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Mitochondrial genome undergoes de novo DNA methylation that protects mtDNA against oxidative damage during the peri-implantation window

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Mitochondrial remodeling during the peri-implantation stage is the hallmark event essential for normal embryogenesis. Among the changes, enhanced oxidative phosphorylation is critical for supporting high energy demands of postimplantation embryos, but increases mitochondrial oxidative stress, which in turn threatens mitochondrial DNA (mtDNA) stability. However, how mitochondria protect their own histone-lacking mtDNA, during this stage remains unclear. Concurrently, the mitochondrial genome gain DNA methylation by this stage. Its spatiotemporal coincidence with enhanced mitochondrial stress led us to ask if mtDNA methylation has a role in maintaining mitochondrial genome stability. Herein, we report that mitochondrial genome undergoes de novo mtDNA methylation that can protect mtDNA against enhanced oxidative damage during the peri-implantation window. Mitochondrial genome gains extensive mtDNA methylation during transition from blastocysts to postimplantation embryos, thus establishing relatively hypermethylated mtDNA from hypomethylated state in blastocysts. Mechanistic study revealed that DNA methyltransferase 3A (DNMT3A) and DNMT3B enter mitochondria during this process and bind to mtDNA, via their unique mitochondrial targeting sequences. Importantly, loss- and gain-of-function analyses indicated that DNMT3A and DNMT3B are responsible for catalyzing de novo mtDNA methylation, in a synergistic manner. Finally, we proved, in vivo and in vitro, that increased mtDNA methylation functions to protect mitochondrial genome against mtDNA damage induced by increased mitochondrial oxidative stress. Together, we reveal mtDNA methylation dynamics and its underlying mechanism during the critical developmental window. We also provide the functional link between mitochondrial epigenetic remodeling and metabolic changes, which reveals a role for nuclearmitochondrial crosstalk in establishing mitoepigenetics and maintaining mitochondrial homeostasis.

mitochondrial DNA | mitochondrial oxidative damage | de novo DNA methylation | DNMT3A/3B | peri-implantation

Mitochondrial remodeling is essential for early embryonic survival and development (1). During the transition from blastocysts to postimplantation embryos, a hallmark event is the notable increase in mitochondrial oxidative phosphorylation (OXPHOS), to support the increased energy demands of postimplantation embryos (1, 2). The developmental transition during peri-implantation is very important for successful pregnancy. In humans, \sim 75% of failed pregnancies are due to developmental defects within this stage (3). Even in mice, implantation is a critical step in establishment of pregnancy (4). During this critical window, however, enhanced OXPHOS generates excessive reactive oxygen species (ROS), which in turn would have damaged the mitochondrial genome, thus threatening embryonic survival (1, 5, 6). Compared with histone-rich nuclear DNA, naked mitochondrial DNA (mtDNA) lacks histone and is more sensitive to oxidative damage (7, 8). Despite the induction of antioxidant defense systems during peri-implantation $(\overline{6}, 9)$, how mitochondria protect their own genome during this stage remains unclear. Coinciding with the metabolic remodeling, a remarkable epigenetic change also occurs in mitochondria during this stage: mitochondrial genome may acquire DNA methylation from the hypomethylated blastocysts, to establish relative hypermethylation of mtDNA in postimplantation embryos. Considering the functional role of DNA methylation in maintaining nuclear genome integrity (10, 11), spatiotemporal coincidence of mitochondrial epigenetic and metabolic changes, lead us to ask if increased mtDNA methylation functions to protect the mitochondrial genome against ROS-induced damage.

During the peri-implantation stage, previously published (12–14), and our own high-throughput DNA methylation data (Fig. 1*D*), implied that mitochondrial genome

Significance

In early mammalian embryos, enhanced mitochondrial oxidative metabolism is a hallmark essential for postimplantation survival and development. However, resultant mitochondrial oxidative stress would have damaged mitochondrial DNA (mtDNA). Until now, how embryonic mitochondria protect their own genome remains unclear. Herein, we reported that the mitochondrial genome undergoes extensive de novo mtDNA methylation that can protect mtDNA against enhanced oxidative damage during the periimplantation stage. DNA de novo methyltransferase DNMT3A and DNMT3B enter mitochondria via their own mitochondrial targeting sequences, and then synergistically catalyze de novo mtDNA methylation. Of note, increased mtDNA methylation plays an important role in preventing oxidative stressinduced mtDNA damage. These findings advance our understanding of mitochondrial epigenetics and its role in maintaining mitochondrial homeostasis during early development.

The authors declare no competing interest. This article is a PNAS Direct Submission.

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may undergo a wave of de novo mtDNA methylation from a hypomethylated state in blastocysts, and thus establishing relatively hypermethylated state in postimplantation embryos. Despite the importance of mitochondrial homeostasis during the developmental transition, however, this possibility has never been functionally determined. The complete mitochondrial genome contains 37 genes but does not encode enzymes with the de novo methyltransferase activity (15). Previous studies based on in vitro cellular experiments, demonstrated that nuclear-encoded methyltransferases can enter mitochondria, and thus maintaining or establishing mtDNA methylation (14, 16). By contrast, the essential role of methyltransferases in catalyzing mtDNA methylation, has never been confirmed via in vivo functional analyses. In particular, whether DNA de novo methyltransferases, i.e., DNMT3A and DNMT3B can translocate into mitochondria and catalyze de novo mtDNA methylation during peri-implantation, as well as the underlying mechanism, remains largely illusive.

More importantly, enhanced OXPHOS during the transition from blastocysts to postimplantation embryos, results in the accumulation of intramitochondrial ROS (1). Given that the naked mtDNA lacks histone yet exists in an environment with high concentrations of ROS, mtDNA is very sensitive to ROS-induced damage (7, 8, 17). Spatiotemporal coincidence of enhanced mitochondrial oxidative stress and increased mtDNA methylation, is reminiscent of the nontranscriptional function of DNA methylation, i.e., maintaining genome stability (10, 11). It has been indicated that global hypomethylation is associated with increased genome instability in mammalian cells (18–20). Thus, we postulated that de novo mtDNA methylation during the developmental transition, might have a role in protecting the mitochondrial genome against oxidative damage in postimplantation embryos.

Herein, using *Dnmt3a* and *Dnmt3b* knockout mouse models, we confirmed the occurrence of de novo mtDNA methylation across the mitochondrial genome during the transition from blastocysts to postimplantation embryos, and identified the underlying mechanism: DNMT3A and DNMT3B can enter mitochondria, via their unique mitochondrial targeting sequences (MTS), and catalyze this process. In vivo and in vitro loss- and gain-of-functional analyses indicated that de novo mtDNA methylation contributes essentially to protect the mitochondrial genome against oxidative damage. Our results reveal the mtDNA methylation dynamics and its physiological role in early embryos, thus presenting insight into the crosstalk between nuclear and mitochondrial genomes that establishes mitoepigenetics, thus maintaining mitochondrial homeostasis.

Results and Discussion

Spatiotemporal Coincidence of De Novo mtDNA Methylation and Mitochondrial Oxidative Stress during the Peri-Implantation Window. Reanalysis results of our previously published transcriptomic data (6, 21), showed the induction of genes controlling OXPHOS during the developmental transition from preimplantation blastocysts (embryonic day [E] 3.5, E3.5) to postimplantation epiblasts (E7.5) (*SI Appendix*, Fig. S1 *A* and *B*). In line with this, OXPHOS-dependent adenosine triphosphate (ATP) production, and its contribution to total ATP content, significantly increased during this stage (Fig. 1*A*). Accordingly, a significantly higher mitochondrial membrane potential, which maintains OXPHOS activity, can be detected in E7.5 epiblasts (Fig. 1*B*). Along with this, intramitochondrial ROS accumulation detected by MitoSOX, a mitochondria-specific probe (Fig. 1*C*), were significantly elevated during the developmental transition. These observations were in line with the up-regulation of genes related to mtDNA repair (*SI Appendix*, Fig. S1*C*), suggesting that mitochondria suffer from considerable oxidative stress during this stage.

Coinciding with notable changes in mitochondrial metabolism during the peri-implantation window, remarkable epigenetic remodeling also occurs in mitochondria. To systematically characterize its dynamics and features, we profiled mtDNA methylation patterns of mouse E3.5 blastocysts, E7.5 epiblasts, and E10.5 embryos using methylated DNA immunoprecipitationsequencing (MeDIP-seq). These time points represent the developmental windows of de novo DNA methylation of the nuclear genome (22-24). Notably, distinct from that in the nuclear genome, 5hmC enrichment in the mitochondrial genome is up to 100-fold higher relative to 5mC enrichment (16). Therefore, bisulfite-based methods are thought to be technically unfavorable to detect dynamic changes of 5mC in mtDNA, because 5mC is indistinguishable from 5hmC by bisulfite modification (25). In addition, the accuracy of bisulfite sequencing to detect mtDNA methylation levels remains controversial because of incomplete C-to-T conversion, which might lead to overestimation (26, 27). Based on these considerations, we selected MeDIP-seq using a specific antibody against 5mC, to analyze mtDNA methylation dynamics and features during the critical developmental window.

Based on the technically reliable MeDIP-seq data (SI Appendix, Fig. S2A), we first profiled mtDNA methylation dynamics using reliable and reproducible reads mapping to the mitochondrial genome (SI Appendix, Fig. S2B and Dataset S1). We found that during the developmental transition from blastocysts to postimplantation embryos, the mtDNA methylation levels increased throughout the mitochondrial genome, albeit to varying degrees (Fig. 1D). In line with this, the heatmap showed more detailed information of mtDNA methylation dynamics: nearly all regions, regardless of their basal levels in blastocysts, displayed a notable increase in mtDNA methylation levels (Fig. 1E), and the increase covered not only the coding and noncoding regions, but also the control region (D-loop) of the mitochondrial genome (SI Appendix, Fig. S2C). It is also noteworthy that mtDNA methylation levels appeared to be inversely correlated with CpG density at each developmental stage (SI Appendix, Fig. S2 D and E), thus de novo mtDNA methylation tended to occur within regions with relatively low CpG density (Fig. 1F). The CpG-density-dependent mtDNA methylation pattern is consistent with results reported for the nuclear genome, particularly in postimplantation embryos and somatic cells (22, 28), and accords with the previous finding that mtDNA methylation is more likely to occur within non-CpG nucleotides (29-31). We next examined the fraction of regions that exhibited relatively high, intermediate, or low methylation scores in each developmental stage, and observed a clear increase in the fraction of regions that exhibited high methylation scores from blastocysts to postimplantation embryos, as well as a dramatic decrease in the fraction of regions that exhibited low methylation scores (Fig. 1*G*).

Finally, we validated the increase in mtDNA methylation from blastocysts to postimplantation embryos using MeDIP-qPCR, using a highly sensitive and specific anti-5mC antibody. During consecutive developmental stages of peri-implantation, all detected regions displayed an increase in the mtDNA methylation level to varying degrees from the hypomethylated state in blastocysts (Fig. 1*H*). This increase occurred irrespective of the type of mtDNA regions we detected: i.e., the protein coding regions (heavy-strand *Nd5*, light-strand *Nd6*), the noncoding region

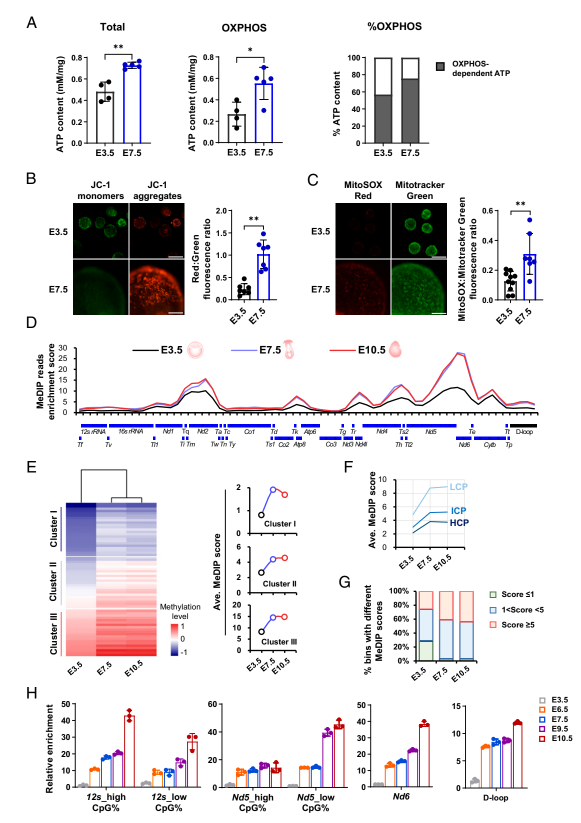


Fig. 1. Spatiotemporal coincidence of de novo mtDNA methylation and mitochondrial oxidative stress during the peri-implantation window. (*A*) Comparisons of total ATP levels (*Left*), as well as OXPHOS-dependent ATP production (*Middle*) and relative contributions (*Right*) between E3.5 and E7.5 embryos. (*B*) Mitochondrial membrane potential in E3.5 and E7.5 embryos. (*B*) Mitochondrial membrane potential in E3.5 and E7.5 embryos. (*B*) Mitochondrial membrane potential by quantifying ratios of red (JC-1 aggregates): green (JC-1 monomer) fluorescence. (Scale bar: 100 μ m.) (*C*) Fluorescent images of intramitochondrial ROS in E3.5 and E7.5 embryos detected by MitoSOX, a mitochondria-specific probe. *Right*: quantification of intramitochondrial ROS levels. (Scale bar: 100 μ m.) (*D*) Methylation profiles of linearized mtDNA detected using MeDIP-seq across the mouse mitochondrial genome in E3.5, E7.5, and E10.5 embryos. The *y* axis shows the normalized enrichment of MeDIP-reads in each bin of 500 bp. The annotations below the linearized mtDNA map show the positions of mtDNA elements. (*E*) Heatmap analysis of all bins of the mitochondrial genome based on their MeDIP enrichment scores at three developmental stages. *Right*: mtDNA methylation dynamics of three subcategories of bins with different average enrichment scores at E3.5. (*F*) mtDNA methylation dynamics of three subcategories of bins with different average enrichment scores at E3.5. (*F*) mtDNA methylation of enriched bins with different enrichment scores at three developmental stages. *(H*) Relative methylation profiles detected using MeDIP-qCR in selected regions during consecutive developmental stages. Data are presented as the mean \pm SD of three independent experiments. **P* < 0.05, ***P* < 0.01.

(12s), or the control region (D-loop). We also found that the increase in mtDNA methylation appeared to be asynchronous among different regions, as shown previously for the nuclear genome (22, 24). Collectively, these results demonstrated that the mitochondrial genome undergoes de novo methylation during the developmental transition from blastocysts to postimplantation embryos, which spatiotemporally coincides with enhanced mitochondrial OXPHOS and oxidative stress. Moreover, by reanalyzing the previously published high-throughput DNA methylation data of human early embryos (32) and mouse early embryonic and extraembryonic lineages (33), we found the occurrence of de novo mtDNA methylation, although limited, seems to be common among species and different lineages during the periimplantation window (*SI Appendix*, Fig. S3 A–D).

DNMT3A and DNMT3B Translocate into Mitochondria and Catalyze De Novo mtDNA Methylation during the Developmental Transition. We next attempted to explore the mechanism underlying de novo mtDNA methylation. Given that the mitochondrial genome does not encode enzymes with the de novo methyltransferase activity, we asked if nuclear-encoded DNMT3A and DNMT3B, the main enzymes catalyzing nuclear de novo methylation, are responsible for de novo mtDNA methylation. To this end, we first detected the mitochondrial localization of DNMT3A and DNMT3B via high-quality 3D-microscopic reconstruction using superresolution immunofluorescence analysis. The results indicated that both DNMT3A and DNMT3B were frequently localized to the mitochondrial matrix or membrane by implantation and postimplantation stages, but rarely by preimplantation stage (Fig. 2A and B). The mitochondrial localization of DNMT3A and DNMT3B was also supported by immunoelectron microscopy, in which they appeared as clustered or single colloidal gold particles (Fig. 2C). To further confirm the mitochondrial localization of DNMT3A and DNMT3B, we purified mitochondria. After excluding possible contaminants from the nuclear and cytosolic fractions, in which DNMT3A and DNMT3B preferentially reside, we detected the presence of these two de novo methyltransferases, albeit at relatively low levels, in the purified mitochondria (Fig. 2D). These results demonstrated that DNMT3A and DNMT3B can enter mitochondria during the process of de novo mtDNA methylation.

Physical binding of DNMT3A or DNMT3B to the hypomethylated DNA region is a prerequisite for catalyzing de novo methylation. Therefore, we next detected whether DNMT3A and DNMT3B could interact directly with mtDNA after entering the mitochondrial matrix. Using high-sensitivity immunoprecipitation, we detected the binding of DNMT3A and DNMT3B peaked by postimplantation stage (Fig. 2E and F), which accords with the relatively hypermethylated mtDNA observed in postimplantation embryos (Fig. 1H). From this stage onwards, most of the binding tended to decline (Fig. 2E and F). Collectively, we proved, using multiple methodologies, that DNMT3A and DNMT3B can translocate into mitochondria and bind to mtDNA during the process of de novo mtDNA methylation. Although DNMT3A and DNMT3B were previously reported to be absent in mitochondria from mouse embryonic fibroblasts and HCT116 human colon carcinoma cells (16), follow-up studies provided evidence that DNMT3A and DNMT3B were localized in the mitochondria or detectable in the mitochondrial protein fraction in human and mouse central nervous tissues, skeletal muscle, as well as in HeLa and 3T3-L1 cells (29, 34, 35). Our results, together with these findings, suggest that DNMT3A and DNMT3B

translocate into mitochondria in a cell type- and developmental stage-dependent manner.

To finally determine the role of DNMT3A or DNMT3B in catalyzing de novo mtDNA methylation, we measured mtDNA methylation levels in $Dnmt3a^{-/-}$ and $Dnmt3b^{-/-}$ E10.5 embryos, in which relatively stable mtDNA methylation patterns should been established. Deficiency of either Dnmt3a or Dnmt3b had no effect on mitochondrial mass (SI Appendix, Fig. S4 A and B); however, it did result in severe mtDNA hypomethylation, irrespective of the type of region detected (Fig. 2G), suggesting that DNMT3A and DNMT3B might catalyze de novo mtDNA methylation in a synergistic manner. These results, together with mitochondrial location of DNMT3A and DNMT3B, suggest that de novo mtDNA methylation depends on both DNMT3A and DNMT3B. This may be explained by the mitochondrial genome-wide distribution of DNMT3A or DNMT3B-preferred sites (SI Appendix, Fig. S4C). In agreement with this, previous studies have demonstrated that not only Dnmt3a knockout alone, but also Dnmt3al Dnmt3b double knockout and Dnmt1/Dnmt3al Dnmt3b triple knockout mouse embryonic stem cells showed a significant reduction in regional or global mtDNA methylation levels (14, 29). Similarly, a recent study clarified that knockdown of each DNMT, especially DNMT3B, in human breast epithelial cells markedly reduced mtDNA methylation levels (31), suggesting DNMT3A and DNMT3B might display functional synergism in catalyzing de novo mtDNA methylation. This notion is somewhat different to that occurring in the nuclear genome. It has been well-established that DNMT3B is the main enzyme responsible for nuclear de novo methylation around the implantation period, and DNMT3A cooperates with DNMT3B to methylate its targets (23, 24). More recently, however, a study revealed that a unique proportion of nuclear genomic regions showed both DNMT3A and DNMT3B-dependent DNA methylation (36). In line with this, earlier studies, focusing on specific sites, also showed the synergistic function of DNMT3A and DNMT3B in methylating the promoters of Oct4, Nanog, and hematopoietic stem cell-specific genes (37, 38), similar to our observations in the mitochondrial genome.

DNMT3A and DNMT3B Contain a Functional MTS that Promotes Efficient Mitochondrial Translocation. We next asked how DNMT3A and DNMT3B enter mitochondria. Nuclear-encoded mitochondrial proteins usually possess an MTS in their N terminus. This MTS forms a cleavable amphiphilic helical structure that facilitates the import of precursor proteins into the mitochondrial matrix (39). Although the mitochondrial location of DNMT3A and DNMT3B has been reported (29, 34, 35), the mechanism responsible for the location has never been identified (14). By examining the 5'-flanking regions of the published coding sequence of mouse Dnmt3a and Dnmt3b, we found an in-frame potential MTS coding region equivalent to 50 codons, starting with the ATG, which might encode a peptide that forms an amphiphilic helix, in the 5'-flanking region of Dnmt3b (Fig. 3A and SI Appendix, Fig. S4A), but not Dnmt3a. Of note, compared with exclusive mitochondrial proteins, mitochondrial proteins that can be located in different subcellular compartments, especially those preferentially residing in the nucleus with a conditional distribution in the mitochondria, always harbor an unconventional MTS. These unconventional MTSs are usually difficult to predict using bioinformatic analysis because they are likely to have lower prediction scores and might be located in the C- terminus (40, 41). Thus, we next examined the 3'-flanking

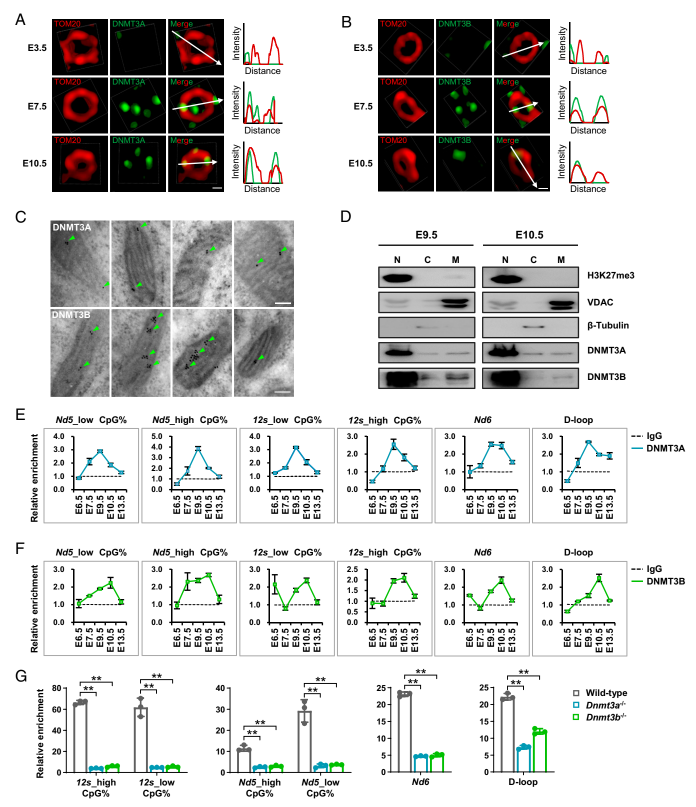


Fig. 2. DNMT3A and DNMT3B translocate into mitochondria and catalyze de novo mtDNA methylation during the developmental transition. (*A* and *B*) Highquality 3D-microscopic reconstruction of the interactions between TOM20-labeled mitochondria (ring-shaped cross section) and DNMT3A (*A*) or DNMT3B (*B*) in E3.5, E7.5, and E10.5 embryos using superresolution immunofluorescence analysis. (Scale bar: 250 nm.) *Right:* Fluorescence intensity profiles cross the white line in the merged picture. (*C*) Mitochondrial localization of DNMT3A and DNMT3B (revealed by clustered or single colloidal gold particles) detected using immunoelectron microscopy with gold-coupled antibodies in E7.5 embryos, (Scale bar: 100 nm.) (*D*) Western blotting analysis of DNMT3A and DNMT3B in purified mitochondria from E9.5 and E10.5 embryos (using earlier stage embryos was technically unfeasible because of the high number of embryos required). Nuclear (N), cytosolic (C), and mitochondrial (M) protein fractions were detected with antibodies against nuclear (H3K27me3)-, mitochondrial (voltage-dependent anion carrier, VDAC)-, and cytosolic (β-tubulin)-specific markers, respectively, to exclude the possible contamination of purified mitochondria with the nuclear or cytosolic fraction. (*E* and *P*) Chromatin immunoprecipitation analysis of the DNMT3A-mtDNA (*E*) and DNMT3B-mtDNA (*F*) physical interactions in selected regions during the peri-implantation stage. (G) Relative methylation levels detected using MeDIP-qPCR in selected regions in wild-type, *Dnmt3a^{-/-}*, and *Dnmt3b^{-/-}* E10.5 embryos. Data show the means \pm SD of three independent experiments. ***P* < 0.01.

regions, and detected three low-scoring potential MTSs in the 3'-flanking region of *Dnmt3a*. Among these, an MTS equivalent to 60 codons was predicted to have significant helical amphiphilicity (Fig. 3B and SI Appendix, Fig. S5A). In addition, the detection of transcripts containing the predicted MTS preferentially in postimplantation embryos (Fig. 3 C and D and SI Appendix, Fig. S5B), as well as the results that these transcripts were initially activated during the transition from blastocysts to epiblasts (SI Appendix, Fig. S5C), not only gives an explanation for the stage-dependent mitochondrial translocation of DNMT3A and DNMT3B isoforms, but also may explain why the mitochondrial genome gain DNA methylation during this developmental transition.

Finally, we functionally determined the efficacy of predicted MTS. When cloned in-frame into pEGFP-N1 with the C-terminal enhanced green fluorescent protein (EGFP), both potential *Dnmt3a* and *Dnmt3b* MTS, similar to the wellestablished CoxVIII MTS, could successfully direct EGFP to the mitochondria in either NIH/3T3 or HEK293T cell, as revealed by the colocalization of MitoTracker Red with EGFP (Fig. 3*E* and *SI Appendix*, Fig. S5*D*). Thus, these results indicated that both DNMT3A and DNMT3B have their own unconventional, but efficient, MTSs, which are responsible for the mitochondrial translocation of DNMT3A and DNMT3B during the process of de novo mtDNA methylation.

De Novo mtDNA Methylation Functions to Protect the Mitochondrial Genome against Oxidative Damage. Having demonstrated the essential role of DNMT3A and DNMT3B in catalyzing de novo mtDNA methylation and its underlying mechanism, we next investigated the developmental significance of de novo mtDNA methylation during early embryogenesis. Many previous studies have focused on the functional link between mtDNA methylation and gene expression or copy number; however, the results varied markedly or even contradicted each other among different studies. This is reminiscent of the important role of DNA methylation in maintaining genome stability (10, 11). Our data have shown that postimplantation embryos underwent an increase in mitochondrial aerobic metabolism, as well as intramitochondrial ROS accumulation arising from enhanced OXPHOS (Fig. 1 A-C and SI Appendix, Fig. S1 A and B). Given that mitochondria are both the major source and the sensitive target of ROS, mtDNA by this stage will suffer from significant oxidative stress that might induce mtDNA damage. This notion is also supported by the result of exogenous H₂O₂ treatment that mimics mitochondrial oxidative stress, which significantly induced mtDNA damage in postimplantation embryos (SI Appendix, Fig. S6 A and B). Thus, spatiotemporal coincidence of de novo mtDNA methylation and mitochondrial oxidative stress, led us to postulated that de novo mtDNA methylation might participate in protecting the mitochondrial genome against oxidative damage in postimplantation embryos. Quantification of mtDNA damage scores supported our hypothesis: compared with their wild-type counterparts, both $Dnmt3a^{-/-}$ and $Dnmt3b^{-/-}$ embryos showed significant increases in mtDNA damage throughout the mitochondrial genome (Fig. 4 A and B). Of note, increased mtDNA damage was not due to the effect of

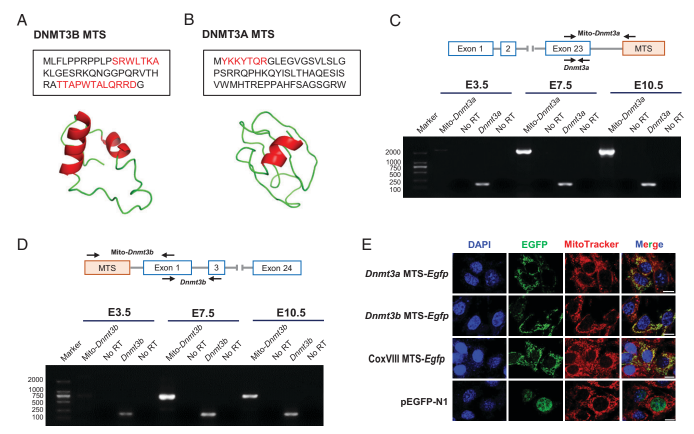


Fig. 3. DNMT3A and DNMT3B contain a functional MTS that promotes efficient mitochondrial translocation. (*A* and *B*) Ribbon models of the representative α -helical structure (red) and corresponding residues (red text) in the MTS of DNMT3B (*A*) and DNMT3A (*B*). The structure was modeled using the I-TASSER server (https://zhanglab.ccmb.med.umich.edu/I-TASSER/). (*C* and *D*) Agarose gel electrophoresis of RT-PCR products amplified from mature transcripts capable of encoding the MTS of DNMT3A (*C*) and DNMT3B (*D*) in E3.5, E7.5, and E10.5 embryos. No RT, no reverse transcriptase control. (*E*) Mitochondrial translocation of EGFP directed by the predicted MTS of DNMT3A and DNMT3B in transiently transfected NIH/3T3 cells. MitoTracker Red was used to label the mitochondria. The commercialized vector containing an MTS from the COX subunit VIII and pEGFP-N1 were used as the positive control and negative control, respectively. (Scale bar: 10 μ m.)

Dnmts deficiency per se. Possible changes that may lead to mtDNA damage were excluded: *i.e.*, OXPHOS was inhibited (*SI Appendix*, Fig. S6 *C–E*) in *Dnmt3b*-deficient postimplantation embryos. In addition, *Dnmt3b* deficiency did not induce mito-chondrial oxidative stress (*SI Appendix*, Fig. S6*F*).

Next, we attempted to further confirmed the role of elevated mtDNA methylation levels in protecting the mitochondrial genome against oxidative damage using an in vitro model. To this end, we established the model via mitochondria-targeted co-overexpression of Dnmt3a and Dnmt3b, as well as targeted overexpression of Dnmt3a or Dnmt3b alone in NIH/3T3 fibroblasts (SI Appendix, Fig. S7 A-C), which are sensitive to H₂O₂induced mitochondrial oxidative stress and mtDNA damage (SI Appendix, Fig. S7 D and E). Our results indicated that hypermethylated mtDNA showed significantly reduced mtDNA damage under H2O2-induced oxidative stress (Fig. 4 C and D). Note that co-overexpression of Dnmt3a and Dnmt3b was more effective in elevating mtDNA methylation levels and protecting against mtDNA damage, compared with overexpressing Dnmt3a or Dnmt3b alone (Fig. 4 C and D), which further supported the concept of functional synergism between DNMT3A or DNMT3B in catalyzing de novo mtDNA methylation.

Additionally, the protective role of de novo mtDNA methylation against oxidative damage, raised the possibility that nuclear de novo DNA methylation may also participate in the protection of nuclear genome. Our results from knockout embryos and in vitro model, indicated that elevated nuclear DNA methylation is also important for maintaining genome stability in the nucleus (SI Appendix, Fig. S8 A and B). In line with this, we noticed that glutathione antioxidant system and DNA repair pathway were notably activated during the periimplantation window (SI Appendix, Fig. S8 C and D), implying that embryos must face a comprehensive intracellular oxidative stress during the critical developmental transition, and a series of redundant mechanisms may be activated by this stage and participate in the protection of both the mitochondrial and nuclear genomes. Although the mechanism responsible for the contribution of cytosine methylation to genome stability remains controversial, possible explanations include increased DNA condensation and flexibility (42-44). In particular, mtDNA methylation may increase the binding affinity of mitochondrial transcription factor A (TFAM), and thus forming the TFAM-DNA complex, which in turn, alters the packaging and deformability of mtDNA (45, 46).

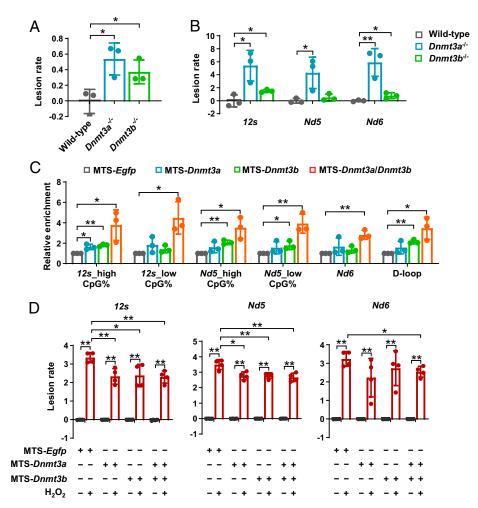


Fig. 4. De novo mtDNA methylation functions to protect the mitochondrial genome against oxidative damage. (*A* and *B*) Quantitative measurement of mtDNA damage within the long-range region (*A*) and selected regions (*B*) in wild-type, $Dnmt3a^{-/-}$, and $Dnmt3b^{-/-}$ postimplantation embryos. MtDNA damage is expressed relative to the wild-type (Zero-class lesions). (C) Effect of mitochondria-targeted co-overexpression of Dnmt3a and Dnmt3b, as well as over-expression of Dnmt3a or Dnmt3b alone, on the relative methylation levels in selected regions in NIH/3T3 cells. Mitochondria-targeted EGFP was transfected in parallel as the control. (*D*) Effect of mitochondria-targeted co-overexpression of Dnmt3a and Dnmt3b, as well as overexpression of Dnmt3a and Dnmt3b. The present expressed relative to the control (normalized to zero-class lesions). Data are presented as the mean \pm SD of three independent experiments. *P < 0.05, **P < 0.01.

Developmental transition from blastocysts to post-implantation embryos

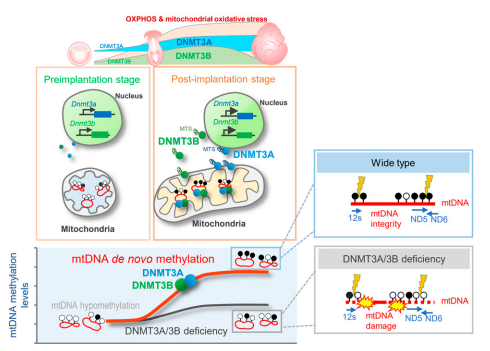


Fig. 5. A model illustrating the dynamics, mechanism, and functional consequence of de novo mtDNA methylation that occurs during the developmental transition from blastocysts to postimplantation embryos.

The presence of 5mC in mtDNA is a hallmark finding that advances the knowledge of mitochondrial biology and supports the novel concept of mitoepigenetics (16). Follow-up studies also confirmed mtDNA methylation among myriads of cell types (12, 16, 29, 47, 48), and evidence from clinical patients and mouse models further suggested that changes in mtDNA methylation are associated with pathogenesis of many diseases (31, 34, 49-52) and aging (53, 54). Despite these findings, the physiological role of mtDNA methylation, especially its developmental significance, remains unknown before the present study. Our results from the in vitro and in vivo models, suggested that de novo mtDNA methylation contributes to protecting the mitochondrial genome against oxidative damage. Our finding is in line with previous observations in the nuclear genome, in which global or regional hypomethylation are associated with an increased frequency of DNA damage of antioncogenes, and are implicated in carcinogenesis (55, 56).

In summary, our data provide direct evidence that the mitochondrial genome undergoes a wave of extensive de novo methylation during the developmental transition from blastocysts to postimplantation embryos. During this developmental window, DNMT3A and DNMT3B translocate into the mitochondrial matrix and bind mtDNA to catalyze de novo methylation of the hypomethylated mitochondrial genome. We also identified that DNMT3A and DNMT3B contain their own functional MTSs that promote efficient mitochondrial translocation. Our results also suggested that established mtDNA methylation in postimplantation embryos functions to protect the mitochondrial genome against oxidative damage (Fig. 5), provide the functional link between mitochondrial epigenetic remodeling and metabolic changes. Given that nearly all previous related studies only focused on mtDNA methylation in differentiated or adult somatic cells, our current findings, focusing on early embryos, answer the question regarding the origins of previously identified 5mC in mtDNA. Furthermore, focusing on

the essential role of nuclear encoded epigenetic enzymes on mitoepigenetics and mtDNA stability, the present study advances current knowledge of the crosstalk that couples the nuclear and mitochondrial genomes, and the role of this crosstalk in maintaining mitochondrial homeostasis.

Materials and Methods

Animal Studies and Ethical Approval. All female ICR mice used in this study were obtained from SPF Biotechnology Co. Ltd. Heterozygous $Dnmt3a^{+/-}$ and $Dnmt3b^{+/-}$ mice that carry one mutant locus of Dnmt3a and Dnmt3b, respectively (23), were obtained from the NIH mutant mouse research and resource center (MMRRC, NIH). All mice were maintained in a climate-controlled room on a 12-h light/dark cycle and allowed food and water ad libitum. The China Agricultural University Institutional Animal Care and Use Committee approved this study, which was performed in accordance with the committee's guidelines. All efforts were made to minimize animal suffering.

Preparation and Collection of Mouse Embryos. Embryos were obtained by natural breeding. ICR female mice (7-8 wk old) were cocaged individually with ICR males (10-12 wk old). The morning of the appearance of a vaginal plug was designated E0.5. Embryos were collected at selected stages. The criteria for sampling embryos were based on the developmental progress and morphology. Embryos showing typical morphological features according to the well-established landmarks were sampled for further analyses, as reported by the well-established guidance (57) and previous studies (21, 24, 58). At E3.5, well-developed late-cavitating blastocysts were obtained by flushing the uterus with M2 medium (Sigma-Aldrich). At E6.5 and E7.5, the conceptuses covered with the decidual mass were gently teased away from the uterus. The decidua in which the conceptus embedded was peeled off, and the parietal yolk sac was opened to expose the visceral yolk sacendoderm layer. The well-developed epiblast was separated from extraembryonic tissues using microdissecting watchmaker's forceps under a stereomicroscope. To ensure accurate sample collection and avoid cross-contamination between embryonic and extraembryonic tissues, pre-experiments have been performed using gRT-PCR and immunostaining for markers specific to the epiblast (59) before large-scale sampling. All sampled embryos were washed serially using phosphate-buffered saline (PBS;

Gibco) containing 0.1% polyvinyl alcohol (PBS-PVA) (Sigma-Aldrich) and stored immediately in liquid nitrogen for further use.

Statistical Analysis. Statistical tests were performed on SPSS 26.0 Windows version of software. Experiments were independently replicated for a minimum of three times unless otherwise specified, and all data are reported as the mean \pm SD. Student's *t* test (for two groups) was used to analyze the significant difference between the treatment and control groups. For multiple comparisons, data were analyzed by one-way ANOVA followed by Tukey's multiple comparison test to compare selected pairs of experimental groups. Statistically significant differences were defined as *P* < 0.05.

Data Availability. All study data are included in the article and/or supporting information.

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