Genome-wide genetic variations are highly correlated with proximal DNA methylation patterns

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507 Mbp (~1% of the human genome was within 10 bp of a CpG site; in these regions, the single nucleotide polymorphism (SNP) rate significantly increased by ~50% (P < 10^-66 by a two-proportion z-test) if the neighboring CpG sites are methylated. To reconfirm this finding in another vertebrate, we compared six single-base resolution methylomes in two inbred medaka (Oryzias latipes) strains with sufficient genetic divergence (3.4%). We found that the SNP rate also increased by ~50% (P < 10^-60), and the substitution rates in all dinucleotides increased simultaneously (P < 10^-48) around methylated CpG sites. In the hypomethylated regions, the “CGCG” motif was significantly enriched (P < 10^-60) and evolutionarily conserved (P = 0.203%), and slow CpG deamination rather than fast CpG gain was seen, indicating a possible role of CGCG as a candidate cis-element for the hypomethylation state. In regions that were hypermethylated in germline-like tissues but were hypomethylated in somatic liver cells, the SNP rate was significantly smaller than that in hypomethylated regions in both tissues, suggesting a positive selective pressure during DNA methylation reprogramming. This is the first report of findings showing that the CpG methylation state is significantly correlated with the characteristics of evolutionary change in neighboring DNA.

[Supplemental material is available for this article.]
**Results**

We examined the relationship between DNA methylation and genetic variation in the human genome using publicly available data. Specifically, we examined the methylome data from human germline sperm cells (Molaro et al. 2011) and the reference single nucleotide polymorphisms (refSNPs) of the CEU population collected by the HapMap Project (The International HapMap Consortium 2003). Although substantial data are available on the human genome, the incidence of genetic variation in the human genome is quite low (~0.1%), which is not sufficient to perform a detailed analysis on the dinucleotide patterns of genetic variations associated with hyper- and hypomethylated regions. For this purpose, the medaka (*Oryzias latipes*) model system provides an ideal resource in vertebrates because two medaka inbred strains, Hd-rR and HNI, have a sufficiently high incidence of genetic variation (~3.4%) (Kasahara et al. 2007). The Hd-rR and HNI medaka inbred strains originated in the southern and northern parts of Japan, respectively, which were separated by a watershed, and the strains diverged ~18 million years ago (Setiamarga et al. 2009).

Another merit of the medaka is its abundance of germline cells in the testes and blastulae (half-day embryos, in which some of the cells remain totipotent [Hong et al. 1998]), which provides information on the primary relationships between the methylation pattern and genomic variation during the course of evolution. Thus, we newly generated six methylomes of medaka genomes at single-base resolution using genomic DNA from three libraries (liver, blastulae, and testes) of the two strains, treating it with bisulphite, and subjecting it to Illumina sequencing. In total, 1.8 billion reads were produced, and 44 billion nucleotides were uniquely mapped to the genome, with an average depth of ninefold for each cell line of each strain (Supplemental Table S1). An average of ~78% of each medaka genome was covered by at least one read (Supplemental Table S1). The bisulphite conversion procedure was highly effective, as validated by a control experiment using yeast (*Saccharomyces cerevisiae* S288C) (Supplemental Table S2). In yeast, cytosine nucleotides are not methylated, and indeed, none of yeast cytosine nucleotides was uncovered to be hypermethylated (methylation level ~0.8) (Supplemental Fig. S1). Data for the entire methylome of medaka at single base resolution can be accessed at [http://utgenome.org/medaka/methylome/](http://utgenome.org/medaka/methylome/).

Substantial cytosines at non-CpG sites are methylated in human ES and *Arabidopsis* (Cokus et al. 2008; Lister et al. 2008, 2009); however, methylation at non-CpG sites was very rare in human sperm (Molaro et al. 2011) and in all of the three medaka tissues (Supplemental Fig. S2); i.e., few non-CpG cytosines (~0.02%) were observed to be hypermethylated. To examine whether cytosine methylation state, unmethylated or methylated, affects proximal genetic variation, both unmethylated and methylated cytosines should be sufficiently available. In CpG sites, the ratio of unmethylated cytosines:methylated cytosines was ~10%:~90%, showing the abundance of unmethylated cytosines. In non-CpG sites, however, the ratio was 99.98%:0.02%, indicating the lack of methylated cytosines. Thus, we focused on methylation patterns of CpG sites in subsequent analyses. The distribution of methylation levels was found to be bimodal, and the vast majority of CpG sites were highly methylated in human sperm (Molaro et al. 2011) and in each medaka library (Supplemental Fig. S2). CpG islands may be appropriate regions in which to measure the effect of methylation pattern on the incidence of genetic variation; however, they occupy a relatively small proportion of the human genome (~1%) as well as the medaka genome (~3%) and exist mostly in CpG-rich regions. Therefore, regions with fewer CpG sites are overlooked. For defining more comprehensive characteristics of CpG sites in a wider range of genome sequences, we, instead, focused on neighboring genome sequences (the 10 bases upstream and downstream) of all the CpG sites, which we hereafter refer to as “CpG site blocks” (see Methods; Supplemental Table S3). Overlapping CpG site blocks were merged, and the combined length of all the CpG site blocks was 507 Mb (~18%) in the human genome and ~188 Mb in the medaka genome, accounting for ~37% of the ~500 Mb that were aligned between the two medaka genomes. We then noticed that one-third of CpG site blocks were covered by <5 reads and were likely to lack accuracy. Thus, we imposed the stringent requirement that each CpG site had a read coverage of ≥5 (see Methods). CpG site blocks that met this condition accounted for ~323 Mb (~11%) in the human genome and ~1.39 Mb (~28%) in the medaka genome, which was sufficient for estimating features of the entire genomes (Supplemental Table S3).

We compared the incidence of genetic variations in hypermethylated regions (methylation level ≥ 0.2) with that in hypermethylated regions (methylation level ≥ 0.8). Because single-nucleotide mutations by deamination of methylated cytosines in CpG dinucleotides are dominant, to examine single-nucleotide substitution rates in this analysis, we excluded SNPs in CpG dinucleotides and SNPs in TpG/CpA dinucleotides which might make TpG/CpA be CpG. Figure 1A shows that, in human sperm, the incidence rate in hypomethylated regions in the entire genome, 0.054%, was significantly lower than that in hypermethylated ones, 0.081% ($P < 10^{-5.46}$ by a two-proportion z-test) (Supplemental Table S4). This significant increase of incidence rate in hypermethylated regions was also detected in intergenic regions ($P < 10^{-3.05}$), in exons ($P < 10^{-2.9}$), and in introns ($P < 10^{-1.51}$) (Supplemental Table S4). We also observed a similar tendency in the medaka system. For example, Figure 1B illustrates a pair of the homologous regions of the human and medaka genomes where the RPS13 gene is encoded. We saw four SNPs in the hypermethylated human region and a higher SNP rate in the hypermethylated medaka region. We will check this characteristic precisely in the entire medaka genome in what follows.

In medaka, the methylation state of each CpG site is not necessarily conserved between the two strains (Fig. 1C,D; Supplemental Table S3A), which is supported by 4.46 × 10^7 methylation polymorphisms per CG site per generation in *Arabidopsis thaliana* (Becker et al. 2011; Schmitz et al. 2011). In general, however, hypomethylated and hypermethylated regions were highly conserved between the two inbred strains (Fig. 1C,D) and were also highly correlated ($R^2 = 0.73$) between testes and blastulae (Fig. 1E). Thus, hereafter, we used hypo- and hypermethylated regions that were conserved between the two strains as the representative regions. Analysis of SNPs between Hd-rR and HNI (Sasaki et al. 2009) around CpG blocks revealed that the SNP rate of 1.81% in hypermethylated regions was significantly lower than the 2.78% rate in hypermethylated regions ($P < 10^{-2.78}$) by a two-proportion z-test (Fig. 1F; Supplemental Table S4). In addition to the entire genome, these characteristics were also significant in intergenic regions ($P < 10^{-2.170}$), in introns ($P < 10^{-2.598}$), and in exons ($P < 10^{-1.13}$) in spite of inherently strong purifying selection in exons (Fig. 1F; Supplemental Table S4). Furthermore, genome-wide regions that were differentially methylated in the two strains exhibited a SNP rate of 3.41%, which is significantly higher than the rate of 2.78% in hypermethylated regions ($P < 10^{-4.45}$) (Supplemental Table S4). This indicates that genomic regions with a relaxed selective pressure display a wider divergence in cytosine methylation.
The high divergence between the two medaka strains provided a detailed analysis of mutations of dinucleotides. Figure 1G shows that the peak substitution rates occurred in CG and TA dinucleotides. The CG substitutions (11% in hypermethylated regions) were predominantly transitions, which can be explained by the deamination of methylated cytosines that dominates point substitutions in vertebrates (Lindahl and Nyberg 1972; Cooper and Krawczak 1989), whereas in hypomethylated regions, the CG substitution rate was moderate. Another commonly mutated dinucleotide, TA, is universally underrepresented because of its high mutation rate (Ohno 1988; Burge et al. 1992). Our novel finding is that substitution rates for all dinucleotides in CpG site blocks change simultaneously when neighboring CG dinucleotides are hypermethylated, hypomethylated, or differentially methylated. These characteristics were significant for any dinucleotide in the entire genome (P < 10^{-441} by a two proportion z-test), in intergenic regions (P < 10^{-263}), in exons (P < 10^{-153}), and in introns (P < 10^{-153}) (Supplemental Table S5).

One might be concerned that the high CG substitution rate could affect the substitution rates for dinucleotides other than CG;
however, the effects are limited to dinucleotides immediately before CG dinucleotides and are considerably small due to the rarity of CG dinucleotides (2.0% in the medaka genome). In fact, the substitution rates of dinucleotides excluding the dinucleotides overlapping CGs by one base remain essentially unchanged (Supplemental Fig. S3). Another possible explanation for the slower mutation rate in evolutionarily conserved hypomethylated regions is that these regions are around transcription start site (TSS) regions that are highly conserved (Taylor et al. 2006; Sasaki et al. 2009); however, evolution in such regions remained approximately the same even after hypomethylated regions near TSSs (those lying in the 500 bp immediately upstream of and downstream from the TSSs for 23,531 predicted medaka genes) were excluded from the analysis (Supplemental Fig. S4; Supplemental Table S3A,B). The higher substitution rate in evolutionarily conserved hypomethylated regions may be influenced by transposable elements which are highly mutated and methylated in vertebrates (Goll and Bestor 2005). However, transposons in the medaka genome are extremely rare and can be ignored (Kasahara et al. 2007). Therefore, methylation-dependent factors other than the influence of gene transcription or transposons may contribute to the slow evolution of hypomethylated regions.

A related problem is that methylation patterns may be affected by cis-acting polymorphisms, such as mutations in CpG sites (Schilling et al. 2009; Shoemaker et al. 2010), but little is known about fundamental cis-elements. It is highly nontrivial to identify cis-elements that induce the CpG methylation state. We attempted to find sequence motifs that were enriched and evolutionarily conserved in hypomethylated regions because such motifs could be candidate cis-elements and would be informative for further studies. We, therefore, searched for potential conserved motifs that were capable of distinguishing two collections of 10,000 hypo- or hypermethylated CpG site blocks using AdaBoost, an established machine learning algorithm (see Methods). The most significant motif was “CGCG,” and the incidence of “CGCG” in 10,000 hypomethylated regions was 61.39%, compared with 21.84% in 10,000 hypermethylated regions, showing the significant relevance of “CGCG” sequence enrichment to the methylation state ($P < 10^{-680}$ by a two-proportion z-test), as illustrated in Figure 2A. We then examined whether the CGCG motif is evolutionarily conserved in hypomethylated regions. For this purpose, it was necessary to predict the ancestor of Hd-rR and HNI strains. Therefore, we assembled a draft genome of an outgroup medaka strain, HSOK. Figure 2B shows that CGCG is significantly more conserved in hypomethylated regions than in hypermethylated regions ($P = ~0.203\%$ by a two-proportion z-test). This evolutionarily preserved CGCG motif in hypomethylated regions may be attributable to slow CpG deamination or fast CpG-gaining. To clarify this issue, we calculated dinucleotide-gaining substitution rates by comparing the genomes of Hd-rR and HNI to their ancestor which was estimated from the outgroup HSOK genome (Fig. 2C; see Methods). The rate of CpG gain in hypomethylated regions, 1.4%, was much smaller than the 3.0% rate observed in hypermethylated ones, supporting the hypothesis that slow CpG deamination is the dominant factor for maintaining CGCG richness in hypomethylated regions (Cohen et al. 2011). Meanwhile, we observed an intriguing property in differentially methylated regions in different strains: The rates of CG/CC gain in

Figure 2. (A) Representative mutation patterns in evolutionarily conserved hypo- and hypermethylated regions and in regions that are differentially methylated in different strains, where substitution rates ascend from top to bottom. The CGCG motif is significantly conserved in hypomethylated regions ($P < 10^{-680}$ by a two-proportion z-test). (B) Number of strain-differentially methylated CpG site blocks with either gain or loss of the CGCG motif. Of 1656 CGCG motif occurrences in multiple alignments of the genomes of the three strains, 52 (24, respectively) were conserved in hypomethylated (hypermethylated) regions of one of Hd-rR or HNI but were mutated in hypermethylated (hypomethylated) regions of the other strain. We then evaluated the significance of the difference between the means in the two groups, 52/1656 vs. 24/1656, to obtain a $P$-value of 0.203% according to a two-proportion z-test. (C) The rates of dinucleotide gain in CpG site blocks. Gain rates in the hypo- or hypermethylated CpG site blocks and each mate in strain-differentially methylated blocks are shown.
the hypomethylated mate are higher than those in the hypermethylated mate (3.3% and 2.4% compared with 2.6% and 2.2%, respectively), suggesting that CG/CC gains are involved in the evolution of methylation among different strains.

Next, we examined the methylation and genetic variations of two germline-like tissue types (blastulae and testes) and somatic liver cells that are fully differentiated and mostly homogeneous. In this analysis, we focused on tissue types in a single strain, Hd-rR. Figure 3A shows an abundance of CpG site blocks with tissue-differentially methylation states such that differences in methylation levels between each of the germline-like tissues and liver are more than 0.5. Notably, ~99% of tissue-differentially methylated regions were hypomethylated in the liver but were hypermethylated in blastulae and testes. Remarkably, as illustrated in Figure 3B, the tissue-differentially methylated regions showed a significantly lower SNP rate than did the hypomethylated ones ($P < 10^{-82}$) (Supplemental Table S6). Further analysis on dinucleotides revealed that the substitution rates of dinucleotides except for CC/GG/CG in differentially methylated regions were also significantly lower than those in hypomethylated regions ($P < 10^{-3}$) (Fig. 3C; Supplemental Table S7A), which was also seen even after excluding regions near TSSs (Supplemental Fig. S6). The high CG substitution rate in these regions could have been a consequence of deamination mutations occurring in primary germline-like cells. These results suggest that, because change in DNA methylation can affect transcription so that it can regulate reprogramming during somatic development, a positive selective pressure of genomic sequence could be involved in these differentially methylated regions (Fig. 3D).

**Discussion**

Because DNA methylation may influence genetic variation, we measured the incidence of genetic variations in methylation states in the human genome as well as in the two medaka inbred strains. We focused on germline-like tissue types, blastulae and testes, to determine the primary effects of methylation and evaluated a total of six single-base-resolution DNA methylomes in the medaka system. Our findings provide novel insights into the relationships between the dynamics of methylation states and genomic variations (Figs. 2A, 3D). Figure 4 illustrates our hypothesis for explaining the difference observed in genetic evolution between hyper- and hypomethylated region. Given that deaminated cytosine (U:G mismatch) is recognized and repaired by base excision repair (BER), deaminated 5-methyl-cytosine (T:G mismatch) would be corrected via more complicated mismatch-repair pathways (Molaro et al. 2011; Fig. 4). Previous studies revealed that two glycosylases, thymine DNA glycosylase (TDG) and methyl-CpG-binding domain...
methylated regions (Fig. 4).

5-methyl-cytosines could lead to a higher mutation rate in hyper-error-prone synthesis of the erased gaps around deaminated fidelity (Loeb and Monnat 2008). Thus, we speculate that the matches, which could involve DNA polymerases with lower mismatch-repair system (MMR) might gain access to T:G mis-

error-prone synthesis of erased gaps

\[\text{DNA replication amplifies errors}\]

Figure 4. A working model for a higher mutation rate in hyper-methylated regions. Deaminated cytosine (U-G mismatch) is repaired by base excision repair (BER), but deaminated 5-methyl-cytosine (T-G mismatch) is corrected by more complicated repair pathways. An alternative mismatch repair system (MMR) might involve low-fidelity DNA polymerase, resulting in the error-prone synthesis of erased gaps.

protein 4 (MBD4), are associated with T-G repair, but processing by these enzymes is biased in the genome and is relatively ineffective (Neddermann et al. 1996; Yoon et al. 2003). An alternative mismatch-repair system (MMR) might involve DNA polymerases with lower fidelity (Loeb and Monnat 2008). Thus, we speculate that the error-prone synthesis of the erased gaps around deaminated 5-methyl-cytosines could lead to a higher mutation rate in hyper-methylated regions (Fig. 4).

In an effort to search for candidate cis-elements for affecting the CpG methylation state, we observed a significant correlation between the methylation state of CpG sites and the presence of CGCG in the neighborhood. This CGCG motif was evolutionarily conserved in hypomethylated regions with statistical significance. A comparative genomic analysis showed slow CpG-gaining in hypomethylated regions, supporting the hypothesis that the evolutionary conservation of CGCG was brought about by slow CpG deamination (Cohen et al. 2011). It is intriguing to ask whether or not CGCG and other candidate motifs are associated with or affect allele-specific methylation. Breeding the F1 generation of the two medaka inbred strains is a promising approach to this problem, and high-throughput short-read bisulfite sequencing is an inexpensive way of observing DNA methylene, though it suffers from a technical difficulty in distinguishing short reads of ~100 bp from two different alleles. The high incidence of genetic variation (~3.4%) between the two inbred strains would facilitate achieving this classification task because an average of 3.4 SNPs per 100-bp read would make it easier to align a read to its originating allele.

Overall, these genetic consequences of methylation, including the slower evolution rate of the cis-element “CGCG” found in hypomethylated regions of the medaka system, should also be explored in other species.

Methods

Construction and sequencing of medaka bisulfite-treated DNA libraries

Genomic DNA from medaka tissue cells (blastulae, testes, and liver from two medaka strains, Hdr-R and HNI) was isolated and sonicated to a desired size range (100–400 bp). The DNA fragments were treated with DNA polymerase to generate blunt ends and were ligated with double-stranded DNA adaptors containing methylated cytosines, which were designed to amplify only those DNA fragments carrying bisulfite-converted adaptor sequences at both ends. Followed by 7–10 cycles of PCR, 250–450-bp size-fractionated DNA was sequenced using an Illumina GAIIx genome analyzer. A validation experiment was performed according to the same procedure using genomic DNA from S. cerevisiae S288C.

Computational and statistical methods

We converted all cytosines in reads and in both the Watson and Crick strands of the reference genome to thymines for primary mapping and used Smith-Waterman alignments between the original sequences of primary best hits. All possible methylation patterns were evaluated, and only uniquely mapped reads were retained for further analyses. The level of methylation of a particular cytosine was estimated by dividing the number of mapped reads reporting a cytosine (C) by the total number of reads reporting a C or T (thymine). The comparison of substitution rate and methylation level was performed among CpG site blocks, which consist of at least one CpG site and its surrounding upstream or downstream 10 bases. Overlapping CpG site blocks were merged. A two-proportion z-test was performed to test the significance of the substitution rate difference observed between hypo-hypermethylated CpG site blocks. The AdaBoost algorithm was used to find conserved motifs in hypo-/hypermethylated CpG site blocks. Estimation of the ancestor sequence of Hdr-R and HNI was given by multiple alignments using an out-group medaka strain HSOK which was assembled using the SOAPdenovo assembler (Li et al. 2010). Details of methods can be found in Supplemental Methods.

Data access

All sequence data are deposited at the NCBI Sequence Read Archive (SRA) (http://www.ncbi.nlm.nih.gov/sra) (accession number SRA026693).

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