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Active DNA demethylation: many roads lead to Rome

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Abstract

DNA methylation is one of the best-characterized epigenetic modifications and has been implicated in numerous biological processes, including transposable element silencing, genomic imprinting and X chromosome inactivation. Compared with other epigenetic modifications, DNA methylation is thought to be relatively stable. Despite its role in long-term silencing, DNA methylation is more dynamic than originally thought as active DNA demethylation has been observed during specific stages of development. In the past decade, many enzymes have been proposed to carry out active DNA demethylation and growing evidence suggests that, depending on the context, this process may be achieved by multiple mechanisms. Insight into how DNA methylation is dynamically regulated will broaden our understanding of epigenetic regulation and have great implications in somatic cell reprogramming and regenerative medicine.

> Eukaryotic chromatin contains a wealth of information required for the growth and development of a multicellular organism. This information is not only stored genetically in the DNA sequence itself but also epigenetically through DNA methylation and posttranslational modifications of histone proteins^{1,2}. Although every nucleotide in the genome has the potential to be transcribed³, the presence or absence of specific epigenetic marks influences gene expression, resulting in a transcriptional programme that specifies for a particular cell type. For example, in embryonic stem (ES) cells, active gene expression marks are found at pluripotent genes and repressive marks are found at lineage-specific genes. Thus, different cell types can be defined by their epigenetic and gene expression profiles.

During development, these transcriptional programmes undergo dynamic changes that ultimately lead to the production of distinct cell types and tissues that make up an organism. Accommodating such a transcriptional programme requires an epigenome that is both dynamic and flexible. Furthermore, the diversity of genetic material to be regulated necessitates the use of marks corresponding to short-term and long-term epigenetic memory, depending on the transcriptional requirements of the cell (as well as those of future generations). Developmental genes that are needed during the later stages of development are transiently held in a repressed state during early development. This is achieved through

FURTHER INFORMATION Yi Zhang's homepage: <http://www.med.unc.edu/~zhangyi/lab.htm>

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DATABASES Entrez Gene: <http://www.ncbi.nlm.nih.gov/gene> BDNF | CYP27B1 | FGF1 | MEDEA | nanog | pS2 | stella **UniProtKB:** <http://www.uniprot.org> AID | AlkB | APOBEC1 | BLIMP1 | DME | DML2 | DML3 | DNMT1 |DNMT3A | DNMT3B | ELP3 | ERα | Gadd45a | GADD45B | JBP1 | JBP2 | MBD2 | MBD4 | MeCP2 | M.EcoRII | M.HpaII | MLL |PRDM1 | PRDM14 | PTH | ROS1 | SMUG1 | TAF12 | TDG | TET1 |XPG | ZFP57

short-term epigenetic marks such as histone modifications, which can be removed before or within a few cell divisions.

By contrast, other regions of the genome are marked with epigenetic information that is stably maintained and heritable after many cell divisions. For example, imprinted genes, transposons and the inactive X chromosome require long-term silencing that is sustained throughout the development and lifespan of an organism. This is generally achieved by DNA methylation, an epigenetic mark that refers to the addition of a methyl group to the fifth carbon of base C. Because DNA methylation provides heritable, long-term silencing that is crucial for an organism, aberrant DNA methylation has been associated with cancer, imprinting-related diseases and psychiatric disorders⁴⁻⁷.

In mammals, DNA methylation occurs predominantly in the context of CpG (C followed by G) dinucleotides, whereas DNA methylation in plants can occur at C bases in diverse sequence contexts⁸. The enzymes responsible for this modification, DNA methyltransferases (DNMTs), are well characterized and conserved in mammals and plants⁸. DNMTs fall under two categories: *de novo* and maintenance⁹. Patterns of DNA methylation are initially established by the de novo DNA methyltransferases DNMT3A and DNMT3B during the blastocyst stage of embryonic development^{10,11} (FIG. 1). These methyl marks are then faithfully maintained during cell divisions through the action of the maintenance methyltransferase, **DNMT1**, which has a preference for hemi-methylated DNA¹²⁻¹⁴. Both the establishment and maintenance of DNA methylation patterns are crucial for development as mice deficient in DNMT3B or DNMT1 are embryonic lethal^{11,15} and DNMT3A-null mice die by 4 weeks of age¹¹.

Although DNA methylation has been viewed as a stable epigenetic mark, studies in the past decade have revealed that this modification is not as static as once thought. In fact, loss of DNA methylation, or DNA demethylation, has been observed in specific contexts (see below) and can occur through active or passive mechanisms (FIG. 1). Active DNA demethylation is the enzymatic process that results in the removal of the methyl group from 5-methylcytosine (5meC) by breaking a carbon-carbon bond. By contrast, passive DNA demethylation refers to the loss of the methyl group from 5meC when DNMT1 is inhibited or absent during successive rounds of DNA replication. whereas passive DNA demethylation is generally understood and accepted, the subject of active DNA demethylation has been controversial¹⁶.

In this Review, we explore what is known about active DNA demethylation and the disputes that are embedded in this field. First, we describe the contexts in which DNA demethylation has been observed and discuss the evidence that supports an active mechanism. we then present the many enzymes that have been proposed to carry out active DNA demethylation. we conclude by discussing emerging themes and highlighting the remaining questions in this exciting field.

Evidence for active DNA demethylation

Even though DNA methylation contributes to stable, long-term and heritable silencing, it has become apparent that in some instances DNA methylation levels can rapidly change by mechanisms involving active DNA demethylation. Genome-wide and gene-specific demethylation events have both been observed, but current evidence suggests that the former only occurs at specific times during early development, whereas the latter occurs in somatic cells responding to specific signals.

Genome-wide DNA demethylation of paternal pronuclei

Prior to fertilization, mammalian gametes are at different stages of the cell cycle and their genomes are organized differently. The egg is meiotically arrested at metaphase II, resulting in a diploid genome that is packaged with histones. Mature sperm, however, have completed meiosis, but their haploid genomes are packaged with protamines instead of histones. when a sperm penetrates the zona pellucida to fertilize the egg, both gametes undergo rapid changes. The egg completes its second meiosis resulting in the extrusion of one copy of the genome as the polar body; the sperm reorganizes its genomic DNA by replacing protamines with histone proteins.

Shortly after the protamine-histone exchange, the sperm-derived paternal pronucleus undergoes genome-wide DNA demethylation^{17,18}, an event that occurs quite rapidly within 4-8 hours post-fertilization (FIG. 2a). Although there are some disputes regarding the timing and synchrony of DNA replication in the zygote¹⁹⁻²⁵, loss of DNA methylation is seen before the completion of the first cell division. Thus, it is unlikely that a passive demethylation mechanism is the cause for this observation. Furthermore, when zygotes were treated with the replication inhibitor aphidicolin, paternal genome demethylation was still detected^{17,26}, further supporting an active demethylation mechanism.

Paternal genome demethylation has been observed in many mammalian organisms, including human, mouse, rat, bovine and $\frac{pi^{17,18,27,28}}{$, but seems to be absent from others, such as sheep²⁹. when sheep sperm are injected into mouse oocytes, demethylation is seen in the sheep-derived paternal genome³⁰, suggesting that the demethylating factor or factors are contributed by the oocyte. However, sheep oocytes injected with mouse sperm also resulted in demethylation of the mouse-derived paternal genome³⁰. Although this occurs to a lesser extent compared to mouse oocytes, it is likely that factors present in the sperm or features unique to the paternal genome also contribute to demethylation. Consistent with this notion, mouse oocytes can demethylate multiple male pronuclei³¹, but are incapable of demethylating the additional maternal genome in parthenogenetic, gynogenetic and digynic triploid zygotes 32 .

Although immunostaining studies suggest that demethylation occurs globally, bisulphite sequencing indicates that some genomic regions are resistant to such a wave of demethylation. These genomic regions include imprinting control regions 33 , intracisternal A-particle (IAP) retrotransposons³⁴ and centric and pericentric heterochromatin^{31,35}. It is not clear why these genomic regions are resistant to this wave of DNA demethylation, but one possibility is that methylation of these regions may be required to ensure transcriptional repression and chromosomal stability. Additionally, the maternal genome remains methylated during this time even though it is exposed to the same cytoplasmic factors. Insight into how some regions in the paternal genome are targeted for DNA demethylation whereas other regions are resistant may also provide clues as to how the maternal genome is protected from active demethylation (BOX 1).

The significance of zygotic paternal genome DNA demethylation is unclear at present. Genome-wide demethylation may facilitate transcriptional activation of the paternal genome³⁶, which has been reported to occur before transcriptional activation of the maternal genome in some species³⁷. Although some transposable elements and repeat sequences have been identified to be resistant to DNA demethylation, it is likely that others are still targets of DNA demethylation, given that these types of sequences account for half of the genome. whether demethylation of transposable elements and repeat sequences results in their reactivation and, if so, what the significance of their reactivation is remains to be determined.

Genome-wide DNA demethylation of primordial germ cells

After fertilization, the one-cell zygote undergoes several cell divisions that ultimately lead to formation of the blastocyst. During this developmental period, the maternal genome undergoes passive DNA demethylation (FIG. 2a) — a gradual loss of DNA methylation occurs with each cell division³⁸ in a replication-dependent manner³⁹. Consistent with this, maternally contributed DNMT1 (also known as DNMT1o) is excluded from the nucleus⁴⁰. Although passive DNA demethylation seems to affect a large part of the genome, imprinted genes still retain their methylation marks. Recent genetic studies indicate that maternal and zygotic DNMT1 (ReF. 41) and the zinc finger protein ZFP57 (ReF. 42) are required to maintain the DNA methylation imprints during pre-implantation development.

At embryonic day 7.5 (E7.5), signals originating from the extraembryonic ectoderm and the visceral endoderm instruct a subset of posterior epiblast cells to become primordial germ cells (PGCs). Specification of PGCs involves the BMP4 and BMP8 signalling pathway and activation of transcription factors BLIMP1 (also known as PR domain zinc finger protein 1 (PRDM1)) and PRDM14 (REFs 43,44). These founder cells of the germ line begin to migrate at E8.5 and arrive at the genital ridge at E11.5. At the beginning of their specification and migration, PGCs are thought to have the same epigenetic marks as other epiblast cells. However, by the time they arrive at the genital ridge, many of these marks including DNA methylation have been erased $45-47$ (FIG. 2b). Given that PGCs have undergone several cell cycles in the presence of DNMT1, this demethylation event is considered to be active. It is thought that global demethylation, including that of imprinted genes, takes place so that new DNA methylation patterns can be re-established, although experimental evidence supporting this remains to be shown.

Loci-specific active demethylation in somatic cells

Active DNA demethylation has also been reported in somatic cells, but only at specific genomic loci in response to certain signals. For example, within 20 minutes of stimulation, activated T lymphocytes undergo active demethylation at the interleukin-2 promoterenhancer region in the absence of DNA replication⁴⁸. Locus-specific demethylation has also been reported to occur at the promoter of brain-derived neurotrophic factor (*BDNF*)⁴⁹, the protein product of which is important for adult neural plasticity (FIG. 3a). In unstimulated neurons, the BDNF promoter is methylated, allowing for the recruitment of the repressive meC-binding protein, MeCP2. when depolarized with KCl, BDNF is upregulated, coinciding with the release of MeCP2 and demethylation of the promoter⁴⁹. Because this event takes place in post-mitotic neurons, active demethylation is thought to be the underlying mechanism. In addition to T cells and neurons, active DNA demethylation has been reported to take place during nuclear hormone-regulated gene activation (FIG. 3b). For example, the $pS2$ (also known as *TFF1*) promoter exhibits periodic methylation and demethylation that coincides with cyclical binding of oestrogen receptor-α (Erα) and expression of pS2 (REFs 50,51). Similarly, active DNA demethylation occurs at the cytochrome p450, subfamily 27B, polypeptide 1 (CYP27B1) promoter in response to parathyroid hormone $(PTH)^{52}$. These studies suggest that DNA methylation may not function solely as a long-term silencing mark, but could also function in the dynamic regulation of genes that require rapid responses to specific stimuli.

Mechanisms of active DNA demethylation

The importance of DNA methylation in diverse biological processes coupled with the observations of active DNA demethylation in embryonic development and somatic cells have led to extensive efforts in identifying DNA demethylases. DNA demethylase activity was first reported in murine erythroleukaemic nuclear extracts⁵³. Although it was determined that 5meC was ultimately replaced by C in a replication-independent manner,

Since then, several studies have led to the proposal of various mechanisms by which active DNA demethylation can occur. These include: enzymatic removal of the methyl group of 5meC, base excision repair (BER) through direct excision of 5meC, deamination of 5meC to T followed by BER of the T•G mismatch, nucleotide excision repair (NER), oxidative demethylation and radical S-adenosylmethionine (SAM)-based demethylation.

Enzymatic removal of the methyl group of 5meC

The simplest way to achieve DNA demethylation is through enzymatic removal of the methyl group of 5meC. This requires an enzyme with enormous catalytic power because of the strength of the carbon-carbon bond that needs to be broken. Methyl-CpG-binding domain protein 2 (MBD2) was the first reported protein to carry out this reaction. It did not require any specific cofactors, and was proposed to release methanol⁵⁶. This thermodynamically unfavourable mechanism was hotly contested for several reasons. First, previous studies had shown that MBD2 can stably bind methylated $DNA^{57,58}$, making it unclear how binding could occur if MBD2 was so efficient at removing the methyl group. Further concerns were raised when MBD2-null mice were not only viable, but also exhibited normal methylation patterns⁵⁹. Importantly, the paternal pronucleus of MBD2-null zygotes still exhibit normal demethylation³¹. These observations, coupled with the fact that no other laboratories could reproduce the reported enzymatic activity, have raised serious doubts on the capacity of MBD2 to serve as a DNA demethylase. Regardless of the controversy surrounding MBD2, it is still conceivable that a *bona fide* DNA demethylation mechanism exists. In fact, numerous histone demethylases that can break a carbon-nitrogen bond have recently been discovered^{60,61}. Although carbon-carbon bonds are inherently more difficult to break than carbon-nitrogen bonds, enzymes that have the capacity to do so have been reported in the thymidine salvage pathway⁶² and the cholesterol synthesis pathway⁶³.

BER through direct excision of 5meC

It has been proposed for some time that DNA demethylation can be achieved through the BEr DNA repair pathway (FIG. 4a). This type of repair involves a DNA glycosylase that removes the target base resulting in an abasic (apurinic and apyrimidinic (AP)) site. The DNA backbone is subsequently nicked by an AP lyase activity to generate a 5['] phosphomonoester and a 3′ sugar phosphate residue. An AP endonuclease then removes the 3′ sugar group leaving a single nucleotide gap that is ultimately filled in by DNA repair polymerases and ligases⁶⁴.

Active DNA demethylation can be accomplished by a DNA glycosylase that directly excises 5meC to initiate BER (FIG. 4a). Strong genetic and biochemical evidence supports the use of this mechanism in plants⁶⁵. In *Arabidopsis thaliana*, DNA demethylation is mediated by the Demeter (Dme) family of DNA glycosylases, which consists of four members: DME, repressor of silencing 1 (ROS1; also known as DML1), DML2 and DML3 (REF. 65). The discovery that these DNA glycosylases suppress DNA methylation initially came from forward-genetic screens in A. thaliana. whereas DME was discovered owing to the loss of expression of the imprinted gene $MEDEA$ in a loss-of-function DME mutant⁶⁶, ROS1 was recovered in a genetic screen for mutants that confer promoter hypermethylation and transgene silencing defects⁶⁷.

DME, ROS1, DML2 and DML3 possess glycosylase activity against oligonucleotides containing $5meC^{67-71}$. In addition, all members of the Dme family possess AP lyase activity

and are thus considered bifunctional glycosylases $69-71$. Besides CpG, DNA methylation in plants can occur in the context of CpNpG (where N is A, T or C) and CpNpN. All members of the Dme family have the capacity to recognize and remove meC bases from doublestranded DNA (dsDNA) oligonucleotides, irrespective of their sequence context in vitro⁷¹. However, attempts to determine the substrate specificity of these enzymes have resulted in conflicting reports owing to the use of different substrates and reaction conditions⁶⁸⁻⁷¹. In vivo studies indicate that mutation of each of these genes results in hypermethylation in all sequence contexts but at distinct genomic $\text{loc}^{69,71-73}$, indicating that each of these enzymes has a unique *in vivo* function.

Although it is clear that plants use BER to achieve DNA demethylation, evidence supporting a similar mechanism in mammals has been less compelling. Despite the lack of an obvious mammalian orthologue of the ROS1 family, the first indication that a repair mechanism could contribute to DNA demethylation came from early studies in chicken embryo extracts⁷⁴, revealing 5meC glycosylase activity against hemi-methylated DNA^{75} . Subsequent purification of this activity showed that it has three components: RNA, an RNA helicase related to the human p68 DEAD-box protein and a homologue of human T DNA glycosylase $(TDG)^{76-78}$. Thus, 5meC glycosylase activity initially detected in chicken embryo extracts was attributed to TDG. However, its excision activity against 5meC was 30-40-fold lower compared with that against T^{78} . Although TDG can flip C and C analogues into its active site, it does not possess the catalytic power to break the N-glycosidic bond⁷⁹. It should be noted that the excision activity of TDG against 5meC is stimulated by the presence of both RNA and the RNA helicase⁷⁸. Similarly, both DNMT3A and DNMT3B have been reported to interact with and stimulate the enzymatic activity of $TDG^{80,81}$. Future work should determine whether these interactions have an effect on substrate preference in vitro and whether loss of function of TDG has an effect on DNA methylation status in vivo.

In addition to TDG, the methyl-CpG-binding protein MBD4 has glycosylase activity against 5meC, but again this activity is 30-40-fold lower than its T•G mismatch glycosylase activity82. Not surprisingly, MBD4-null zygotes exhibit normal demethylation of the zygotic paternal pronucleus⁸³, and MBD4-null mice have an increased number of C to T mutations regardless of whether the C is methylated or not 84 . Despite its unfavourable biochemical properties, MBD4 was reported to carry out active DNA demethylation of the CYP27B1 promoter in response to PTH52. Interestingly, phosphorylation by protein kinase C enhanced MBD4 glycosylase activity against $5meC^{52}$, which may partially explain earlier enzymatic studies showing MBD4's preference for C over 5meC⁸⁵.

Deamination of 5meC to T followed by BER

DNA demethylation can also be achieved by deamination of 5meC to produce T, followed by BER to replace the mismatched T with unmethylated C (FIG. 4b). Both cytidine deaminases and DNMTs have been proposed to carry out the first step of this mechanism. on deamination of 5meC, T glycosylases such as TDG and MBD4 (see above) may function by repairing the mismatch.

Cytidine deaminases are important players in diverse biological processes such as the generation of antibody diversity, RNA editing and retroviral defence 86 . These processes require the production of mutations in DNA and RNA, which is achieved, in part, through the deamination of cytidine to uridine by the activation-induced deaminase (AID) and apolipoprotein B mRNA editing enzyme, catalytic polypeptide (APOBEC) family of proteins. APOBEC1, the founding member of this family, is involved in editing apolipoprotein B pre-mRNA $87,88$. The related deaminase AID was discovered to be essential for somatic hypermutation and class switch recombination of immunoglobulin genes in B

cells89,90. Consistent with its role in the diversification of antibodies, AID-deficient mice are viable and fertile and significant phenotypic abnormalities are seen only in B cells^{89,90}.

Despite the lack of developmental defects in AID-knockout mice, both AID and APOBEC1 have been shown in vitro and in an E. coli assay to have the capacity to deaminate 5meC to T in the context of single-stranded DNA^{91} . AID and APOBEC1 are also expressed in mouse oocytes, ES cells and PGCs, which may be a consequence of their genomic location in a cluster of pluripotency genes that include nanog and stella (also known as DPPA3 and $PGC7⁹¹$. Nevertheless, expression of AID in PGCs and the early embryo points to a possible role in global DNA demethylation. Indeed, a recent large-scale bisulphite sequencing study indicates that DNA methylation levels of male and female PGCs derived from AID-null embryos increased about 4% (from 18% to 22%) and 13% (from 7% to 20%), respectively, when compared to their wildtype counterpart⁹², suggesting that AID may contribute to PGC demethylation. However, because the DNA methylation levels in AID-null PGCs (~20%) are still relatively low compared with ES or somatic cells (70-80%), considerable demethylation still occurs in the absence of AID, indicating that other factors responsible for PGC demethylation remain to be identified.

Nevertheless, studies in zebrafish embryos have suggested that Aid, Mbd4 and the DNA repair protein Gadd45a (growth arrest and DNA-damage-inducible 45α) can cooperate in demethylating a methylated DNA duplex 93 . In this study, when a methylated linear dsDNA of ~740 bp was injected into a zebrafish embryo, demethylation of the injected DNA was seen when Aid and Mbd4 were co-expressed. The authors postulated that Aid de aminated 5meC, allowing Mbd4 to excise the T•G mismatch. Indeed, the T•G mismatch was detected using a PCR strategy, but only when Aid was expressed with a catalytic mutant of Mbd4 because the wild-type version excised the mismatch too quickly for it to be detected. Furthermore, when Aid and Mbd4 were titrated to levels that did not cause demethylation, the inclusion of Gadd45a elicited demethylation, indicating that these three proteins act cooperatively⁹³.

Although the above studies have provided some evidence that AID may contribute to mammalian DNA demethylation, decisive biochemical and genetic evidence supporting a major role in this process is still lacking. Biochemically, AID can act on 5meC in the context of single-stranded DNA but not dsDNA 91 . Genetically, AID-knockout mice exhibit the expected B cell and immunological defects^{89,90}, but no gross developmental or reproductive defects. Similarly, APOBEC1-knockout mice are also viable and fertile^{94,95}. Although genetic redundancy may be a possible cause of the lack of expected developmental and reproductive phenotypes, such explanation needs to be confirmed by generating combinational knockouts. Furthermore, because DNA methylation occurs symmetrically, deamination of both strands would give rise to a TG•GT double mismatch. There is no evidence indicating that either TDG or MBD4 can use a double mismatch as a substrate. Furthermore, processing of a double mismatch by the AP endonuclease would generate a DNA double-strand break. This would put tremendous pressure on the repair machinery if such a mechanism were used for global demethylation. However, for locus-specific DNA demethylation, such a mechanism would not present a big problem.

In addition to AID and APOBEC, DNMTs have recently been implicated in 5meC deamination, even though they are commonly known for their ability to catalyse DNA methylation. Evidence indicating their involvement in the deamination process initially came from studies in bacteria where the methyltransferases \underline{M} . HpaII⁹⁶⁻⁹⁸ and \underline{M} . Ecor $\Pi^{99,100}$ were shown to possess deaminase activities. Consistent with bacterial studies, the mammalian counterparts, DNMT3A and DNMT3B, have also been shown to possess deaminase activity in vitro⁵¹. As discussed above, the promoters of oestrogen-responsive

genes undergo cyclical rounds of methylation and demethylation. Thus, the participation of DNMT3A and DNMT3B in both methylation and demethylation would facilitate rapid transcriptional cycling (FIG. 3b). Interestingly, ERα associates with and stimulates the activity of TDG^{101,102}, allowing for the repair of the T \bullet G mismatch. DNMT3A and DNMT3B also associate with TDG and this interaction stimulates glycosylase activity $80,81$. Indeed, DNA demethylation was found to coincide with the recruitment of TDG and other BER enzymes⁵¹.

However, it is surprising that DNMTs possess two opposing enzymatic activities. Although the methyltransferase activity of DNMT3A is inhibited by $T\text{DG}^{81}$, the 5meC deamination reaction can only occur under conditions where SAM concentrations are very low or nonexistent⁵¹. In order for DNMT3A to carry out efficient methylation and demethylation during transcriptional cycling, levels of SAM must fluctuate rapidly. Given that SAM is crucial for many essential biochemical and biological processes, it is difficult to imagine how this could be achieved without serious biological consequences.

Nucleotide excision repair

Another DNA repair pathway, NER, has also been proposed to carry out DNA demethylation. This type of repair is generally used to repair DNA containing bulky lesions, which form after exposure to chemicals or radiation. After damaged DNA is recognized, dual incisions flanking the lesion are made and a 24-32 nucleotide oligomer is released. The resulting gap is then filled in by repair polymerases and sealed by a ligase⁶⁴.

In an assay aimed at identifying proteins required for activation of a reporter that is silenced by DNA methylation, Niehrs and colleagues uncovered a novel function for GADD45A 103 , which is encoded by a p53- and breast cancer type 1 susceptibility protein (BRCA1)inducible gene and participates in diverse biological processes, including DNA damage response, cell cycle progression, apoptosis and NER¹⁰⁴. Overexpression of GADD45A in mammalian cell lines leads to loci-specific and global demethylation, whereas knockdown results in DNA hypermethylation¹⁰³. Because GADD45A had previously been implicated in $NER^{105,106}$, Barretto *et al.* explored the role of NER in DNA demethylation and found that loss of DNA methylation is accompanied by DNA synthesis and requires the NER endonuclease xeroderma pigmentosum group G-complementing protein (XPG), which interacts with GADD45A¹⁰³. The recruitment of GADD45A and other components of the NEr repair machinery to ribosomal RNA (rRNA) genes is facilitated by TBP-associated factor 12 (TAF12) and leads to DNA demethylation and rRNA gene activation¹⁰⁷. However, it is not clear how the demethylation process is initiated and whether GADD45A is directly involved. More importantly, two independent studies have raised doubt on the role of GADD45A in the active DNA demethylation process. In the first study, the Pfeifer group carried out a series of experiments that were similar to those carried out by the Niehrs group, but obtained no evidence indicating that GADD45A had any effect on DNA methylation¹⁰⁸. In the second study, analysis of the GADD45A-null mice indicated that loss of GADD45A function had neither loci-specific nor global effects on DNA methylation levels¹⁰⁹.

GADD45B, another member of the GADD45 family, has also been implicated in active demethylation of genes that are crucial for adult neurogenesis¹¹⁰. Loss of GADD45B results in defects in neural progenitor proliferation and dendritic growth. This was attributed to promoter hypermethylation and the repression of BDNF and fibroblast growth factor 1 $(FGFI)$, two genes crucial for neurogenesis¹¹⁰. However, GADD45B is not involved in zygotic DNA demethylation as GADD45B-null zygotes undergo normal paternal genome demethylation¹¹¹. Because GADD45B has not been biochemically characterized, it is unknown whether it is directly involved in the active demethylation of neurogenesis genes.

Oxidative demethylation

Another possible mechanism by which DNA demethylation can be carried out is through oxidative demethylation. The $E.$ coli enzyme \overline{AlkB} is a member of the 2-oxoglutarate (2OG)-dependent dioxygenases and is involved in the bacterial response to alkylation damage to DNA. using oxygen, iron and 2OG as cofactors, AlkB is able to carry out oxidative demethylation of 1-methyladenine and 3meC by releasing the methyl group as formaldehyde^{112,113}. The same mechanism is used by the JmjC family of enzymes to demethylate histon e substrates^{60,114}.

Although breakage of a carbon-carbon bond is energetically difficult, enzymes that catalyse such reactions do exist. As shown in FIG. 5a, thymine 7-hydroxylase can catalyse the conversion of T to iso-orotate through three consecutive oxidation reactions using oxygen, iron and 2OG as cofactors¹¹⁵. Iso-orotate can be further converted to C through a decarboxylation reaction. Although thymine 7-hydroxylase and iso-orotate decarboxylase have been isolated from fungi, such as Rhodotorula glutinis, Neurospora crassa and Aspergillus nidulans⁶², no homologue of thymine 7-hydroxylase has been found in mammals. Interestingly, the trypanosome base J-binding proteins, JBP1 and JBP2, have properties similar to that of thymine 7-hydroxylase 116,117, prompting the Rao group to search for mammalian homologues with similarity to the dioxygenase domains of the JBP proteins. This effort led to the identification of the ten-eleven translocation (TET) family of proteins¹¹⁸, we have also independently characterized the mouse TET family¹¹⁹.

TET1, the founding member of the TET family, was initially discovered in acute myeloid leukaemia (AMI) as a fusion partner of the histone H3 Lys4 methyltransferase $MLL^{120,121}$. Subsequent studies *in vitro* and in cultured cells showed that human TET1 is capable of hydrolysing 5meC to produce 5-hydroxymethylcytosine (5hmC) in DNA118. Similarly, all three members of the mouse TET family possess this enzymatic activity¹¹⁹. Consistent with the presence of a dioxygenase domain in the proteins and the predicted reaction mechanism, the putative iron-binding sites are required for their enzymatic activities^{118,119}. Furthermore, TET1 is capable of acting on both fully methylated and hemi-methylated DNA^{118} .

Although 5hmC has previously been reported to exist in animal DNA122, this modified base is not found in some cell types and tissues^{118,123}, thus raising the question of whether 5hmC is present in mammalian DNA at physiologically relevant levels. This issue was directly addressed in two cell types. In Purkinje neurons, 5hmC is ~40% as abundant as $5 \text{me} C^{124}$, whereas the frequency of 5hmC in ES cells was estimated to be approximately 1 in every $3,000$ nucleotides¹¹⁸. Thus, it is evident that 5hmC constitutes a large fraction of mammalian DNA in some cell types.

The consequences of 5hmC in genomic DNA are currently unclear. Because 5hmC seems to be stable, it may function like other modifications by altering local chromatin structure or contributing to the recruitment or exclusion of other factors that influence transcription. For example, the transcriptional repressors MeCP2, MBD1, MBD2 and MBD4 bind to methylated DNA, but do not recognize 5hmC125,126. It is also possible that the TET proteins may facilitate passive demethylation in dividing cells such as ES cells as 5hmC is not recognized by DNMT1 (REF. 127); thus, newly replicated DNA would not maintain patterns of methylation. Alternatively, 5hmC may be an intermediate in an active demethylation pathway that ultimately leads to the replacement of 5meC with C (FIG. 5b). This could be achieved by several ways that include: BER by a 5hmC-specific DNA glycosylase (as such activity has been previously reported to exist in calf thymus extracts¹²⁸), deamination of 5hmC to generate 5hmu followed by BER initiated by a 5hmuspecific glycosylase such as single-strand-selective monofunctional u DNA glycosylase 1 $(SMUG1)¹²⁹$, conversion of 5hmC to C through loss of formaldehyde on ultraviolet light

exposure¹³⁰ or high pH¹³¹ (or possibly carried out by DNMTs)¹³², and two consecutive oxidation steps followed by decarboxylation similar to that used by the thymidine salvage pathway (FIG. 5a). It is not clear why TET proteins cannot catalyse consecutive reactions such as that of thymine 7-hydroxylase. Because all *in vitro* assays carried out so far used recombinant TET proteins alone, it is possible that association of TET proteins with their in vivo partners is necessary to confer such a capability. In this case, a decarboxylase may eventually remove the carboxyl group to complete the demethylation process.

Consistent with the relative enrichment of 5hmC in ES cells, recent studies have shed light on the role of TET1 in ES cell biology. During ES cell differentiation, TET1 mRNA levels decline, coinciding with a decrease in 5hmC levels¹¹⁸, which suggests that TET1 may be important for ES cell identity. Indeed, knockdown of TET1, but not TET2 or TET3, in mouse ES cells results in impairment of ES cell self-renewal and maintenance¹¹⁹. Analysis of the differentiated TET1-knockdown ES cells revealed a bias towards the trophoectoderm and primitive endoderm lineages. Furthermore, knockdown of TET1 at two-cell stage embryos followed by cell lineage tracing revealed that the knockdown cells are biased towards the trophoectoderm¹¹⁹, indicating that TET1 is required for inner cell mass cell specification. Consistent with its role in ES cell self-renewal and maintenance, knockout of TET1 resulted in embryonic lethality (K. Hong and Y.Z., unpublished observations), making the evaluation of the role of TET1 on global demethylation of the paterna l genome difficult.

With regard to the mechanism underlying the role of TET1 in ES cells, TET1 maintains nanog expression in ES cells by directly binding to the nanog promoter and protecting it from becoming hypermethylated, as knockdown of TET1 in ES cells resulted in downregulation of nanog expression concomitant with increased nanog promoter methylation¹¹⁹. Nanog seems to be one of the main TET1 targets as the phenotypes associated with TET1 knockdown can largely be rescued by ectopic expression of nanog¹¹⁹.

Although TET2 is also expressed in ES cells, it seems that TET2 does not play a significant part in ES cell biology as knockdown of TET2 does not confer any obvious phenotype ¹¹⁹. However, a flurry of recent studies have uncovered that dysfunction of human TET2 may be a key event in leukaemogenesis as human TET2 is mutated in a range of human myeloid malignancies, including myelodysplastic syndromes (MDSs), myeloproliferative disorders (MPDs) and acute myeloid leukaemias $(AMls)^{133-140}$. Currently, *TET2* is the most frequently mutated gene that has been identified in patients with MDS and these mutations have been suggested to occur early during the pathogenesis of the disease¹³⁷. Consistent with a role for TET2 in regulating DNA demethylation, aberrant DNA methylation is frequently found in patients with $MDS¹⁴¹$. Indeed, mutations of TET2 that mimic mutations identified in patients with MDS abolished the enzymatic activity of TET2 (A. C. D'Alessio and Y.Z., unpublished observations). Furthermore, the DNA methyltransferase inhibitor 5 azacytidine (5-azaC) has been shown to be an effective treatment for patients with high-risk MDS and secondary $AML^{142,143}$, indicating that aberrant DNA methylation plays a crucial part in MDS development and progression. The participation of TET2 in DNA demethylation may provide a molecular basis for the effectiveness of using methyltransferase inhibitors in the treatment of patients with MDS, thus setting the stage for understanding the molecular mechanism underlying the pathogenesis of leukaemias.

Radical SAM mechanism

Although many proteins have been proposed to carry out active DNA demethylation, none of the proteins discussed above have been shown to have a role in paternal genome demethylation in zygotes. To identify proteins involved in paternal genome demethylation, our laboratory used a candidate gene knockdown approach coupled with live-cell imaging. To facilitate a screen of candidate proteins, we developed a probe that consists of the Cys-X-

X-Cys domain of MLL fused to enhanced green fluorescent protein (EGFP). Because the Cys-X-X-Cys domain has high affinity for unmethylated CpG^{144} , injection of mRNA encoding the probe into zygotes results in the accumulation of the probe at the demethylated paternal pronucleus¹¹¹, allowing live-cell imaging of the paternal genome demethylation process. Using this imaging system, we screened several candidate proteins by injecting small interfering RNAs (siRNAs) against each of the candidates into eggs before carrying out intracytoplasmic sperm injection (ICSI), and monitored the effect of the siRNA on the accumulation of the probe at the paternal pronucleus. This screen uncovered a role for elongator complex protein 3 (ELP3) in paternal genome demethylation. ELP3 knockdown prevented the accumulation of the probe in the paternal pronucleus at pronuclear stages 4 and 5 (REF. 111). In addition, immunostaining and bisulphite sequencing of selected retrotransposon elements further support a role for ELP3 in paternal genome demethylation¹¹¹.

ELP3 is a member of the core elongator complex (ELP1-ELP3), which combines with another subcomplex (ELP4-ELP6) to form the holo-elongator complex^{145,146}. Because knockdown of the ELP1 and ELP4 components also impaired paternal genome demethylation, it is likely that the entire elongator complex may be involved in the demethylation process¹¹¹. Interestingly, the Fe-S radical SAM domain of ELP3, but not the histone acetyltransferase (HAT) domain, is required for paternal genome demethylation 111 . Although this may provide a clue regarding the enzymatic mechanism of ELP3, recent studies in yeast suggest that the Cys-rich domain of Elp3 is required for the integrity of the elongator complex^{147,148}, raising the possibility that the Fe-S radical SAM motif may have a structural rather than an enzymatic role. Thus, direct biochemical evidence of the enzymatic activity of the elongator complex and genetic evidence using ELP3-null oocytes remain to be shown.

Interestingly, a recent study confirmed the presence of an Fe-S cluster in the bacteria *Methanocaldococcus jannaschii* Elp3 protein¹⁴⁹. The assumption that mammalian ELP3 is a radical SAM protein has led to a potential mechanism for ELP3-catalysed DNA demethylation as outlined in FIG. 6 (S. J. Booker, personal communication). like every radical SAM enzyme, the reaction is initiated by the generation of a powerful oxidizing agent, the 5′-deoxyadenosyl (5′-dA) radical, from SAM. The 5′-dA radical could extract a hydrogen atom from the 5-methyl group to generate a 5meC radical. In the next step, an electron is returned back to the Fe-S cluster to generate a third intermediate, which can be converted to the relatively stable 5hmC by the addition of a water molecule. In order to break the carbon-carbon bond, the next step requires the generation of an intermediate, the resulting carbanion of which would be stabilized. This can probably be achieved by a thymidylate synthase or methyltransferase type of mechanism, whereby a Cys residue carries out a nucleophilic attack at carbon 6, leading to the release of formaldehyde. In the absence of an external nucleophile, an alternative pathway leading to the release of formaldehyde can also take place. Finally, an elimination at the formaldehyde release step results in the final product of C.

Although future studies are required to validate or refute this proposed mechanism, we note that this work is not trivial for three reasons. First, the identities of mammalian ELP5 and ELP6 still need to be determined as an apparent orthologue of yeast Elp5 and Elp6 cannot be identified by sequence homology searches. Second, the radical SAM reaction occurs under anaerobic conditions and reconstitution of the elongator complex under anaerobic conditions is challenging. Finally, given that zygotic DNA demethylation occurs only on the paternal genome, some unique features of the paternal genome may be required in order for it to serve as a substrate. Despite these challenges, identification of the elongator complex as an

important factor for paternal genome demethylation in zygotes allows for further studies towards understanding the molecular mechanisms of active DNA demethylation.

Concluding remarks

Observations of active DNA demethylation during embryonic development and in somatic cells have opened the door for many questions to be answered. In particular, how DNA demethylation is achieved in mammalian cells remains debatable as no single enzyme or mechanism has gained decisive biochemical and genetic support (see Supplementary information S1 (table)). It is possible that multiple mechanisms exist to carry out DNA demethylation and that the use of each one is dictated by the specific biological context.

Although repair-based mechanisms, particularly deamination of 5meC followed by BER, have offered an attractive mechanism for active DNA demethylation, genetic evidence is still lacking. Furthermore, the involvement of a repair-based mechanism in global DNA demethylation would put tremendous pressure on the repair machinery when considering that paternal pronucleus demethylation is completed within 4 hours^{17,18}.

Although AID seems to contribute to active demethylation in PGCs, it is only responsible for a small part of it as considerable demethylation still takes place in the AID-null $PGCs^{92}$. Nevertheless, this mechanism does provide a reasonable explanation for loci-specific demethylation in response to gene-activation signals. Although AID deficiency has some effect on PGC demethylation, there is no evidence that it affects paternal DNA demethylation in zygotes. Similarly, MBD4-null zygotes still experience paternal genome demethylation⁸³. It seems that although repair-based mechanisms may be responsible for loci-specific DNA demethylation and partial demethylation in PGCs, their role in zygotic paternal genome demethylation is less likely. To date, the only factor shown to have a role in zygotic paternal genome demethylation is the elongator complex, although it is unclear whether its role is direct or indirect¹¹¹. Future work should focus on gaining additional genetic evidence using elongator-null zygotes and elucidating its enzymatic activity. The recent demonstration that the TET family proteins are capable of catalysing the conversion of 5meC to 5hmC has raised the possibility that these proteins may have a role in active DNA demethylation^{118,119}, we anticipate that work evaluating their role in demethylation of the zygotic paternal genome and PGCs is forthcoming. Furthermore, analysis of the fate and function of 5hmC will also attract a lot of attention.

In addition to determining the mechanism of active demethylation, one open question that remains is to what extent the paternal genome and PGCs are demethylated. Although this event is considered to be global, as determined by 5meC immunostaining, it is evident that some regions of the paternal genome are protected from this wave of demethylation. The advent of high-throughput analyses including chromatin immunoprecipitation-on-chip (ChIP-chip), ChIP sequencing (ChIP-Seq) and bisulphite sequencing (BS-Seq; bisulphite treatment followed by high-throughput sequencing) has allowed for genome-wide profiling of epigenetic marks such as DNA methylation $92,150-153$. using single-molecule, realtime sequencing, a recent study showed the feasibility of direct detection of modified nucleotides in DNA, including N6-meA, 5mC and 5hmC154. Future studies using these tools will undoubtedly determine precisely which genomic regions are demethylated and which regions are protected. However, improvements in the sensitivity of these technologies will be necessary for such experiments, given that paternal genomic DNA would need to be obtained from individual zygotes.

As well as being fundamental to our knowledge in epigenetics, a better understanding of how DNA demethylation occurs will allow for the development of techniques and approaches that will improve somatic cell reprogramming (BOX 2) and cancer treatment.

Tumour suppressor gene silencing by promoter DNA methylation is thought to play an important part in cancer development¹⁵⁵. Consistently, inhibitors of DNMTs have been used in the treatment of certain cancers¹⁵⁶. Owing to the reversible nature of epigenetic modifications, developing drugs that target epigenetic factors is becoming one of the top priorities for many biotechnology and pharmaceutical companies¹⁵⁷. It is anticipated that targeted demethylation of tumour suppressor genes may reactivate the silenced tumour suppressor genes, which can lead to cellular differentiation or halt uncontrolled cell proliferation.

The mechanism underlying the regulation of DNA methylation is a question that has elicited much attention and controversy over the past decade. Although recent studies have proposed numerous ideas as to how active DNA demethylation is carried out, many aspects are still contentious and a consensus has yet to be achieved. with the development of new technology and the studies described above, our continued and collective efforts in this field will hopefully provide clearer answers in the coming decade.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Glossary

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Box 1

Protection of the maternal genome from demethylation

Whereas the paternal genome undergoes extensive demethylation, the maternal genome remains methylated even though it is exposed to the same cytoplasmic factors. This may be due to a mechanism that protects the maternal genome from this wave of demethylation or to a putative DNA demethylase that is specifically recruited to the paternal genome.

Sperm DNA is packaged with protamines, which are exchanged for canonical and noncanonical histones on fertilization. Interestingly, deposition of the histone variant H3.3 occurs asymmetrically, with a strong preference for the paternal pronucleus^{158,159}. This raises the possibility that asymmetric H3.3 deposition may trigger the paternal genome-specific demethylation process. Asymmetric patterns of histone modifications have also been seen in the maternal and paternal pronuclei and may also contribute to the asymmetric demethylation process. For example, methylation, dimethylation and trimethylation at H3 Lys27 (H3K27me1, H3K27me2 and H3K27me3, respectively) and at Lys9 (H3K9me2 and H3K9me3) are clearly seen in the maternal pronucleus of zygotes, but are virtually undetectable in the paternal pronucleus¹⁵⁹⁻¹⁶⁴. Thus, the maternal genome may use a protective mechanism against demethylation by carrying specific histone variants or modifications.

Alternatively, a recent study has suggested that non-histone factors present in the oocyte might protect the maternal genome from demethylation¹⁶⁵. Zygotes lacking stella (also known as *DPPA3* and *PGC7*), a maternal effect gene required for early development¹⁶⁶, exhibited demethylation of both pronuclei. Although stella can directly bind DNA in *vitro*, it seems to lack specificity for methylated DNA^{165} . Therefore, how stella protects the maternal genome from demethylation remains to be determined.

Box 2

Implications of active DNA demethylation in reprogramming

Induced pluripotent stem (iPS) cells can be generated by introducing four transcription factors — octamer-binding protein 3 (OCT3; also known as OCT4 and POU5F1), SRY box-containing factor 2 (SOX2), krüppel-like factor 4 (KLF4) and MYC — into somatic cells^{167,168}. Successful reprogramming requires the activation of endogenous $OCT4$ and nanog genes, which are known to be silenced by DNA methylation in somatic cells¹⁶⁹⁻¹⁷². Demethylation of the *OCT4* and nanog promoters is thus an integral event in iPS cell generation¹⁷³. In fact, inefficient DNA demethylation is thought to be one of the causes of the low efficiency in iPS cell generation because the use of the DNA methylation inhibitor 5-azacytidine can increase the efficiency of iPS cell generation by converting partially reprogrammed cells to fully reprogrammed iPS cells¹⁷³.

Transcription factor-based iPS cell generation is a slow process compared to somatic cell nuclear transfer (SCNT) and cell fusion^{174,175}. One possible explanation for this difference may be the mechanisms used to reactivate endogenous *OCT4* and nanog. Epigenetic reprogramming of somatic cells to pluripotent iPS cells may necessitate several cell divisions¹⁷⁶ owing to the absence of the DNA demethylase or demethylases required for demethylation of the OCT4 and nanog promoters. By contrast, reactivation of OCT4 and nanog can occur quickly during SCNT and cell fusion because the DNA demethylase or demethylases may already be present in eggs and embryonic stem (ES) cells. Consistent with this notion, reprogramming by cell fusion requires activationinduced deaminase (AID)-dependent demethylation and reactivation of OCT4 and nanog¹⁷⁷. Surprisingly, although AID was present at the *OCT4* and nanog promoters in fibroblasts, these promoters are methylated, suggesting that other factors or regulatory events are required for demethylation. Given that genetic evidence does not support an important role for AID in ES cells (see main text), it is unclear whether AID directly participates in promoter demethylation of these genes during somatic cell reprogramming. Regardless, it is evident that activation of pluripotent genes through DNA demethylation is an important step during the somatic cell reprogramming process. Identification and characterization of the enzymes involved should improve protocols of somatic cell reprogramming.

Figure 1. Mechanisms of DNa methylation and demethylation

During early development, methylation patterns are initially established by the *de novo* DNA methyltransferases DNMT3A and DNMT3B. When DNA replication and cell division occur, these methyl marks are maintained in daughter cells by the maintenance methyltransferase, DNMT1, which has a preference for hemi-methylated DNA. If DNMT1 is inhibited or absent when the cell divides, the newly synthesized strand of DNA will not be methylated and successive rounds of cell division will result in passive demethylation. By contrast, active demethylation can occur through the enzymatic replacement of 5 methylcytosine (5meC) with C.

Figure 2. Dynamics of DNa methylation during development

a | Active demethylation in the zygotic paternal genome. Shortly after a sperm fertilizes an egg, the paternal genome rapidly undergoes genome-wide active DNA demethylation and remains demethylated following multiple rounds of cell division. During this time, the maternal genome experiences gradual, passive demethylation. De novo methylation patterns are established by the DNA methyltransferases DNMT3A and DNMT3B during the development of the blastocyst. **b** | Active demethylation in primordial germ cells (PGCs). After implantation of the blastocyst at embryonic day 7.5 (E7.5), the extraembryonic ectoderm (ExE) and visceral endoderm (VE) produce signals that specify a subset of epiblast cells (Epi) to become PGCs. This process requires two key transcription factors, BLIMP1 (also known as PR domain zinc finger protein 1 (PRDM1)) and PDRM14, which are expressed during this stage of development. Following specification, PGC founder cells divide in the presence of the DNA methyltransferase DNMT1 and migrate towards the genital ridge. During this migration and on arrival at the genital ridge, 5-methylcytosine (5meC) is erased through an active mechanism. ICM, inner cell mass; TE, trophectoderm.

a | Active demethylation at the brain-derived neurotrophic factor (*BDNF*) promoter. In neurons, BDNF is maintained in a repressed state through DNA methylation and binding of the repressive methylcytosine (meC)-binding protein MeCP2. On depolarization with KCl, DNA methylation and MeCP2 binding are lost, concomitant with increased BDNF expression. This demethylation event is considered to be active because it occurs in postmitotic neurons. **b** | Active demethylation at nuclear receptor target promoters. The promoter of the oestrogen receptor (ER) target gene $pS2$ (also known as *TFF1*) undergoes cyclical rounds of methylation and demethylation that correspond to the repression and expression of the gene, respectively. Transcriptional activation of $pS2$ occurs in the presence of oestrogens (E2) and coincides with demethylation of the promoter. This is achieved by deamination of 5meC by DNA methyltransferase 3 (DNMT3) followed by base excision repair (BER) of the T•G mismatch by T DNA glycosylase (TDG). To revert to repression, DNMT3 remethylates the promoter. Although DNMT3 is involved in both methylation and demethylation, it is important to note that DNMT3 can only carry out the deamination step in the absence or at low concentrations of the methyl donor S-adenosylmethionine (SAM).

Figure 4. Base excision repair-based mechanisms for DNa demethylation

a | Base excision repair (BER) through direct excision of 5-methylcytosine (5meC). Initiation of the BER pathway can be carried out by a glycosylase that directly excises 5meC to generate an abasic (apurinic and apyrimidinic (AP)) site. The DNA backbone is nicked by an AP lyase (or by the glycosylase itself if it is bifunctional). The 3′ sugar group is then cleaved by an AP endonuclease and the resulting single nucleotide gap is filled in with an unmethylated C by an unknown polymerase and ligase. It has been well established in plants that the demeter (Dme; also known as repressor of silencing 1 (ROS1)) family of enzymes can carry out the 5meC glycosylase reaction, but to date no mammalian enzymes have been reported to be capable of carrying out this step efficiently. **b** | Deamination of 5meC followed by BER. In contrast to direct excision of 5meC, deamination of 5meC produces T, which can be repaired by BER by a T•G mismatch glycosylase such as T DNA glycosylase (TDG) or methyl-CpG-binding domain protein 4 (MBD4) to regenerate an unmethylated C. DNMT, DNA methyltransferase.

Figure 5. Oxidative demethylation by TET proteins

a | Part of the thymidine salvage pathway. Direct removal of the methyl group of 5 methylcytosine (5meC) involves breaking a carbon-carbon bond, which requires an enzyme with great catalytic power. Such an enzyme exists in the thymidine salvage pathway. Starting with T, thymine-7-hydroxylase (THase) carries out three consecutive hydroxylation reactions to produce iso-orotate, which is processed by a decarboxylase to produce U. A similar mechanism may be used in active DNA demethylation, particularly by the ten-eleven translocation (TET) family of proteins. **b** | The fate of 5-hydroxymethylcytosine (5hmC). The TET family of proteins catalyses the conversion of 5meC to 5hmC, which may be an intermediate that can be further processed by one of the following mechanisms. BER may be initiated by a 5hmC glycosylase (1); 5hmC may undergo deamination to produce 5hmU (2), which is repaired by BER through a 5hmU glycosylase such as SMUG1 (single-strandselective monofunctional U DNA glycosylase 1); 5hmC may directly be converted to C by DNA methyltransferases (DNMTs), ultraviolet (UV) exposure or high pH (3); or consecutive hydroxylation reactions followed by a decarboxylation reaction similar to the thymidine salvage pathway may be used to ultimately replace 5hmC with C (4). Alternatively, 5hmC itself may be a functional modification. α-KG, α-ketoglutarate.

Figure 6. Proposed mechanism for elP3-mediated DNa demethylation

Mammalian elongator complex protein 3 (ELP3) contains an Fe-S radical Sadenosylmethionine (SAM) domain that is important for active DNA demethylation of the zygotic paternal genome. If ELP3 is indeed a functional radical SAM protein, it may directly carry out DNA demethylation through the following mechanism. First, ELP3 uses SAM to generate a 5′-deoxyadenosyl radical, which could extract a hydrogen atom from the 5 methyl group of 5-methylcytosine (5meC; 1) to form a 5meC radical (2). After an electron is donated back to the Fe -S to create the third intermediate (3), a water molecule would promote the formation of 5-hydroxymethylcytosine (5hmC) (4). A nucleophilic attack at carbon 6 can result in the carbon-carbon bond breaking to release formaldehyde (5-7). In the absence of an external nucleophile, an alternative pathway $(4'-6')$ that leads to the release of formaldehyde can also take place. Finally, an elimination step would produce an end product of C (8).