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The Mechanisms of Generation, Recognition, and Erasure of DNA 5-Methylcytosine and Thymine Oxidations*

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One of the most fundamental questions in the control of gene expression in mammals is how the patterns of epigenetic modifications of DNA are generated, recognized, and erased. This includes covalent cytosine methylation of DNA and its associated oxidation states. An array of AdoMet-dependent methyltransferases, Fe(II)- and α-ketoglutarate-dependent dioxygenases, base excision glycosylases, and sequence-specific transcription factors is responsible for changing, maintaining, and interpreting the modification status of specific regions of chromatin. This review focuses on recent developments in characterizing the functional and structural links between the modification status of two DNA bases 5-methylcytosine and thymine (5-methyluracil).

In the DNA of higher organisms, cytosine exists in several chemical forms, including unmethylated cytosine (C), 5-methylcytosine (5mC), 5-hydroxymethylcytosine (5hmC), 5-formylcytosine (5fC), and 5-carboxylcytosine (5caC) (1–5). These forms are genetically equivalent in terms of base-pairing and protein-coding, but differ in how they interact with macromolecules and influence gene expression. There is much interest in the effects of these modifications in epigenetic regulation, in development and differentiation, in neuron function, and in diseases. In general, the modifications (or “marks”) are added to cytosine in situ, following its incorporation into DNA in the unmodified form. DNA methyltransferases (Dnmt) convert certain cytosines to 5mC, usually within the sequence context CpG (6, 7) or CpA (8). A subset of these 5mC residues is then converted to 5hmC, 5fC, and 5caC in consecutive Fe(II)- and α-ketoglutarate-dependent oxidation reactions by the tet-11en translocation (Tet) dioxygenases (2–4). The Tet dioxygenases are widely distributed across the eukaryotic tree of life, from mammals to the amoeboflagellate Naegleria gruberi (9), mushroom (Coprinopsis cinerea) (10), and honey bee (Apis mellifera) (11). High-throughput methods for characterizing 5mC oxidation states at single base resolution are becoming available, so our understanding of 5mC oxidation is expected to develop rapidly. In this minireview, we focus on the mechanisms of generating, recognizing, and possibly erasing 5mC and its oxidative forms in DNA.

Tet Proteins Are 5-Methylpyrimidine Dioxygenases, but Not Demethylating Enzymes

Chromatin regulates transcriptional processes through post-synthetic modifications of both of its components: DNA and histones (Fig. 1a). Much remains to be learned about how the combination of these modifications (or lack thereof) facilitates or silences transcription. One broad theme has emerged that a web of interactions tightly coordinates the modification of a segment of DNA and its associated histones, affecting local chromatin structure and determining the functional states. The Tet enzymes belong to a family of Fe(II)- and α-ketoglutarate-dependent dioxygenases that also includes the Jumonji domain-containing histone lysine demethylases, the N-methyl nucleic acid demethylase including Escherichia coli AlkB and its mammalian homologs, and many others. To comprehend the unique features of Tet-driven reaction products, we first briefly review the mechanisms of other dioxygenases considered as demethylases of amino (N)-methylation. The N-demethylation reaction catalyzed by most of these enzymes involves the transient formation of a N-hydroxymethyl intermediate followed by the spontaneous (non-enzymatic) release of formaldehyde.

In the case of protein lysine demethylation, the FAD-dependent amino oxides LSD1/2 and Fe(II)- and α-ketoglutarate-dependent Jumonji dioxygenases (12) generate a hydroxymethyl intermediate for each reaction that subsequently decomposes to release a formaldehyde spontaneously (without additional enzymatic activities) and the demethylated lysine (Fig. 1b) (with one methyl group reduced). The same basic mechanism applies to demethylation of N-methylated nucleic acids (e.g. N6-methylcytosine, N4-methylthymine, and N3-methyladenine) by the AlkB family of DNA repair enzymes (13, 14) (Fig. 1c) as well as the recently described demethylation of N6-methyladenine (N6mA) in mRNA by ALKBH5 and fat mass and obesity-associated protein (FTO) (15, 16) (Fig. 1d). In fact, during the search for enzymes capable of reversing methylated lysines in histones, it was initially hypothesized that Fe(II)- and α-ketoglutarate-dependent dioxygenases might reverse lysine methylation via a mechanism similar to that of AlkB (17), and the purification of JHDM1, the first identified Jumonji domain-
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(a) DNA modifying enzymes

(b) Protein (histone) lysine methylation and demethylation

(c) DNA N3mC and demethylation

(d) mRNA N6mA and demethylation

(e) DNA 5mC and thymine oxidations

(f) 5fu --- 5caU

(g) DNA replication
containing histone demethylase, used a biochemical assay based on the detection of formaldehyde, one of the predicted reaction products (18).

In stark contrast, formaldehyde is not released during Tet-mediated hydroxylation of 5-methylcytosine (Fig. 1e). Unlike methylation/demethylation of monoamines, the main mechanism of polymerization and demethylation of carbons is that the C5 atom of the cytosine ring is an inert carbon. DNA cytosine methyltransfersases solve this problem by flipping the target base into a concave active site and forming a transient covalent adduct at cytosine C6 (19, 20). However, the 5-hydroxymethyl modification at C5 (5mC) does not change the nature of the C5-CH3 carbon-carbon bond and thus either stays as a stable modification or is further converted to 5fC and 5caC in consecutive Tet-mediated oxidation reactions, generating stable modifications further away from 5mC. That Tet-mediated 5mC remains a stable mark, rather than serving as an intermediate in direct demethylation, is supported by the observation that 5mC loss in the paternal genome, immediately after fertilization during mouse development, is accompanied by concurrent increase in 5hmC (21).

Very recently, it has been shown that Tet enzymes are also active on thymine, generating 5-hydroxymethyluracil (5hmU) paired with adenine that could be specifically bound by potent protein readers (22, 23) (Fig. 1f). This is distinct from G:5hmU, the deamination product of G:5hmC. Like 5mC, thymine contains a methyl group at C5, and thus the Tet-mediated hydroxylation of 5-methyluracil (thymine) provides an opportunity to establish a pseudo-symmetric, modified CpA/TpG dinucleotide (Fig. 1g). CpA/TpG could be considered an intrinsically “hemi-methylated” DNA element. Methylation at CpA sites by Dnmt3 would then generate a “fully methylated” mCpA/TpG dinucleotide (Fig. 1f). Indeed, the levels of mCpA/TpG undergo dynamic changes during differentiation of the germ line and in brain development from fetus to young adult (24, 25). Some transcription factors recognize 5mCpG and TpG in the same way (26) (see below). Perhaps TpG dinucleotides are selected for when it is advantageous for a particular DNA sequence to be treated as if it were permanently (hemi)-methylated. Further modification by Tet enzymes on both 5mC and T would generate fully modified CpA/TpG sites with 5-hydroxymethyl or even 5-formyl and 5-carboxyl modification on both 5mC and T (23). Such fully modified CpA/TpG might even be maintained during DNA replication through the combined activity of Dnmt3 and Tet enzymes (Fig. 1g). A maintenance methylation function has been proposed for both Dnmt3a and Dnmt3b (27). Assuming that hemi-hydroxymethylated and hemi-methylated sites are substrates for modification by Tet proteins, there is also the potential for regenerating a fully modified hydroxymethylated site, recapitulating the parental DNA state after replication (Fig. 1g).

DNA methylation patterns are reprogrammed at two distinct stages during early vertebrate development: in the zygote and early embryo when DNA methylation patterns contributed by the sperm and egg are largely erased and subsequently re-established, and again during germ cell development, giving rise to germ cell-specific patterns that will be carried into the next generation (21). The Tet enzymes and oxidized 5mC derivatives, presumably through their role as intermediates in 5mC turnover (see below), play a critical role in both processes. Although Tet1 and Tet2 single knock-out mice are viable and reproduce normally (28–30), the majority of Tet1 and Tet2 double knock-out mice die perinatally, and female survivors exhibit reduced fertility (31), implying some functional redundancy between Tet1 and Tet2. In contrast, deletion of Tet3 leads to neonatal lethality (32), and loss of all three Tet enzymes restricts normal differentiation of mouse embryonic stem cells (33). Consistent with a critical role of Tet3 in early embryonic reprogramming, there are high levels of Tet3 and 5hmC in the zygote (34) and conditional deletion of Tet3 in the oocyte results in aberrant hydroxylation and delayed demethylation of the paternal genome upon fertilization, leading to impaired development and reduced fertility (32). It is interesting to note that although Dnmt3a (but not Dnmt3b or Dnmt1) is expressed in both the male and the female pronucleus of the mouse zygote, Tet3 (but not Tet1 or Tet2) is only expressed in the male pronucleus, as observed in an earlier study (32). Thus, Dnmt3a and Tet3 selectively converge in the male pronucleus and coincide with the replication-independent loss of 5mC and gain of 5hmC that occur in the paternal genome in the early mouse embryo (34, 35). However, more recent work indicates that there may also be low levels of Tet3 expressed in the female pronucleus (36, 37), suggesting that 5mC oxidation may also play a role in the erasure of the oocyte-specific DNA methylation in the zygote. Together, the accumulated data speak to a role for both active (via Tet3) and passive (via DNA replication) demethylation in the zygotes, in the early embryo, and in primordial germ cells (38).

Despite the widespread role of 5mC in the regulation of vertebrate gene expression, the situation may be different for invertebrates. For example, there is currently no evidence for the presence of 5mC in Drosophila melanogaster (39). Never-
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theless, the *Drosophila* genome harbors a well conserved homolog of the mammalian Tet gene, the function of which is currently unknown (40). It is possible that *Drosophila* Tet may oxidize other methylated bases, such as thymine in DNA or 5mC in tRNA (generated by Dnmt2 (41)) or N6mA in mRNA or DNA. A very recent publication (100) has appeared demonstrating that *Drosophila* Tet catalyzes demethylation of N6mA in DNA both in vivo and in vitro as well as catalyzing 5mC in vitro.

Structures of Tet Dioxygenases in Complex with 5mC DNA

Two x-ray structures are currently available for Tet enzymes: the catalytic domain of human TET2 (42) and a *Naegleria* Tet-like dioxygenase, NgTet1 (9). Like DNA methyltransferases, Tet enzymes use a base-flipping mechanism to access 5mC. Structurally, NgTet1 represents the core structure of the catalytic domain of the mammalian Tet enzymes (Fig. 2a). Like other structurally characterized α-ketoglutarate-dependent dioxygenases, NgTet1 has a core double-stranded β-helix fold that binds Fe(II) and α-ketoglutarate (Fig. 2b). Two twisted β-sheets (a four-stranded minor sheet and an eight-stranded major sheet) pack together with five helices on the outer surface of the major sheet to form a three-layered structure (Fig. 2b). The unequal number of strands of the two sheets creates the active site located asymmetrically on the side of the molecule where the extra strands of the major sheet are located.

There is considerable sequence conservation within the catalytic core among Tet enzymes from different species (Fig. 2c). When compared with NgTet1, mammalian Tet enzymes (human and mouse Tet1–3) and the honey bee Tet protein have eight insertions and one deletion scattered throughout the catalytic core domain (Fig. 2c). Two zinc ion centers, each coordinated by three cysteines and one histidine, bring together the loop inserted between strands 3 and 4 to the N-terminal region, and the three loops between strands 1 and 2, helix 2 and strand 5, and strands 6 and 7, respectively. These intramolecular interactions likely confer stability to the molecule. Invariant residues are involved in structural integrity, hydrophobic core, DNA binding, base-specific interactions, and Fe(II) and α-ketoglutarate binding.

The mammalian and honey bee Tet proteins have their catalytic domains located in the C-terminal part of the proteins with an atypical insertion of ~300 residues between strands β8 and β9, which is not found in other α-ketoglutarate-dependent dioxygenases, nor in NgTet1. The insertion separates the two halves of the ferrous binding motif, HXD . . . H. Interestingly, the large insertion in TET1 shares significant sequence similarity to the C-terminal domain of RNA polymerase II (43), suggesting that it might impart a regulatory function. Additionally, in human TET2, part of this insertion forms a fourth layer on the outer surface of the minor sheet (Fig. 2d).

Recognition of 5mCpG and TpG and Their Oxidized Derivatives

DNA 5mC is a major epigenetic signal that acts to regulate chromatin structure and ultimately gene expression. The oxidized derivatives, 5hmC, 5fC, and 5caC, might also act as distinct epigenetic signals. These modifications protrude into the major groove of DNA, the primary recognition surface for proteins, and change its atomic shape and pattern of electrostatic charge. In principle, such changes can alter the way in which proteins bind to their recognition sequences in DNA by strengthening the interactions, weakening them, or abolishing them altogether. This, in turn, can modulate gene expression and control cellular metabolism and is believed to be one of the principal mechanisms underlying epigenetic processes such as differentiation, development, aging, and disease.

Three well characterized classes of mammalian proteins interact with DNA in a methylation-dependent manner. Methy-binding domain proteins (MBDs) recognize fully methylated CpG sequences in which both DNA strands contain 5mC. The MBD domain of MeCP2 binds the symmetrical, fully methylated CpG site using two arginines; each hydrogen bonds to one guanine (44). The two arginines are also engaged in van der Waals contacts with the methyl group of the neighboring 5′-5mC of the same DNA strand and form two “5mC-Arg-G triads” (Fig. 3a) to symmetrically bind the fully methylated CpG palindromic duplex. Very recent work suggests that MeCP2 binds methylated CpA sites with similar affinity to that of fully methylated CpG (45, 46). Likewise, the MBD domain of MBD4 binds dinucleotides containing fully methylated 5mCpG/5mCpG and G:T mismatches occurring in the CpG context (5mCpG/TpG) (47). In all three examples, there are two methyl groups present symmetrically (5mCpG/5mCpG, 5mCpA/TpG, and 5mCpG/TpG), suggesting that the MBD proteins simultaneously recognize the two methyl groups on opposite strands.

The SET and RING finger-associated (SRA) domain of UHRF1 recognizes hemi-methylated CpG sequences containing 5mC in only one strand, such as those that arise during DNA replication (48). UHRF1 binds 5hmC DNA with ~10-fold weaker affinity than 5mC DNA (49), whereas the SRA domain of UHRF2 has a slightly stronger affinity for 5hmC DNA than 5mC DNA by a factor of ~1.5 (50). Recently, it was suggested that UHRF22 can directly and specifically bind A:5hmU (relative to A:T) in vitro (22). The SRA domain uses a DNA recognition mode vastly different from that of MBD and zinc finger (ZnF) proteins (see below). Rather, the SRA domain, as a non-enzymatic, sequence-specific DNA-binding domain, utilizes a base-flipping mechanism to interact with DNA, similar to that of DNA-modifying enzymes such as DNA methyltransferases, DNA glycosylases, and Tet dioxygenases.

Almost 20 years ago, Holliday argued that “sequences longer than CpG would be necessary for the regulation of gene expression by methylation” (51). Indeed, the recognition of some sequence-specific transcription factors is blocked by cytosine methylation (for example, CCCTC-binding factor (CTCF) (52, 53)). In contrast, certain Cys2-His2 (C2H2) ZnF proteins bind preferentially to DNA when CpG sites embedded within their recognition sequences are methylated (54). The structures of five ZnF domains bound to 5mC-containing DNA have been solved, including the transcription factors Kaiso, Zfp57, Krüppel-like factor 4 (Klf4), growth response protein 1 (Egr1), and Wilms tumor protein 1 (WT1) (55–58). Kaiso recognizes either methylated CpG dinucleotides (59) or an unmodified sequence with a TpG in the place of 5mCpG (60). Structures of the ZnF
FIGURE 2. Structures of the Tet enzymes. a, schematic representation of human Tet1 (hTet1) C-terminal catalytic domain and NgTet1. b, structure of NgTet1-DNA complex (Protein Data Bank (PDB) 4LT5). The NgTet1 protein folds in a three-layered jelly-roll structure. c, sequence alignment of human TET1, TET2, and TET3 (NP_085128.2, NP_001120680.1, and NP_001035490.2), mouse Tet1, Tet2, and Tet3 (NP_081660.1, NP_001035490.2, and NP_081660.1), honey bee (A. mellifera) AmTet (GB52555 in BeeBase OSGv3.2), and NgTet1 (XP_002667965.1). d, structure of human TET2-DNA complex (PDB 4NM6). The secondary structure elements are labeled according to NgTet1 structure (panel b). Note the large insertion in human TET2 between strands 8 and 9 (magenta), which is indicated by a magenta arrow in panel c.
A methyl-Arg-G triad forms during recognition of methyl-CpG and TpG dinucleotides in double-stranded DNA. a, MeCP2 forms two 5mC-Arg-G triads to bind the palindromic fully methylated CpG duplex symmetrically (PDB 3C2I). b, Kaiso recognizes either a specific unmethylated DNA element containing a TpG dinucleotide (top) or a methylated CpG dinucleotide (bottom) (PDB 4F6M and 4F6N). In both cases, a methyl-Arg-G triad is involved. c, Zfp57 uses a pair of methyl-Arg-G triads to recognize the TpG dinucleotide on the top strand and a methyl-CpG on the bottom strand (PDB 4GZN). A third methyl-Arg-G triad recognizes the TpG on the bottom strand (not shown). d, example of transcription factor recognition sequences containing a CpA/TpG site (taken from Ref. 93). e-i, examples of protein domains with specificity for unmodified cytosine (e), 5mC/H (f), 5hmC/H (g), 5fC/F (h), or 5caC (i). The Tet3 CXXC domain binds to an unmodified cytosine in any sequence context (94), which is distinct from the CXXC domains of MLL (95), CFP1 (96), and Dnmt1 (97), which are restricted to unmodified CpG sites. CHTOP (chromatin target of PRMT1) binds to 5hmC (98). The modification of the CG dinucleotide (underlined) of TGACGCAA to 5mC, 5fC, or 5caC enhances DNA binding by the basic leucine zipper protein C/EBP (99). The carboxylation of cytosine (5caC) in a CpG dinucleotide adjacent to the consensus recognition sequence (underlined) of the basic-helix-loop-helix transcription factor proteins Tcf3/Ascl1 (CAGCANNNTG) enhanced binding of the heterodimer by ~10-fold (64).
domain of Kaiso revealed that Arg-511 of Kaiso interacts with the 5mCpG and TpG dinucleotides in a similar fashion (55), forming a methyl-Arg-G triad (26) (Fig. 3b), just like that described above for the two MBD arginines. The triad is maintained for TpG because the thymine methyl group is in the equivalent position (5-carbon) to that in 5mC. Similarly, Zfp57 uses a pair of methyl-Arg-G triads to recognize 5mCpG and TpG dinucleotides on the two strands, respectively (Fig. 3c).

Klf4 shows the strongest binding to fully methylated DNA, with slightly higher affinity (1.5-fold) than that of the unmodified DNA, and each oxidation event, from 5mC to 5hmC to 5fC to 5caC, results in progressively weaker binding (by factors of 2, 3, and 6, respectively) (57). Like Kaiso, the consensus-binding sequence element for Klf4 contains either CpG, which can be methylated, or TpG, which is intrinsically methylated on one strand (Fig. 3d). Klf4 is one of the four Yamanaka reprogramming factors (61), and along with the other three (Myc, Oct4, and Sox2), all contain TpG/CpA in their consensus recognition sequences (Fig. 3d). It may not be a coincidence that non-CG (mainly CpA) methylation disappears upon induced differentiation of embryonic stem cells and is restored in induced pluripotent stem cells resulting from expression of the reprogramming factors (24), which recognize CpA-containing sequences.

Both Egr1 and WT1 bind the same consensus DNA sequence containing two CpG sites and display high affinity for the sequence with C or 5mC but much reduced affinity when 5hmC or 5fC to 5caC, results in progressively weaker binding (by factors of 2, 3, and 6, respectively) (57). Like Kaiso, the consensus-binding sequence element for Klf4 contains either CpG, which can be methylated, or TpG, which is intrinsically methylated on one strand (Fig. 3d). Klf4 is one of the four Yamanaka reprogramming factors (61), and along with the other three (Myc, Oct4, and Sox2), all contain TpG/CpA in their consensus recognition sequences (Fig. 3d). It may not be a coincidence that non-CG (mainly CpA) methylation disappears upon induced differentiation of embryonic stem cells and is restored in induced pluripotent stem cells resulting from expression of the reprogramming factors (24), which recognize CpA-containing sequences.

The second mechanism is an active DNA demethylation/demodification of 5mC and its derivatives that involves DNA glycosylases and base excision repair. Arabidopsis ROS1 and mammalian TDG are the two DNA glycosylases currently implicated in this process, which involves the removal of the modified cytosine base, subsequent processing by the base excision repair machinery, and ultimately the incorporation of an unmodified C or T status.
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5caC (69–71), whereas TDG removes 5fC and 5caC but not 5mC and 5hmC (4, 72–74) (Fig. 4).

Arabidopsis ROS1 is a multi-domain bifunctional DNA glycosylase/lyase, which excises 5mC and 5hmC as well as thymine and 5hmU (i.e. the deamination products of 5mC and 5hmC) when paired with a guanine, leaving an apprimarydic site that is subsequently incised by the lyase activity (75). ROS1 is slow in base excision and fast in apyrinoid lyase activity in vivo (71, 76), indicating that the recognition of pyrimidine modifications might be a rate-limiting step or that other cofactors might be needed to stimulate ROS1 enzymatic activity (in a manner similar to that described for Dnmt3a activity stimulation by Dnmt3L (77)). Mammalian DNA glycosylases that excise 5mC or 5hmC have not been identified, but such activities have been reported (78, 79).

Mammalian TDG excises the mismatched base from G:X mismatches, where X is uracil, thymine, or 5hmU. These are, respectively, the deamination products of cytosine, 5mC and 5hmC. In addition, TDG excises the Tet protein products 5fC and 5caC when paired with a guanine, but not 5hmC or 5mC. It is worth noting that ROS1 is inactive on G:U (71, 76, 80), whereas TDG, although named as a thymine DNA glycosylase, has much faster activity on G:U mismatches (74, 81). Furthermore, the structurally related E. coli mismatch uracil glycosylase (eMUG) can excise 5caC and 5fC as well (74, 82). Both 5fC and 5caC exhibit an intrabase hydrogen bond between their formyl or carboxyl oxygen atoms, respectively, and the adjacent cytosine N4 exocyclic amine nitrogen atom, both in the free form (83) and in the protein-bound form (58) (Fig. 3, h and i, top panels). It is unknown whether this intrabase hydrogen bond contributes to the previously observed mutagenic potential of 5fC and 5caC in vivo and in vitro (84, 85). Both DNA and RNA polymerases sometimes misincorporate adenine opposite of the 5fC or 5caC, whereas little misincorporation is observed on 5mC- and 5hmC-containing templates. It is possible that the tendency for mismatches to arise opposite 5fC and 5caC, whereas little misincorporation is observed on 5mC- and 5hmC-containing templates. It is possible that the tendency for mismatches to arise opposite 5fC and 5caC (but not 5mC and 5hmC) is somehow correlated with differential excision by TDG. It is still not clear whether the differential excision of 5hmC versus 5fC and 5caC by TDG lies in the early steps of intrahelical interrogation to detect the C5 modification and the initiation of base flipping, or in the post-flipping steps arising from differences in the active-site transition states of the enzyme-bound complexes. It is possible that TDG/eMUG might recognize or even promote a mismatch-like “wobble” pair of G:5fC and G:5caC (with the aid of the intrabase hydrogen bond) and turn them into substrates, whereas ROS1 is sensitive to pyrimidine modifications at C5 position.

In light of the recent observation that 5hmU could also be the product of Tet-mediated thymine hydroxilation (22, 23), it is interesting to note that SMUG1 (named after single-strand-specific monofunctional uracil-DNA glycosylase) can excise 5hmU when mispaired with a guanine as well as paired with an adenine (86) or 5FU in any base pair context (87). In addition, members of the NEIL family of DNA glycosylases have been found to bind oxidized 5mC derivatives 5hmC, 5fC, or 5caC (88) and 5hmU (22), and partially rescue the loss of TDG in a gene reactivation assay (89).

Perspective

Although it is well accepted that DNA methylation patterns are replicated in a semi-conservative fashion during cell division via the selective recognition of hemi-methylated Cpg dinucleotides at DNA replication forks, one of the unresolved fundamental questions is how, and indeed whether, the pattern of oxidized 5mC derivatives is similarly “inherited” at CpG and CpA sites. Many transcription factors (e.g. Klf4 and MeCP2) recognize consensus-binding elements, containing either CpG, which can be methylated, or TpG (or CpA), which is intrinsically methylated on one strand and can be methylated on the other strand and further modified on both strands. Transcription factors may have adapted to respond to different states of cytosine modification. These states can affect binding affinity, and so gene activity could plausibly be controlled on a much finer scale by these modifications than a simple “on” or “off.” This hints, perhaps, at new levels of subtlety and versatility in epigenetic regulatory processes.

Finally, members of base excision DNA glycosylases remove the modified cytosine and thymine bases. ROS1/DME, TDG/eMUG, and SMUG1 excise modified bases from “natural” base pairs, but have little sequence specificity for the surrounding DNA context. Another enzyme, R.Pabi from Pyrococcus abyssi, initially identified as a restriction enzyme, was recently determined to be a sequence-specific adenine DNA glycosylase (90). It is therefore possible that modification-dependent and sequence-specific DNA glycosylases might exist to recognize the modification within specific sequences, similar to the family of modification-dependent restriction endonucleases (91). Methylpurine glycosylases (MPGs) from E. coli, Saccharomyces cerevisiae, and human have been reported to remove normal bases (G>A>C~T) from DNA (92). Methylpurine glycosylase protein was recently identified as a 5hmC binder in mouse embryonic stem cells and adult mouse brain using a DNA pull-down approach combined with quantitative mass spectrometry (88). Thus, not all DNA base excision glycosylases are strictly involved in mismatch DNA repair; some actually excise modified bases with proper Watson-Crick base pairing to accomplish DNA demodification. Moreover, intriguing recent work showing that the demethylation of male and female pronuclear DNA is unaffected by the deletion of TDG from the zygote (36) suggests that other demethylation mechanisms (or other DNA glycosylases) must exist downstream of Tet3-mediated oxidation.

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