains diffused throughout the nucleoplasm, but associates with replication foci during S phase (Leonhardt et al. 1992; Liu et al. 1998). Movement of the replication machinery transiently disrupts the chromatin fiber and leads to the random segregation of the parental nucleosomes among the two daughter strands (Gruss and Sogo 1992). Additional nucleosomes are added onto the newly synthesized DNA via deposition of newly synthesized histone [H3/H4] tetramers followed by H2A/H2B dimers (Jackson 1990; Gruss et al. 1993). Furthermore, the new nucleosomes contain acetylated histone H4 and H3, which is a prerequisite for nuclear transport (Mosammaparast et al. 2002). Thus, nucleosomes in the newly synthesized DNA are expected to contain new acetylation but lack other chromatin-specific epigenetic marks such as methylation. To re-establish the original epigenetic marks, the new histones must undergo removal of some modifications while allowing other modifications to occur. Indeed, acetyl groups on new histones are rapidly removed in mammalian cells (Taddei et al. 1999), perhaps allowing the patterns of modification existing prior to replication to be re-established. Although the core regions of the histone octamer remain inside the nucleosome, the N-terminal tails are exposed, and thus are amenable to events such as methylation, acetylation, phosphorylation, sumoylation, ADP-ribosylation, and ubiquitination (Van Holde 1988; Wolffe 1998).

The N-terminal regions of histones H3 and H4 are highly modified. Lysine residues at positions 4, 9, 27, and
36 of histone H3 and 20 of histone H4 are methylated [Lachner et al. 2003]. Methylation of Lys 9 of histone H3 (H3K9) has been extensively studied and shown to correlate with transcriptional gene silencing. Furthermore, the methylation of lysines on histones is reversible and shown to regulate gene expression [Wysocka et al. 2005]. The methylated H3K9 serves as a specific binding site for heterochromatin protein 1 (HP1). The Suv39h class of enzymes maintains methylation of heterochromatin [Rea et al. 2000], while in mammals G9a is likely to be the major euchromatic H3K9 histone methyltransferase. Gene disruption of G9a in mice significantly decreases H3K9 methylation in euchromatic regions [Tachibana et al. 2002]. Using a panel of antibodies specific for mono-, di-, and trimethylated H3K9 with G9a-null embryonic stem cells or Suv39h1 and Suv39h2 double-null mouse embryonic fibroblast cells, it was established that G9a is specific for mono- and dimethylation, whereas Suv39h1 and Suv39h2 are largely responsible for trimethylation [Rice et al. 2003]. However, recent biochemical studies with purified full-length G9a or its SET domain, along with a substrate peptide bearing the N terminus of histone H3, suggest that G9a is capable of mono- and dimethylation, whereas Suv39h1 and Suv39h2 are largely responsible for trimethylation [Patnaik et al. 2004; Collins et al. 2005]. G9a-null mice display severe growth retardation and die between embryonic days 9.5 and 12.5 due to deregulation of developmental genes [Tachibana et al. 2002]. Thus, G9a deficiency is not complemented by other known H3K9 methyltransferases. A similar growth retardation and embryonic lethality was observed in DNMT1-null mice [Li et al. 1992]. These observations support a direct link between epigenetic gene regulation and mammalian development.

DNA replication in eukaryotic cells stimulates a complex array of responses [Stillman 2005]. During DNA replication, DNMT1 is loaded onto replication fork via PCNA (Chung et al. 1997) and participates in methylation of CpG sites [Leonhardt et al. 1992]. Histone methyltransferase-mediated DNA methylation is demonstrated for EZH2 target promoters [Vire et al. 2006], IAP elements, and Mage genes [Dodge et al. 2004; Tachibana et al. 2002]. In a previous study, we demonstrated that doxorubicin-induced repression of the anti-apoptotic survivin gene is caused by DNA and histone methylation of its proximal promoter, which is mediated by the physical recruitment of DNMT1 [Estève et al. 2005a]. However, the mechanism underlying coordinated DNA and histone methylation during DNA replication has not been previously reported. In this report, we demonstrate a physical association and functional cooperation between two epigenetic factors, DNMT1 and G9a, during DNA synthesis.

Results

Interaction between maintenance DNMT1 and G9a in vivo

The observation that DNA and histone methylation on euchromatic regions is carefully preserved during replication raised the possibility that the histone and DNA methyltransferases engage in direct cross-talk. To determine whether DNMT1 associates with G9a in cells, we cotransfected a GFP-G9a fusion construct along with 6xHis-tagged DNMT1 into COS-7 cells. After 48 h, we immunoprecipitated protein complexes from cell extracts using anti-GFP and anti-DNMT1 antibodies. The GFP and DNMT1 antibodies both precipitated GFP-G9a (Fig. 1A, lanes 2, 3, top and middle), and DNMT1 (Fig. 1A, lanes 2, 3, top and bottom). Precipitation of DNMT1 required G9a, as cotransfection of the GFP empty vector along with the DNMT1 construct did not yield a detectable complex formation (Fig. 1A, lane 1). In a similar experiment, we were also able to coimmunoprecipitate endogenous DNMT1 and G9a from HCT116, but not in DNMT1-null HCT116 cells (Fig. 1B). Endogenous coimmunoprecipitation of DNMT1 and G9a was also observed in HEK293 cells (data not shown). We further verified the existence of DNMT1 and G9a in the same complex by gel filtration chromatographic separation of Jurkat cell nuclear extract. The estimated molecular mass of DNMT1 and G9a are ~183 and 140 kDa, respectively. A complex between DNMT1 and G9a would result in a combined mass >320 kDa, thus the complex can be resolved from DNMT1 or G9a monomer and be detected by Western blot analysis. Indeed, the chromatographic peak containing DNMT1, G9a, and H3K9me2 eluted approximately at a molecular mass of 449 kDa (Fig. 1C) as determined by gel filtration elution of the standard protein. All of the G9a was in complex with DNMT1 since no G9a protein was detected eluting at the region of the 158-kDa protein standard. To further validate this observation, we performed immunoprecipitation of DNMT1 and G9a using anti-DNMT1 or anti-G9a antibodies. A microgram of anti-DNMT1 captured some DNMT1, but all of the G9a. The unbound fraction of the nuclear extract did not display the presence of free G9a. When a microgram of anti-G9a was used for immunoprecipitation, all of the G9a was captured, but only a fraction of DNMT1 was found in the captured complex (Fig. 1D). A nonspecific anti-GFP antibody did not immunoprecipitate either G9a or DNMT1 (data not shown). We conclude from these experiments that DNMT1 and G9a exist in a stable complex in vivo and almost all of the G9a is in the form of a complex with DNMT1. However, it remained possible that the association was mediated by other proteins that bridged the interaction.

To determine whether DNMT1 directly interacts with G9a we used pull-down assays using purified proteins. Fragments of G9a encompassing the entire 1263-amino-acid protein were fused to GST and purified on glutathione beads. The bound GST fusion proteins were incubated with purified recombinant full-length DNMT1 (purified using a baculovirus expression system) [Pradhan et al. 1999]. The N-terminal region of G9a encompassing the first 463 amino acids bound tightly to DNMT1, whereas a deletion bearing amino acids 291–633 did not (Fig. 1E). When the GST-G9a full-length and DNMT1 full-length enzymes were incubated together, DNMT1 re-
mained bound to G9a (Fig. 1E). A reciprocal GST pull-down assay was performed using GST fusion fragments of DNMT1 encompassing the entire 1616-amino-acid enzyme and recombinant full-length G9a. G9a bound to a region of DNMT1 containing the first 446 amino acids (Fig. 1F). These data suggest that DNMT1 and G9a directly interact and that the N termini of DNMT1 and G9a mediate this interaction. To unequivocally demonstrate the binding between the N-terminal region of DNMT1 and G9a, additional in vitro binding assays were performed. We incubated the full-length DNMT1 with either a full-length G9a or an N-terminal deletion MBP fusion mutant lacking the first 633 amino acids (MBP-delG9a), the putative DNMT1-binding motif. The incubation mixture was subjected to immunoprecipitation using either anti-DNMT1, anti-G9a, or anti-MBP antibodies and examined for the presence of both G9a and DNMT1 by Western blot analysis. Indeed, a strong complex between DNMT1 and G9a full-length proteins was observed (Supplementary Fig. 1, lanes 1,3) and this binding was disrupted in the presence of MBP-delG9a (Supplementary Fig. 1, lanes 2,4).

The physical interaction of DNMT1 and euchromatic histone methyltransferase G9a (Tachibana et al. 2002) represents a potential mechanism by which DNA and histone methylation could be coordinated on euchromatic regions in vivo during replication. One prediction of this idea is that G9a and DNMT1 should colocalize at replication foci during DNA replication in early S phase, when the bulk of the euchromatic region of the genome undergoes replication (Gilbert 2002).
Colocalization of DNMT1 and G9a during DNA replication and their association with methylated H3K9

DNMT1 associates with replication foci during DNA replication via direct interaction with PCNA, a replication processivity factor [Chuang et al. 1997]. Our findings that DNMT1 interacts with G9a suggested the possibility that DNMT1 recruits G9a to active sites of replication. To address the functional significance of the complex of DNMT1 and G9a and to understand the sequence of events occurring during replication, we cotransfected a GFP-G9a fusion along with a DsRed-DNMT1 fusion construct into either HeLa or COS-7 cells. The cells were synchronized with aphidicolin at the G1/S border and visualized for the localization of the fusions in the nuclei by fluorescence microscopy. The quality of the synchronization at different time points was monitored by bromodeoxyuridine (BrdU) (pulse chase) incorporation, immunostaining, and scoring for BrdU-incorporated cells. At the 2–4 h time point, ~90% of the cells displayed BrdU incorporation. The percentage of BrdU-incorporated cells increased to 98% by the eighth hour. The replication foci, as revealed by BrdU incorporation, appeared from small spotty clusters to large red punctate nuclear spots as the replication time progressed [Supplementary Fig. 2]. In the colocalization studies, distinctive green spots appeared in the nuclei of the cells transfected with the GFP-G9a fusion construct. A similar pattern of red punctate nuclear spots was observed in cotransfected cells, revealing that DNMT1 is localized in the nucleus [Fig. 2A], as observed previously [Leonhardt et al. 1992]. Superimposition of GFP-G9a and DsRed-DNMT1 resulted in yellow nuclear spots throughout the nucleus, excluding the nucleolus, which remained dark [Fig. 2B]. Similarly, a DsRed fusion construct containing the first 502 amino acids of DNMT1 was cotransfected with a GFP fusion of G9a containing the first 464 or 598 amino acids. We observed yellow spots in the DNMT1–G9a merged panels [Fig. 2C,D] similar to Figure 2A, panels II and III, demonstrating the colocalization of the N terminus of both DNMT1 and G9a. Thus, the cumulative results from GST pull-down analysis [Fig. 1E,F] and colocalization studies between DNMT1 and G9a confirm recruitment of both epigenetic factors.

To validate that G9a and DNMT1 colocalize at active sites of DNA synthesis, transfected cells were pulse-chased with BrdU. BrdU becomes incorporated into newly synthesized DNA, thus macromolecules at the vicinity of newly synthesized DNA may be colocalized along with the BrdU signal. Previous studies showed that DNMT1 colocalized with BrdU in cycling cells [Leonhardt et al. 1992]. Thus, we initially examined whether G9a would also colocalize with BrdU. As shown in Figure 2E, the yellow punctate pattern indicates that GFP-G9a was almost exclusively colocalized with BrdU (red) in the replicating nuclear DNA. A parallel experiment with GFP-DNMT1 and BrdU displayed a similar pattern of colocalization as observed previously [Fig. 2F, Leonhardt et al. 1992]. The data of Figure 2A showing G9a and DNMT1 colocalization, coupled with the data of Figure 2, E and F, provide strong support for the idea that G9a and DNMT1 localize close to sites of DNA synthesis.

If DNMT1 recruits G9a to the chromatin and the interaction is indeed functional, there should be a correlation between the DNMT1–G9a association and histone methylation. We addressed this hypothesis via two sets of immunocytochemistry experiments. In the first set, cells were transfected with the GFP-G9a fusion construct and probed with H3K9me2-specific antibodies [Fig. 2G]. In the second set, cells were transfected with the DsRed-DNMT1 fusion construct and probed with H3K9me2-specific antibodies [Fig. 2H]. The majority of H3K9me2 was in close proximity to G9a [Fig. 2G] and DNMT1 [Fig. 2H] as evidenced by a similar punctate colocalization pattern. Thus, DNMT1, G9a, and H3K9me2 are present in the vicinity of DNA synthesis during cell division.

Association and interaction between DNMT1 and SUV39H1 occurs on pericentric heterochromatin before DNA replication

Mammalian heterochromatin is enriched with H3K9me3, the catalytic product of the Suv39h1 class of histone methyltransferases [Lehnertz et al. 2003]. Genetic deletion of Suv39h1, a mouse homolog of human SUV39H1, resulted in reduction of H3K9me3, but not mono or H3K9me2, confirming its catalytic role in trimethylation [Lehnertz et al. 2003; Rice et al. 2003]. Furthermore, genetic deletion of G9a resulted in drastic reduction of mono and H3K9me2, suggesting that G9a maintains some mono and the majority of H3K9me2 [Peters et al. 2003; Rice et al. 2003]. Additionally, Suv39h1 and H3K9me3 colocalize on hetero-
chromatin (Fig. 3A; Peters et al. 2001) as was observed for DNMT1 and H3K9me2 localization with either DNMT1 or G9a. Fusion proteins are shown at the top of the panels along with immunocytochemical detection of BrdU or H3K9me2 as indicated. Panels are designated A–H. In the left-hand corner of B–H, the identity of the fusion proteins and the antibody used for immunocytochemistry is noted. (Nu) nucleus stained by hoechst staining; [BrdU] bromodeoxyuridine; [H3K9me2] dimethylated histone H3 Lys 9. The DNMT1 and G9a fusion constructs used are either fused with DsRed (DsRed-DNMT1) or GFP (GFP-DNMT1). A fusion protein containing either a GFP or DsRed partner along with a portion of G9a or DNMT1 as indicated in bracket is used. (A) G9a and DNMT1 co-localization during cell cycle. On the left side of the panel, cytochemical observation after synchronization in hours and percent of cells where DNMT1 and G9a fusions are colocalized are given. For each time [at 0 time [panel I], between 2 and 4 h [panel II], between 8 and 12 h [panel III], and after 15 h [panel IV] post-aphidicolin treatment], 20–50 cells were counted. (B) DNMT1 (1–446) and G9a. (C) DNMT1 (1–502) and G9a (1–598). (D) DNMT1 (1–502) and G9a (1–464). (E) G9a and BrdU. (F) DNMT1 and BrdU. (G) H3K9me2 and G9a. (H) DNMT1 and H3K9me2. Amino acids numbers are in parentheses.
Figure 3. Colocalization and interaction between DNMT1 and SUV39H1. (A) Cytochemical localization of SUV39H1 and H3K9me3 in heterochromatin. (B) Cytochemical localization of DNMT1 and H3K9me3 in heterochromatin. (C) Mapping of the SUV39H1-binding region on DNMT1 using overlapping GST fusion fragments of DNMT1. Various GST fusions of DNMT1 are indicated above with amino acids in parentheses and antibody used shown at right. (D) Schematic diagram of DNMT1 with SUV39H1-binding region on it shown below. (E) SUV39H1 and DNMT1 do not colocalize during DNA replication, but are associated in heterochromatin. On the left side of the panel cytochemical observation after synchronization in hours and percent of cells where DNMT1 and SUV39H1 fusions are colocalized are given. For each stage [at 0 time [panel I], between 2 and 4 h [panel II], between 8 and 12 h [panel III], and after 15 h [panel IV] post-aphidicolin treatment], 20–50 cells were counted.
were blotted and probed with anti-SUV39H1. The first 446 amino acids of DNMT1 strongly associated with SUV39H1 (Fig. 3C,D). A similar association between DNMT1 and SUV39H1 was observed previously and the complex was catalytically competent (Fuks et al. 2003).

To elucidate whether DNMT1 and SUV39H1 are associated during DNA replication, additional colocalization experiments using GFP-SUV39H1 and DsRed-DNMT1 fusion constructs were performed in synchronized HeLa cells. Fluorescent localization revealed that, indeed, both DNMT1 and SUV39H1 are colocalized in the vicinity of heterochromatin and pericentric heterochromatin regions of 75% of the cells, as observed by the yellow merged pattern around the nucleolus that was densely stained by the nuclear staining (Fig. 3E, panel I). However, during DNA replication, when DNMT1 is located at the replication foci and visualized as the red punctate spots, SUV39H1 did not colocalize with it (Fig. 3E, panels II,III,IV). This observation was confirmed by a BrdU pulse-chase experiment, where SUV39H1 did not colocalize with BrdU at the replication forks although SUV39H1 appeared throughout the chromatin [data not shown]. In a control immunocytochemical experiment, SUV39H1 and H3K9me3 were colocalized on the heterochromatin as was observed in Figure 3A. Exclusion of SUV39H1 from the replication foci and colocalization of it on heterochromatin with DNMT1 suggests a role for these enzymes in post-replicative H3K9 trimethylation and DNA methylation. This hypothesis gains further support by the presence of a chromodomain on SUV39H1 that can bind dimethylated H3K9 tails with high affinity (Jacobs et al. 2004). Perhaps this chromodomain-mediated anchoring of SUV39H1-dimethylated H3K9 facilitates H3K9 trimethylation as proposed earlier (Peters et al. 2003; Rice et al. 2003). Furthermore, mutation of the SUV39H1 chromodomain was shown to render the enzyme catalytically inactive (Chin et al. 2006). It is also plausible that the SUV39H1–DNMT1 binary complex may remain bound to methylated H3K9 and recruit a methylation-independent chromatin repressor complex onto the heterochromatin.

Molecular mechanisms of DNMT1 and G9a cooperation

To validate the cytochemical colocalization of DNMT1 and G9a at the vicinity of replication foci and to understand the events associated with replication-dependent chromatin methylation, we examined the connection between DNMT1–G9a loading and Lys 9 dimethylation of H3. HeLa cells were synchronized by thymidine and aphidicolin and at different time points, cells were harvested, and chromatin fractions were Western blotted to determine whether DNMT1 and G9a were loaded onto the chromatin. Indeed, the chromatin fractions displayed a gradual increase in G9a and DNMT1 in a time-dependent manner corresponding with increased amounts of H3K9me2. Significantly, the loading control H3 and SUV39H1 remained very similar in all of the time points (Fig. 4A). We further immunoprecipitated the synchronized cells with anti-PCNA antibodies to find out whether the docking of DNMT1–G9a binary complex is mediated by PCNA. We observed a gradual increase in DNMT1–G9a loading with a normalized PCNA level, suggesting that more DNMT1–G9a binary complexes are being formed as replication progresses (Fig. 4B). This suggests a possible formation of a trimmeric complex during DNA replication. As replication proceeds, the loading and increase in PCNA–DNMT1–G9a trimeric complexes in the nucleus (Fig. 4B) is observed, demonstrating a constant supply of active enzyme complexes to the replication site. To validate the role of DNMT1 in maintaining G9a-mediated H3K9 methylation, small interfering RNA (siRNA)-mediated knockdown of the endogenous DNMT1 or G9a were performed in HeLa cells and total cellular extracts and chromatin fractions were examined for H3K9me2 changes. Knockdown of G9a resulted in ~65% reduction of H3K9me2 in both total extract and chromatin fraction as compared with the control knockdown without impairment of DNMT1 loading on the chromatin (Fig. 4C, top). Moreover, DNMT1 knockdown resulted in ~50% reduction of H3K9me2 and correlated with a decrease in G9a loading onto the chromatin (Fig. 4C, right). These results indicate that knockdown of either G9a or DNMT1 results in impairment of total H3K9 methylation. A similar decrease in H3K9 di- and trimethylation was observed previously in DNMT1-null HCT116 cells (Espada et al. 2004).

To validate whether the impaired H3K9 methylation in DNMT1 knockdown cells is due to poor loading of G9a onto the replicating chromatin, we examined and compared the pattern of GFP-G9a and BrdU incorporation in siRNA-mediated DNMT1 knockdown COS-7 cells and DNMT1-null HCT116 cells. BrdU incorporation in both DNMT1 knockdown or DNMT1-null cells appeared throughout the nucleus, as did G9a [Fig. 4D,E]. However, the majority of G9a did not colocalize with BrdU [Fig. 4D,E, merged]. Therefore, these observations suggest that DNMT1 participates in localization and loading of G9a onto chromatin during DNA replication.

Chromatin methylation coordinated by DNMT1 and G9a

To demonstrate that DNMT1–G9a indeed coordinates chromatin methylation during DNA replication on specific gene sequences, we investigated the intergenic spacer (IGS) elements of rDNA. The IGS-rDNA contained six CpG dinucleotides in a span of 78 nucleotides [nt] that are a potential target for DNMTs. Recent studies indicate that an epigenetic network mediates the transcriptional state of rDNA and DNA methylation plays a major role in rDNA silencing (Santoro and Grummt 2001; Santoro et al. 2002). To establish whether IGS of rDNA is methylated and that DNMT1 is responsible for IGS-rDNA methylation, methylation-sensitive PCR (MSP) was performed on the siRNA-mediated DNMT1 knockdown genome or on cells treated with zebularine, a DNMT1-specific mechanistic inhibitor (Karpf and Jones 2002; Cheng et al. 2004). MSP can dis-
Figure 4. DNMT1 facilitates G9a-mediated methylation of the chromatin. (A) Concurrent DNMT1 and G9a loading onto the chromatin during S phase. HeLa cells were synchronized by thymidine and aphidicolin and released into regular medium. Time points in hours after the release is indicated at the top with antibody used for Western analysis indicated at the bottom. Chromatin fractions were used in the blot as indicated. (B) Analysis of DNMT1 and G9a recruitment on PCNA during early S phase. Time points in hours after the release is indicated at the top with antibody used for Western analysis at the bottom. Loading for Western analysis is normalized for PCNA to determine DNMT1 and G9a recruitment. (C) DNMT1 knockdown impairs G9a loading and H3K9 methylation. Western blot analysis of the total cell extract (left) and chromatin (right) are shown. The antibodies used are indicated at the bottom and the siRNA-mediated knockdown of DNMT1 (DNMT1 KD) and G9a (G9a KD) are indicated at the top. Control knockdown (Control KD) was performed with siLitmus. (D) Localization profile of G9a and BrdU in DNMT1 knockdown COS-7 cells. BrdU is stained red and GFP-G9a is visualized as green. A portion of knockdown cells was used for Western blot analysis to determine the degree of knockdown for DNMT1 as shown in the right panel. Histone H3 was used as control. (E) Same as in D, except that the cells were genetically knockout for DNMT1 in HCT116 background.
Methylation-specific PCR analysis of IGS-rDNA. (Fig. 5) demonstrated that knockdown of G9a by siRNA resulted as observed before (Fig. 4C). Previously, we have though PCNA loading on IGS-rDNA remained similar loading, and decreased H3K9me2 concentration, although concentration of H3K9me2. Indeed, knockdown of impair G9a loading to IGS-rDNA, thus decreasing local formed to elucidate whether DNMT1 knockdown will tin immunoprecipitation (ChIP) experiments were per-
served (Fig. 5A, lane 1 vs. lane 3). Furthermore, chroma-
tinct unmethylated gene-specific PCR product was ob-
resulted in partial demethylation of IGS-rDNA as a dis-
ated primer will yield a PCR product. Indeed, IGS-rDNA et al. 1996). If the gene is methylated, only the methyl-
ylated or unmethylated gene-specific primers (Herman methylating status of a gene via a pair of meth-
ylated or unmethylated gene-specific primers [Herman et al. 1996]. If the gene is methylated, only the methylated primer will yield a PCR product. Indeed, IGS-rDNA is methylated and knockdown or depletion of DNMT1 resulted in partial demethylation of IGS-rDNA as a distinct unmethylated gene-specific PCR product was observed (Fig. 5A, lane 1 vs. lane 3). Furthermore, chromatin immunoprecipitation (ChIP) experiments were performed to elucidate whether DNMT1 knockdown will impair G9a loading to IGS-rDNA, thus decreasing local concentration of H3K9me2. Indeed, knockdown of DNMT1 severely down-regulated DNMT1 and G9a loading, and decreased H3K9me2 concentration, although PCNA loading on IGS-rDNA remained similar (Fig. 5B) as observed before (Fig. 4C). Previously, we have demonstrated that knockdown of G9a by siRNA resulted in global reduction of di- and trimethylation of H3K9 and coexpression of H3 and G9a in Escherichia coli resulted in H3K9me3 formation (Estève et al. 2005b). Biochemical data also suggested that G9a performs mono-, di-, and trimethylation on H3K9 [Patnaik et al. 2004; Collins et al. 2005]. Thus, a disruption of G9a loading via DNMT1 knockdown would impair H3K9 methylation. To investigate if chromatin methylation is impaired in IGS-rDNA loci, we performed Q-PCR analysis of the ChIP DNA from DNMT1 knockdown and control knockdown cells using H3K9me, H3K9me2, H3K9me3, and a control H4K20me2-specific antibody to investigate whether chromatin methylation is impaired in IGS-rDNA loci. Indeed, depletion of DNMT1 resulted in reduction of H3K9me, H3K9me2, and H3K9me3, but not for H4K20me2 (Fig. 5C) and H4K20me3 [data not shown]. These results suggest that the PCNA-DNMT1 binary complex guides G9a for H3K9 methylation on IGS-rDNA loci, and loss of DNMT1 impairs this loading.

**G9a and DNMT1 stimulate each other’s enzymatic activity**

The colocalization of SUV39H1 or G9a with DNMT1 coupled with their direct interactions in pull-down and immunoprecipitation assays suggested two possibilities: that the histone and DNA methylases helped each other bind to the newly replicated chromatin, and that they influenced each other’s catalytic activity. If interaction of DNMT1 with G9a or SUV39H1 stimulated its activity via a conformational change, then the effect should be evident on naked DNA templates. Figure 6A shows the results of a DNA methyltransferase activity assay where unmethylated or hemimethylated DNA was incubated with DNMT1 alone or in the presence of SUV39H1 and/or G9a. Remarkably, both de novo and maintenance DNA methylation, on unmethylated or hemimethylated substrates, respectively, were stimulated only in the presence of G9a. Importantly, a deletion but catalytically active mutant of G9a [delG9a] [Estève et al. 2005b] lacking the N-terminal 633 amino acids [DNMT1-binding region] was unable to activate DNMT1 activity [Fig. 6A]. SUV39H1 either alone or with G9a did not affect DNMT1 activity. These data suggest that the association of G9a strongly stimulates DNMT1 activity, implying that the interaction has functional consequences.

Unlike DNMT1, the methylation activity of G9a was only modestly stimulated by full-length DNMT1 [~25%] on nucleosomes isolated from HCT116 cells, and this modest stimulation was abolished if a catalytically active deletion mutant of DNMT1 [delDNMT1], lacking the N-terminal 501 amino acids [DNMT1-binding domain] was substituted for the full-length DNMT1 in the assay [Fig. 6B]. A slightly higher methylation (~35%) was observed with nucleosomes assembled in vitro with recombinant histones [data not shown]. If G9a stimulate DNMT1 activity, loss of G9a may result in loss of DNA methylation in IGS-rDNA. To examine this hypothesis, MSP assays were performed on G9a knockdown HCT116 genomic DNA. Indeed, a small but significant amount of demethylation was observed (Supplementary Fig. 3a).
Subsequently, bisulphate sequencing of IGS-rDNA was conducted on DNA derived from control knockdown, 5 AZC-treated, and G9a knockdown cells. In the control knockdown genomic DNA, the CpG site was methylated in all of the clones corresponding to the MSP primer-binding site as compared with ∼60% for G9a knockdown or 5 AZC-treated genomic DNA (Supplementary Fig. 3b). These data suggest that DNMT1 may aid histone methylation catalyzed by G9a on nucleosomal substrates during DNA replication (Fig. 2A) and G9a may aid DNMT1 activity on certain loci.

A second prediction of the interaction between G9a and DNMT1 is that it should aid binding of DNMT1 to chromatin substrates. To test this idea, chromatin was assembled using recombinant histones and attached to streptavidin beads. The immobilized chromatin was incubated with G9a or delG9a in the presence of AdoMet. The subsequent incubation was carried out with DNMT1 and 3H-AdoMet as shown. Fluorography of the methylated DNA is shown at the top and Western blot of G9a loaded onto the chromatin is shown at the bottom with lane numbers.

and this enhanced recruitment was abolished if delG9a was used (Fig. 6C, lanes 1–6, bottom). We conclude that the interaction between full-length G9a and DNMT1 leads to enhanced recruitment of DNMT1 and improved DNA methyltransferase activity in vitro, demonstrating the functional significance of the complex. Additionally, the decreased affinity of DNMT1 for chromatin pretreated with G9a and AdoMet (Fig. 6C, lanes 1–3) could be due to rapid turnover of DNMT1 on substrate.

To validate the idea that G9a can stimulate DNA methylation catalyzed by DNMT1 on chromatin in a replication-independent manner, we prebound G9a onto the chromatin array in the presence or absence of cold AdoMet. The subsequent incubation was carried out with DNMT1 and 3H-AdoMet as shown. Fluorography of the methylated DNA is shown at the top and Western blot of G9a loaded onto the chromatin is shown at the bottom with lane numbers.
DNA band was almost invisible after a short exposure (Fig. 6D, lane 1). However, in the presence of prebound G9a without AdoMet, DNA methylation on the chromatin was increased significantly (Fig. 6D, lane 2). The presence of unlabeled AdoMet during prebinding of full-length G9a enhanced DNMT1 activity severalfold as compared with G9a alone (Fig. 6D, lane 3 vs. lane 2) suggesting either an AdoMet-induced conformational change of G9a facilitated DNMT1 catalysis or that methylated H3K9 stimulated DNMT1 catalysis. To investigate whether G9a remains bound to the chromatin array during the time course of the experiment, especially in the presence of AdoMet, we Western blotted identical sets of experiments and probed the bound G9a with anti-G9a. Indeed, the majority of G9a remained bound to the chromatin in the presence of AdoMet (Fig. 6D, bottom) coinciding with enhanced DNA methylation.

**Discussion**

The genome and epigenome are copied at the replication fork. Therefore, events occurring during S phase provide a window of opportunity for eliciting genetic changes or maintaining the existing genetic state. The mechanisms underlying the re-establishment of the epigenetic state are not well understood. Based on our results, DNMT1 binds to G9a and SUV39H1 directly in cells and in vitro. A key question is understanding the precise timing of DNMT1 and G9a recruitment. Our data suggest that the context determines the timing. For example, DNMT1 copurifies with polymerase (POLα), supporting a model of concomitant DNA replication and maintenance methylation (Vertino et al. 2002). In such a case, the recruitment of DNMT1 may assist G9a. In the current study, we showed that purified chromatin from DNMT1 knockdown cells displayed a significantly reduced amount of G9a loading relative to HeLa wild-type cells, indicating that DNMT1 participates in chromatin loading of G9a. Importantly, the binary complex of DNMT1 and G9a may aid the maintenance DNA methylation process as observed in our in vitro assay. Stimulation of DNMT1 activity is particularly important during DNA replication since biochemical studies suggest that DNMT1 displays low catalytic activity in vitro (Pradhan et al. 1999). Additionally, DNMT1 may reciprocally aid H3K9 methylation by G9a as indicated in our in vitro assays. The colocalization of DNMT1 and G9a by PCNA immunoprecipitation and immunocytochemical analysis places both proteins in close proximity during S phase, validating the biochemical data, suggesting that H3K9 and CpG methylation are concurrent events. However, from the PCNA immunoprecipitation data we cannot conclude that DNMT1–G9a binary complex is directly recruited by PCNA.

The binding of DNMT1, G9a, and SUV39H1 may also follow a sequential path with DNMT1 binding first to form a platform to recruit the histone methylases. Based on the known ability of PCNA to load DNMT1, we propose it would bind first during DNA replication and then recruit G9a or a preformed PCNA–DNMT1–G9a complex. The DNMT1–G9a binary complex is catalytically competent and can perform both histone and DNA methylation. Concurrent with maintenance DNA methylation, H3K9 residues get predominately dimethylated. Either DNMT1 or G9a can be released first from the methylated chromatin. Furthermore, we have demonstrated that DNMT1 interacts directly with G9a and SUV39H1 by coprecipitation from cell extracts and by GST pull-down assays. Detailed analysis of the interaction in vitro on DNA and nucleosomal substrates indicates that the complexes are functional. G9a stimulates DNMT1 activity, while DNMT1 has a modest effect on G9a. Furthermore, knockdown of G9a resulted in DNA methylation impairment on IGS-rDNA loci. An important prediction of such a complex is that one partner should help the other bind chromatin cooperatively. Indeed, prebound G9a significantly stimulates DNMT1 binding to chromatin. Additionally, the complexes colocalize in the cell and at replication foci during S phase and participate in chromatin methylation of IGS-rDNA. A unifying model for these data is presented in Figure 7.

A related hypothesis has been proposed previously with CAF-I (chromatin assembly factor-I), MBD1 (methyl DNA-binding protein 1) and a histone methyltransferase, SETDB1 complex (Sarraf and Stancheva 2004). The trimeric CAF1-MBD1 and SETDB1 complex is associated with histones H3 and H4 at S phase, implying that methylation of H3K9 may occur during replication. Immunocytochemical detection of mono-, di-, and trimethylated Lys 9 of histone H3 using highly specific antibodies demonstrated that H3K9me1 is distinct at early DNA synthesis, followed by accumulation of H3K9me2 at the nuclear and nucleolar periphery at the mid S phase (Wu et al. 2005). These observations support the presence of G9a during DNA synthesis, since G9a is the predominant enzyme for euchromatic H3K9 methylation. Thus, before the packaging of the chromatin to higher order structures, chromatin modification must occur. Heterochromatin assembly also involves HP1, which binds to H3K9 di- and trimethylated (Jacobs et al. 2004; Wallace and Orr-Weaver 2005). HP1 interacts with p150, the large subunit of CAF-I, suggesting that CAF-I may participate in HP1 redistribution on heterochromatin during replication (Murzina et al. 1999). Furthermore, the presence of G9a-GLP/Eu-HMTase heteromeric complex may aid in H3K9 methylation (Tachibana et al. 2005). However, the precise role of Eu-HMTase in chromatin methylation during cell division is yet to be established. The functional role of other H3K9 methyltransferases, such as Eu-HMT and SETDB1, do not complement with G9a, as genetic deletion of G9a leads to embryonic lethality (Tachibana et al. 2002). It is possible that each histone methyltransferase, such as Eu-HMTase and G9a, may have different chromatin segments as substrate for methylation that needs further investigation. Thus, the role of chromatin assembly factors, histone modification enzymes, and DNA methyltransferases is to participate in the reinstatement of heterochromatin at the replication fork during S phase via
chromatin modification. However, many of these cooperative functions are likely to be required outside S phase for maintenance of heterochromatin. A recent study by Okuwaki and Verreault (2004) demonstrated that DNMT1 is capable of de novo methylation of nucleosomal DNA of 5S rRNA genes. This demonstrates a new capability of DNMT1 outside S phase. Similarly, Easwaran et al. (2004) have demonstrated replication-independent chromatin loading of DNMT1 during G2 and M phases of mammalian cells to constitutive heterochromatin. This observation has implications in chromatin silencing. The recruitment of G9a to DNMT1 perhaps coordinates histone and DNA methylation long after DNA replication. Furthermore, due to the low catalytic turnover of DNMT1 (Pradhan et al. 1999) stimulation by G9a may aid DNA methylation. This hypothesis is evident in genetic knockout of G9a, resulting in reduction of H3K9 methylation in Prader-Willi syndrome imprinting center and loss of maternal-specific CpG methylation of Prader-Willi syndrome imprinting center in mouse ES cells [Xin et al. 2003]. These findings support a possible role for both G9a and DNMT1 in the establishment of histone and DNA methylation.

In summary, we show that two global epigenetic enzymes, DNMT1 and G9a, coordinate synchronized DNA and histone H3K9 methylation during DNA replication.

Materials and methods

Cell culture and constructs
All cell lines (HeLa, HEK 293, Jurkat, and COS-7) were obtained from the American Type Culture Collection (ATCC) and were grown as per recommendations. Parental HCT116 and DNMT knockdown cell lines (HCT116 DNMT1<sup>−/−</sup>, Clone 1C1) were grown as described previously [Rhee et al. 2000]. Cells were synchronized using double-thimidine block, followed by aphidicolin treatment according to O’keefe et al. [1992].

All GST-DNMT1 constructs have been described previously [Kim et al. 2002], as well as the GST-G9a constructs [Estève et al. 2005b]. Details of the fusion constructs of G9a and DNMT1 with GFP or DsRed are available upon request.

Immunoprecipitation, GST pull-down, and Western blot analysis
Antibody for histone H3 was from Cell Signaling Technology. Anti-DNMT1 and G9a antibodies were from New England Bio-labs, and BD Pharmingen, respectively. Antibodies anti-monok9 (#07-450), anti-dimek9 (#07-441; lot: 27271), or -trimeK9 (#07-442; lot: 24416) of histone H3; anti-dimeK20 (#07-367) and anti-trimeK20 (#07-749) of histone H4; PCNA; and anti-SUV39H1 were from Upstate Biotechnology. Anti-GFP antibody was purchased from Roche. Nuclear extracts were made as described previously [Andrews and Faller 1991]. Coimmunoprecipitation and GST pull-downs were performed as described previously (Kim et al. 2002) except the washes performed with 1× phosphate-buffered saline (PBS) supplemented with 500 mM NaCl. For activity assays, coimmunoprecipitation was performed in the presence of 50 mM Tris-Cl (pH 7.5) and 100 mM NaCl. Densitometric analysis was performed using NIH ImageJ.

Nuclear extracts were prepared from HCT116 wild-type and HCT116 DNMT1<sup>−/−</sup> colon adenocarcinoma cells (obtained from Dr. Bert Vogelstein, The Johns Hopkins University, Baltimore, MD). Immunoprecipitation of the nuclear extract samples were performed using G9a-specific antibody (MBL). The nuclear extracts were prepared from HEK293 cells using IP buffer. Thirty microliters of Protein G/A agarose beads were washed twice...
with IP buffer and equilibrated using the same buffer. Nuclear extract protein equivalent to 1 mg was added to agarose beads (MBL), and then 5 µg of antibody specific to G9a was added. Normal mouse IgG [Santa Cruz Biotechnology] was used to immunoprecipitate the same amount of nuclear extract as a negative control. Samples were rotated overnight at 4°C. Agarose beads were washed three times with 500 µL of IP buffer. The protein complex was eluted in 2× SDS loading buffer (50 µL) at 70°C for 10 min. The supernatant was collected and analyzed by Western blot using DNMT1 antibody [Novus].

**Gel filtration chromatography**

Gel filtration of nuclear extract was performed on HiPrep 26/60 Sephacryl S-200 HR [GE Healthcare Life Sciences] equilibrated in 10 mM Tris (pH 7.5), 1 mM EDTA, and 500 mM NaCl using an FPLC apparatus [GE Healthcare Life Sciences]. Three milliliters of nuclear extract were loaded (60 µg) onto the column at a flow rate of 0.5 mL/min with 3-ML fractions collected. Additional runs using proteins from HMW Gel Filtration Calibration Kit [GE Healthcare Life Sciences] were used to determine the molecular weight of the eluted nuclear extract peaks. Blue dextran was used for determination of the void volume of the column. Ferritin (449 kDa) and aldolase (158 kDa) were used for gel filtration profile monitoring.

**DNA and histone methyltransferase assay**

DNMT1, MBP-delG9a [delG9a], and G9a were obtained from New England Biolabs. SUV39H1 was cloned into pVic1 and purified as described by Chin et al. (2006). DNA methyltransferase assays were carried out at 37°C for 30–60 min in duplicate with a total volume of 25 µL of reaction mix as previously described [Pradhan et al. 1999]. SNRPN exon 1 duplex DNA (4 µM) was used for methylation assays with 25 nM DNMT1, 40 nM G9a, 40 nM delG9a [MBP-delG9a], and 40 nM SUV39H1. The reactions were spotted on DE81 [Millipore] membrane and processed as described previously [Pradhan et al. 1999]. The histone methyltransferase assay was performed with a synthetic H3 tail representing the first 17 amino acids along with an N-terminal cys (C-ARTKQTARKSTGGKAPR) as described previously [Pataki et al. 2004]. Methylation of chromatin by G9a and DNMT1, 2 U of DNase I [New England Biolabs] was used to destroy methylated DNA.

**Immunocytochemistry**

COS-7 and HeLa cells were cultured on coverslips and respectively transfected with a mixture of plasmid and FuGENE6 (Roche) or Lipofectamine 2000 (Invitrogen) at a ratio of 3 µL/µg. The cells were visualized after 48 h using a Zeiss 200M microscope with a 63× oil objective lens at 488 nm for GFP-DNMT1, GFP-G9a, and GFP-SUV39H1 fusion proteins, 568 nm for DsRed-DNMT1 and DsRed-G9a detection; and 460 nm for GFP-G9a, and GFP-SUV39h1 fusion proteins; 568 nm for GFP-DNMT1 antibody (Novus). HeLa cells were washed twice in PBS (pH 7.4) and scrapped in 15 mL of PBS. After centrifugation [1500 rpm, 5 min at 4°C], the supernatant was discarded and the cells were resuspended in 10 mL of buffer A [15 mM Tris-CI at pH 7.5, 5 mM MgCl2, 60 mM KCl, 0.5 mM DTT, 15 mM NaCl, 300 mM sucrose]. Ten milliliters of buffer A containing 1% NP-40 was added and mixed by inverting. After 10 min on ice, the nuclei were pelleted and resuspended in 5 mL of ice-cold buffer A. After centrifugation, the pellet was resuspended in buffer B [15 mM Tris-CI at pH 7.5, 1 mM CaCl2, 60 mM KCl, 0.5 mM DTT, 15 mM NaCl, 300 mM sucrose] and 500 U of micrococcal nuclease (Roche) were added for 1.5 h at 4°C. The reaction was stopped by adding 50 mM EDTA. The nuclei were pelleted, washed with ice-cold PBS, and then resuspended in 0.5 mL of buffer C [1% SDS, 10 mM EDTA, 50 mM Tris-CI at pH 8]. Thirty micrometers of protein was loaded on SDS-polyacrylamide gels for Western blot analysis. All buffers contained PMSF and protease cocktail inhibitors (Sigma).

**Binding and chromatin array recruitment assay**

Chromatin arrays were prepared and biotinylated as described previously [Ikeda et al. 1999]. Chromatin was prepared using recombinant Xenopus histones assembled into octamers and bound to arrays by step-wise salt dialysis as described previously [Logie and Peterson 1997; Luger et al. 1999]. After salt dialysis, chromatin was immobilized on M280 streptavidin beads [DYNAL] in chromatin-binding buffer (300 mM KCl, 10% glycerol, 4 mM MgCl2, 1 mM DTT, 200 µg/mL BSA). Immobilized chromatin templates were validated by limited MNase digestion and native PAGE of EcoRI-digested fragments as described previously [Ikeda et al. 1999]. G9a was bound to the chromatin array in reaction mix as described above, in the presence or absence of the cofactor AdoMet for 30 min at room temperature. After washing to remove unbound proteins, the templates were incubated with increasing amounts of DNMT1 at 37°C in reaction buffer as previously described above. After washing, the templates were analyzed for DNMT1 recruitment by SDS-PAGE gel electrophoresis and immunoblotting. For fluorography analysis of 3H-methylated DNA, the films were exposed at ~80°C for 5 d.

**Gene knockdown, methylation-sensitive PCR, and ChiP assay**

HeLa cells were transfected twice using RNAiFECT reagent (Qiagen) with 15 nM of siRNA mixes The mixes used were specific for G9a or DNMT1 [New England Biolabs]. For control transfection, litmus siRNA was used [New England Biolabs]. Seventy-two hours after the second transfection, ChiP or MSP were performed [Estève et al. 2005a]. For each IP, 5 µg of purified rabbit IgG [Calbiochem], anti-PCNA [Upstate Biotechnology], anti-DNMT1 [New England Biolabs], anti-G9a [Upstate Biotechnology], anti-H3K9me2, anti-H3K9me, anti-H3K9me3, anti-H4K20me2, and anti-H4K20me3 [Upstate Biotechnology] were used. For ChiP assay, DNA was amplified using the following primers: for IG5-rDNA sense, 5'-CTCTTACACAGGACGTGAGAACG-3; antisense, 5'-TCGACCTCCGCAATCGTACA-3. The MSP primer sets for IG5-rDNA were unmethylated gene-specific sense, 5'-ATGAGAGTGAGAAAAGT-3; unmethylated gene-specific antisense, 5'-GCTCAGAGGCATGCCAGAT-3.

**DNMT1 and G9a coordinate chromatin methylation**

DNMT1 and G9a coordinate chromatin methylation
TGTGGTTTG-3', and antisense, 5'-TCAACCTCCAAAAA TCATAACCCAAA-3'; methylated gene-specific sense, 5'-AC GAGCAGGAAGCCTTGTCG-3', and antisense, 5'- TCGACCTTCGGGAAATCTGACACCCG-3'. 5-aza-2-deoxycytidine-treated Jurkat genomic DNA was obtained from New England Biolabs. Primer for bisulphate sequencing are available upon request.

For quantitative ChIP assay, IGS-rDNA quantification was performed by real-time PCR, in a Bio-Rad iCycler using SYBR Green Supermix (Bio-Rad). The SYBR green incorporation into double-stranded DNA was measured during each elongation step. Relative values were calculated using actin as an endogenous control. The percentage of DNA remaining upon the IP step. Relative values were calculated using actin as an endogenous control. The percentage of DNA remaining upon DNMT1 knockdown was calculated with respect to the mock-treated cells [Litmus siRNA] using the ΔCt method. Primer sets are available upon request. Error bars on the graph represent the standard deviation of triplicates.

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Direct interaction between DNMT1 and G9a coordinates DNA and histone methylation during replication

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