Influence of DNA methylation on positioning and DNA flexibility of nucleosomes with pericentric satellite DNA

Akihisa Osakabe1,†, Fumiya Adachi1,†, Yasuhiro Arimura1, Kazumitsu Maehara2, Yasuyuki Ohkawa3 and Hitoshi Kurumizaka1

1Laboratory of Structural Biology, Graduate School of Advanced Science and Engineering, Waseda University, 2-2 Wakamatsu-cho, Shinjuku-ku, Tokyo 162-8480, Japan
2Department of Advanced Medical Initiatives, Faculty of Medicine, Kyushu University, Fukuoka 812-8582, Japan

DNA methylation occurs on CpG sites and is important to form pericentric heterochromatin domains. The satellite 2 sequence, containing seven CpG sites, is located in the pericentric region of human chromosome 1 and is highly methylated in normal cells. In contrast, the satellite 2 region is reportedly hypomethylated in cancer cells, suggesting that the methylation status may affect the chromatin structure around the pericentric regions in tumours. In this study, we mapped the nucleosome positioning on the satellite 2 sequence in vitro and found that DNA methylation modestly affects the distribution of the nucleosome positioning. The micrococcal nuclease assay revealed that the DNA end flexibility of the nucleosomes changes, depending on the DNA methylation status. However, the structures and thermal stabilities of the nucleosomes are unaffected by DNA methylation. These findings provide new information to understand how DNA methylation functions in regulating pericentric heterochromatin formation and maintenance in normal and malignant cells.

1. Introduction

DNA methylation is an important epigenetic mark that regulates the formation of chromatin domains, such as heterochromatin [1–5]. In mammals, DNA methylation occurs in the CpG dinucleotide and is considered to affect the structure and stability of the nucleosome, which is the basic architecture in chromatin [6–10]. In the nucleosome, about 150 base pairs of DNA are left-handedly wrapped around the histone octamer, composed of two each of the core histones H2A, H2B, H3 and H4 [11–13].

DNA methylation is reportedly correlated with nucleosome positioning in plant and mammalian genomes [14,15]. The genomic DNA regions with high CpG content are known as CpG islands, and the CpG methylation apparently plays pivotal roles in gene regulation and genomic DNA maintenance [4,16,17]. Abnormal DNA methylation statuses have been detected in various cancer cells [18,19]. CpG islands are mostly hypomethylated in normal cells, but are hypermethylated in cancer cells, especially in the promoters of tumour suppressor genes [4,20,21]. In contrast, large-scale CpG island demethylation has been detected at the tissue-specific gene promoters in lung cancers [22]. These previous findings suggested that DNA methylation functions in proper gene expression and genomic DNA stability [23,24].

Heterochromatin instability in pericentromeric satellite regions has also been detected as an early and frequent event during human carcinogenesis [25]. Interestingly, this heterochromatin instability occurs concomitantly with the hypomethylation of the CpG sites on the satellite DNA [25–28]. However, the
2. Results

2.1. Nucleosome formation on the human satellite 2 sequence

We first prepared a 160 base-pair human satellite 2 DNA fragment. This satellite 2 fragment contained seven CpG sites, TTCGAT, TTCGAT, TTCGAT, TCCGAG, TTCGAT, TCCGAT and TCCGAG (from 5’ to 3’), which are potentially methylated in normal cells (figure 1a, upper panel). To ensure that these CpG sites are fully methylated, all of the CpG sites were replaced by TCCGAA, which can be cleaved by the restriction enzyme BstBI (figure 1a, lower panel). In this study, this satellite 2 derivative was named Sat2. As shown in figure 1b (lane 1), all of the CpG sites in the Sat2 160 base-pair fragment were digested by BstBI. As anticipated, the BstBI cleavage was completely inhibited when the Sat2 160 base-pair fragment was treated with the DNA methyltransferase M.SssI (figure 1b, lane 2), indicating that all seven CpG sites of Sat2 were fully methylated.

We then reconstituted the nucleosomes with methylated or unmethylated 160 base-pair Sat2 DNA fragments, by the salt dialysis method. The reconstituted nucleosomes were treated with microccocal nuclease (MNase), which preferentially cleaves the linker DNA segments detached from the histone surface, and the resulting approximately 145 base-pair DNA fragments were purified (figure 1c, lanes 4 and 5). We then performed massively parallel sequencing (deep sequencing) with these MNase-treated DNA fragments and found one major (right, denoted as R) and two minor (centre and left, denoted as C and L, respectively) nucleosome positions on the Sat2 sequence (figure 1d,e). The major R position was mapped on the right edge of the Sat2 DNA fragment, and the minor C and L positions were shifted by about 7 and 13 base pairs from the right edge, respectively (figure 1d). In both the methylated and unmethylated Sat2 DNAs, about 70% of the nucleosomes were formed at the R position, although a slight decrease was observed with the methylated Sat2 (figure 1e). Similarly, upon the DNA methylation, the nucleosome population at the C position was decreased (figure 1e). In contrast, the population of the L position was increased 1.5-fold when the methylated Sat2 was used as the substrate (figure 1e).

2.2. Crystal structures of the nucleosomes containing the methylated Sat2R and Sat2L DNAs

We crystallized the nucleosomes containing the methylated Sat2L (145 base pairs) and Sat2R (146 base pairs) DNA fragments and determined their structures at 2.63 Å and 3.15 Å resolutions, respectively (table 1 and figure 2a,b). For a reference, we also determined the structure of the nucleosome containing the unmethylated Sat2R DNA sequence at 2.90 Å resolution (table 1 and figure 2c). The histone octamer structures in the nucleosomes containing the methylated Sat2R and Sat2L DNAs were the same as that in the nucleosome containing the unmethylated Sat2R DNA (figure 2a–c). In addition, the DNA binding path in the methylated Sat2R nucleosome was not different from that in the unmethylated R nucleosome (figure 2d). The DNA binding path in the methylated Sat2L nucleosome was also the same as that in the unmethylated Sat2R nucleosome (figure 2e). Therefore, these results indicate that the hypermethylation at the seven CpG positions of the Sat2 DNA does not affect the intrinsic DNA wrapping property of the histone octamer.

Since the nucleosomes were packed in a nested manner in the crystals, the additional methyl groups of the 5-methyl-cytosines were not visible in these nucleosome structures. Therefore, we mapped the 5-methyl-cytosine locations on these nucleosome structures in two nested orientations (figure 3a,b,c). Six out of the seven CpG sites were incorporated into each Sat2R or Sat2L nucleosome (figure 3c). Interestingly, in the Sat2R nucleosome, most of the 5-methyl-cytosines tended to be exposed to the solvent (figure 3c). In contrast, two 5-methyl-cytosines are buried in the histone–DNA contact surface in the Sat2L nucleosome (figure 3c). These structural differences may affect the accessibility of the methyl-DNA binding proteins to the nucleosomal 5-methyl-cytosine [31].

2.3. DNA methylation changes the accessibility of the DNA ends of the nucleosome, without affecting its thermal stability

We next tested the MNase sensitivity of the nucleosomes containing the methylated and unmethylated DNAs with two R and L positions. To do so, four types of nucleosomes, containing methylated Sat2R (146 base pairs), unmethylated Sat2R (146 base pairs), methylated Sat2L (145 base pairs), and unmethylated Sat2L (145 base pairs) DNAs, were reconstituted and purified by native polyacrylamide gel electrophoresis (PAGE) (figure 4a).

The quantitative MNase assay revealed that, under the unmethylated conditions, the nucleosome containing the Sat2L DNA was quite susceptible to MNase, as compared to the nucleosome containing the Sat2R DNA (figure 4b,c). DNA methylation drastically reduced the MNase susceptibility of the Sat2L nucleosome (figure 4b,c). In contrast, the MNase susceptibility of the Sat2R nucleosome was enhanced upon DNA methylation (figure 4b,c). In nucleosomes, MNase is known to preferentially degrade the DNA segments that are detached from the histone surface. We confirmed that MNase equally degraded the non-nucleosomal Sat2L and Sat2R DNAs (figure 4d), indicating that the enzyme did not exhibit any sequence specificity to these DNAs. In addition, the DNA methylation did not affect the MNase susceptibility of the non-nucleosomal Sat2L and Sat2R DNAs (figure 4d). Therefore, these data indicate that DNA hypermethylation enhances the DNA end flexibility of the Sat2R nucleosome, but reduces that of the Sat2L nucleosome.

Since the thermal stabilities of these four nucleosomes were exactly the same, the differences in their MNase susceptibilities were not due to changes in the nucleosome stability upon DNA methylation (figure 4e). In this thermal stability
assay, nucleosome disruption by heating was monitored as histone dissociation from the nucleosome, by using SYPRO Orange, a fluorescent dye that specifically binds to denatured proteins, as a probe. Therefore, DNA hypermethylation influenced the DNA end flexibility of the nucleosomes without affecting their thermal stabilities, and this may depend on the translational positioning of the nucleosomes.

3. Discussion

The human satellite 2 repeats located in pericentric heterochromatin regions are highly methylated in normal cells, but are reportedly hypomethylated in cancer cells [32–34]. However, the question remained as to whether the DNA methylation status affects the structure and stability of the nucleosome on the satellite 2 sequence. To answer this question, we reconstituted the satellite 2 nucleosomes with or without DNA methylation and studied the impacts of the DNA methylation on the positioning, structure, stability and DNA end flexibility of the nucleosomes.

We identified the major and minor nucleosome positions on the satellite 2 sequence (figure 1d,e). We found that the nucleosome population of the minor position (L) significantly increased upon DNA methylation (figure 1d,e). It is intriguing that the Sat2L nucleosome was more easily degraded.
by MNase in the absence of DNA methylation, without affecting the nucleosome structure and stability (figures 2–4). Therefore, the methylation of the satellite 2 DNA may function to accommodate the DNA ends of the Sat2L nucleosome more tightly. This is consistent with a previous report that DNA methylation reportedly facilitates the wrapping of DNA ends [6,8,35].

However, the DNA methylation oppositely affected the DNA end flexibility of the major Sat2R nucleosome. This is consistent with the results reported by Jimenez-Useche & Yuan [7], who found that the DNA methylation does not compact the nucleosomal DNA [7]. Therefore, the previous controversial observations regarding whether the DNA methylation reduces or enhances the nucleosomal DNA end flexibility may be reconciled, by considering the translational positions of the nucleosome. Consistent with this idea, DNA methylation reportedly affects the DNA end flexibility differently, depending on the nucleosomal locations of the CpG dinucleotides [9].

We found that the DNA end flexibilities of the Sat2R and Sat2L nucleosomes became similar when the satellite 2 DNA was fully methylated. This finding suggests that DNA methylation may reduce the differences in the nucleosome characteristics and may function to facilitate well-organized, regular chromatin folding in heterochromatin.

In this study, we determined the physical characteristics of satellite 2 nucleosomes with or without DNA methylation. Our results have led to a new question: how are these structural and physical characteristics of the satellite 2 nucleosomes with or without DNA methylation linked to the chromosome instability frequently observed in cancer cells? Further cell-biological and genetic studies are awaited.

4. Material and methods

4.1. Purification of recombinant human histones

Human histones H2A, H2B, H3 and H4 were purified by the method described previously [36–39]. In this method, bacterially expressed human histones with an N-terminal His$_6$-tag were purified with Ni-NTA agarose (Qiagen). After removal of the His$_6$-tag portion by the addition of thrombin protease (1 unit mg$^{-1}$ protein), the histones were further purified with MonoS column chromatography (GE Healthcare), freeze-dried and stored at 4°C.

The freeze-dried histones (1 : 1 : 1 : 1 stoichiometry) were dissolved in 20 mM Tris–HCl buffer (pH 7.5), containing 7 M guanidine hydrochloride and 20 mM 2-mercaptoethanol. The sample was dialysed against 10 mM Tris–HCl buffer (pH 7.5), containing 2 M NaCl, 1 mM EDTA and 5 mM 2-mercaptoethanol, and the resulting histone octamers were purified by Superdex200 (GE Healthcare) gel filtration column chromatography.

4.2. Preparation of Sat2 DNA fragments for nucleosome reconstitution

Four 160 base-pair Sat2 DNA fragments, each bearing seven BstBI (New England BioLabs) recognition sites, were inserted

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Table 1. Data collection and refinement statistics (molecular replacement).

|                         | unmethylated Sat2R nucleosome | methylated Sat2R nucleosome | methylated Sat2L nucleosome |
|-------------------------|-------------------------------|-------------------------------|-----------------------------|
| resolution range (Å)    | 50 – 2.90                    | 50 – 3.15                    | 50 – 2.63                   |
| space group             | P2$_1$,2,2$_1$               | P2$_1$,2,2$_1$               | P2$_1$,2,2$_1$              |
| cell parameters         | a = 105.431 Å; b = 109.331 Å; c = 175.771 Å; α = 90.0°; β = 90.0°; γ = 90.0° | a = 103.452 Å; b = 108.990 Å; c = 173.446 Å; α = 90.0°; β = 90.0°; γ = 90.0° | a = 105.197 Å; b = 109.297 Å; c = 173.686 Å; α = 90.0°; β = 90.0°; γ = 90.0° |
| total number of reflections | 44 980 | 34 975 | 59 234 |
| Rmerge (%)              | 9.0 (48.6)                   | 8.7 (48.3)                   | 7.1 (48.9)                  |
| completeness (%)        | 98.7 (97.5)                  | 99.3 (98.3)                  | 98.8 (97.7)                 |
| l/cr (l)                | 12.6 (2.2)                   | 11.6 (2.4)                   | 14.1 (2.3)                  |
| redundancy              | 5.0 (3.5)                    | 5.9 (4.2)                    | 5.3 (3.5)                   |
| refinement              |                               |                               |                             |
| resolution (Å)          | 37.9 – 2.90                  | 24.9 – 3.15                  | 19.9 – 2.63                 |
| Rwork/Rfree (%)         | 25.12/29.49                  | 23.52/29.55                  | 22.38/28.42                 |
| r.m.s.d. bonds (%)      | 0.010                        | 0.010                        | 0.010                       |
| r.m.s.d. angles (%)     | 1.158                        | 1.331                        | 1.369                       |
| Ramachandran plot       |                               |                               |                             |
| most favoured (%)       | 97.96                        | 96.58                        | 97.95                       |
| allowed (%)             | 2.04                         | 3.42                         | 2.05                        |
| disallowed (%)          | 0                            | 0                            | 0                           |
| PDB code                | 5CPI                         | 5CPJ                         | 5CPK                        |

$^a$Rmerge = $\sum_{hkl} \sum_{i} |I_i(hkl) – \langle I(hkl) \rangle|/\sum_{hkl} \sum_{i} I_i(hkl)$.

$^b$Rwork = $\sum_{hkl} ||F_{obs}|| – |F_{calc}||/\sum_{hkl} ||F_{obs}||$. Rfree was calculated with 5% of the data excluded from the refinement.
into the pGEM-T Easy vector (Promega). The plasmid was amplified in *Escherichia coli* cells and was purified by the method described previously [40]. The 160 base-pair Sat2 DNA fragment was isolated from the plasmid by digestion with *EcoR*V. The vector DNA portion was removed by PEG-6000 precipitation, and the 160 base-pair Sat2 DNA fragment was then purified by chromatography on TSKgel DEAE-5PW (TOSOH). For the Sat2L and Sat2R DNA fragments, eight Sat2L (145 base pairs) or Sat2R (146 base pairs) DNA fragments were tandemly ligated into the pGEM-T Easy vector. The DNA fragments were purified by the same methods as described above.

The DNA sequences of Sat2L and Sat2R were as follows. Sat2L: 5′−ATCAT TTCCA TTCGA AGATT CCATT CGAAT CCATT CCAAA ATGAT TACAT TCGAA TCCAT TCGAA GATTC CATTT GAGCC TGTTC GAAAA TTCCA TTTGA GTCCA ACCAA TGATT CCATT CATTT CCATT CAATG ATTCC ATGAT−3′

Sat2R: 5′−ATCAG ATTCC ATTCG AATCC ATTCG AAAAA TATTA CATTC GAATC CATTC GAAGA TTCCA TTGGA GCCTG TTCGA AAATT CCATT TGAGT CCAAC CAATG ATTCG ATTCG ATTCG ATTCG ATTCG TCCAT TTGGA T−3′

CpG methylation was introduced by an incubation with the *bacterial* DNA methyltransferase *M.SssI* (New England BioLabs), in the presence of 160 μM S-adenosylmethionine (2 units μg⁻¹ DNA) at 37°C for 16 h. The reaction was terminated by an incubation at 65°C for 30 min. The unmethylated satellite 2 DNA was cleaved with *BstBI* (10 units μg⁻¹ DNA) at 65°C for 4 h, and the resulting methylated Sat2 DNA was purified by chromatography on TSKgel DEAE-5PW.

### 4.3. Reconstitution of nucleosomes containing Sat2 DNA fragments

Nucleosomes containing the Sat2 DNA fragments were reconstituted by the salt dialysis method, as described previously [37–39]. DNA fragments were mixed with histone octamers in the presence of 2 M KCl. The KCl concentration was gradually reduced from 2 M to 0.25 M, using a peristaltic pump. The reconstituted nucleosomes were incubated at 55°C for 2 h and were further purified by non-denaturing PAGE, using a Prep Cell apparatus (Bio-Rad).

### 4.4. Deep sequencing analysis of the nucleosome positioning

Purified nucleosomes, containing the unmethylated or methylated Sat2 DNA fragment, were treated with MNase
The library was prepared using an NEBNext Ultra DNA Library Prep Kit and was then sequenced using an Illumina HiSeq 1500 system (Illumina K.K.; USA). The sequenced reads were uniquely mapped onto the target DNA sequence, using the BOWTIE 2 program (v. 2.2.2) with default parameters. The proportions of the mapped reads of the nucleosome dyad position, which was estimated as the position shifted by 73 base pairs from the 5'-end of the reads, on the target Sat2 DNA (1–160 base pairs), were calculated.

4.5. Thermal stability assay for nucleosomes

The nucleosome stability was monitored by a thermal stability assay, as described previously [41–43]. Purified nucleosomes (1.1 μM) were mixed with SYPRO Orange dye (Sigma-Aldrich) in 20 mM Tris–HCl buffer (pH 7.5), containing 1 mM DTT. The SYPRO Orange fluorescence was detected with a StepOnePlus™ Real-Time PCR unit (Applied Biosystems), using a temperature gradient from 25°C to 95°C, in steps of 1°C min⁻¹.

4.6. Crystallization and structure determination

Purified nucleosomes containing unmethylated or methylated Sat2R DNA (146 base pairs) and methylated Sat2L DNA (145 base pairs) fragments were dialysed against 20 mM potassium cacodylate buffer (pH 6.0), containing 1 mM EDTA. The nucleosome solution (3.5 mg ml⁻¹ DNA concentration) was mixed with an equal volume of 20 mM potassium cacodylate buffer (pH 6.0), containing 50–70 mM KCl and 70–105 mM MnCl₂. The drops were equilibrated against 500 μl reservoir solution (20 mM potassium cacodylate buffer (pH 6.0), containing 35–45 mM KCl and 45–60 mM MnCl₂), and crystals were obtained by the hanging drop method. The resulting nucleosome crystals were cryoprotected by soaking in a solution containing 20 mM potassium cacodylate (pH 6.0),...
The refinement of the 145 base-pair DNA (PDB ID: 3UT9) was used as the search model for molecular replacement [30]. The refinements for the nucleosome containing some containing the methylated 145 base-pair Sat2L DNA (200 ng DNA) were incubated with MNase (0.04 units) in 10 μl of 50 mM Tris–HCl (pH 8.0) buffer, containing 2.5 mM CaCl$_2$ and 0.9 mM dithiothreitol, at 25°C for 1, 3 and 5 min. For the experiments with naked DNAs, unmethylated Sat2R, unmethylated Sat2L, methylated Sat2L, methylated Sat2R, methylated Sat2L and methylated Sat2R DNA (200 ng DNA) were incubated with MNase (0.04 units) in 10 μl of 50 mM Tris–HCl (pH 8.0) buffer, containing 2.5 mM CaCl$_2$ and 0.9 mM dithiothreitol, at 25°C for 1, 3 and 5 min. For the experiments with naked DNAs, unmethylated Sat2R, unmethylated Sat2L, methylated Sat2L, methylated Sat2R, methylated Sat2L and methylated Sat2R DNA (200 ng DNA) were incubated with MNase (0.04 units) in 10 μl of 50 mM Tris–HCl (pH 8.0) buffer, containing 2.5 mM CaCl$_2$ and 0.9 mM dithiothreitol, at 25°C for 1, 3 and 5 min. The MNase susceptibility assay with naked DNAs. Unmethylated Sat2L DNA (lanes 6–9), methylated Sat2L DNA (lanes 10–13) and methylated Sat2R DNA (lanes 14–17) were treated with MNase (0.8 units) for 0 (lanes 2, 6, 10 and 14), 1 (lanes 3, 7, 11 and 15), 3 (lanes 4, 8, 12 and 16) and 5 min (lanes 5, 9, 13 and 17). Lanes 1 and 18 indicate the 10 base-pair DNA ladder markers. (b) Graphic representation of the experiments shown in panel (a). Standard deviation values are shown (n = 3). (d) The MNase susceptibility assay with naked DNAs. Unmethylated Sat2L DNA (lanes 2–5), unmethylated Sat2R DNA (lanes 6–9), methylated Sat2L DNA (lanes 10–13) and methylated Sat2R DNA (lanes 14–17) were treated with MNase (0.8 units) for 0 (lanes 2, 6, 10 and 14), 1 (lanes 3, 7, 11 and 15), 3 (lanes 4, 8, 12 and 16) and 5 min (lanes 5, 9, 13 and 17). Lanes 1 and 18 indicate the 10 base-pair DNA ladder markers. (e) Thermal stability curves of the nucleosomes. The normalized fluorescence intensity was plotted against the temperature (from 60°C to 90°C).

Figure 4. The MNase assay. (a) Purified nucleosomes containing unmethylated Sat2L DNA (lane 5), methylated Sat2L DNA (lane 6), unmethylated Sat2R DNA (lane 7) and methylated Sat2R DNA (lane 8) were analysed by 6% non-denaturing PAGE with ethidium bromide staining. Lanes 1–4 indicate the naked DNAs of unmethylated Sat2L DNA (lane 1), methylated Sat2L DNA (lane 2), unmethylated Sat2R DNA (lane 3) and methylated Sat2R DNA (lane 4), respectively. (b) The MNase susceptibility assay with nucleosomes. Purified nucleosomes (200 ng DNA) containing unmethylated Sat2L DNA (lanes 2–5), unmethylated Sat2R DNA (lanes 6–9), methylated Sat2L DNA (lanes 10–13) and methylated Sat2R DNA (lanes 14–17) were treated with MNase (0.8 units) for 0 (lanes 2, 6, 10 and 14), 1 (lanes 3, 7, 11 and 15), 3 (lanes 4, 8, 12 and 16) and 5 min (lanes 5, 9, 13 and 17). Lanes 1 and 18 indicate the 10 base-pair DNA ladder markers. (c) Graphic representation of the experiments shown in panel (b). Standard deviation values are shown (n = 3). (d) The MNase susceptibility assay with naked DNAs. Unmethylated Sat2L DNA (lanes 2–5), unmethylated Sat2R DNA (lanes 6–9), methylated Sat2L DNA (lanes 10–13) and methylated Sat2R DNA (lanes 14–17) were treated with MNase (0.8 units) for 0 (lanes 2, 6, 10 and 14), 1 (lanes 3, 7, 11 and 15), 3 (lanes 4, 8, 12 and 16) and 5 min (lanes 5, 9, 13 and 17). Lanes 1 and 18 indicate the 10 base-pair DNA ladder markers. (e) Thermal stability curves of the nucleosomes. The normalized fluorescence intensity was plotted against the temperature (from 60°C to 90°C).
dithiothreitol, at 25°C for 1, 3, and 5 min. After the incubation, the reactions were stopped by the addition of stop solution (60 μl), composed of 20 mM Tris–HCl (pH 8.0), 20 mM EDTA, 0.25% SDS, and 0.5 mg ml⁻¹ protease K (Roche). The reaction mixtures were further incubated at 25°C for 15 min. The DNA was then extracted with phenol–chloroform, and the resulting DNA fragments were analysed by 10% non-denaturing PAGE in 0.5 × TBE buffer (45 mM Tris base, 45 mM boric acid, and 1 mM EDTA). The DNA bands were visualized by ethidium bromide staining.

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