Melatonin prevents deterioration in quality by preserving epigenetic modifications of porcine oocytes after prolonged culture

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ABSTRACT

Prolonged culture of metaphase II oocytes is an in vitro aging process that compromises oocyte quality. We tested whether melatonin preserves epigenetic modifications in oocytes after prolonged culture. The porcine oocytes were matured in vitro for 44 h, and then metaphase II oocytes were continuously cultured in medium supplemented with or without melatonin for 24 h. We found that the parthenogenetic blastocyst formation rate of prolonged-culture oocytes was lower than in fresh oocytes. We further observed that methylation at H3K4me2 and H3K27me2 of oocytes enhanced after prolonged culture. However, 5mc fluorescence intensity was lower in prolonged-culture oocytes than in fresh oocytes. Moreover, the promoter of the imprinted gene NNAT exhibited a higher level of DNA methylation in prolonged-culture oocytes than in fresh oocytes, which was associated with a reduced expression level and glucose uptake capability. Conversely, melatonin improved blastocyst formation rate and preserved histone and DNA methylation modifications, as well as NNAT function in the oocytes after prolonged culture. Notably, DNA methyltransferase inhibitor 5-aza significantly attenuated the protective role of melatonin on genomic DNA methylation. In summary, our results revealed that epigenetic modifications are disrupted in oocytes after prolonged culture, but the changes are reversed by melatonin.

INTRODUCTION

Factors within oocytes are essential for successful reproduction. However, the oocyte is vulnerable to the adverse endogenous and/or exogenous environment, leading to impaired quality and compromised development. For example, in postovulatory oocyte aging, a process where fertilization does not occur within the ideal window period of post ovulation, the unfertilized metaphase II (MII) oocyte will undergo a time-dependent deterioration in vivo or in vitro [1, 2], thereby leading to declined fertilization capability [3, 4] and compromised early embryo development [5]. Moreover, oocyte aging also has detrimental effects on fetus developmental during mid-term gestation [6] and fitness of the offspring [7, 8], demonstrating that postovulatory oocyte aging has profound impacts on offspring health. Therefore, it is important to fertilize the oocyte when it is fresh or to protect oocytes from deterioration when delayed fertilization is performed.

In view of the long-term effects associated with oocyte aging that originates from defects within oocytes, various attempts have been carried out to prevent oocyte deterioration during postovulatory aging, including the modification of the components of culture medium or supplementation with specific compounds. For instance, melatonin (N-acetyl-5-methoxytryptamine), a major
product secreted by the pineal gland, which participates in the entrainment of the circadian rhythms and seasonal reproduction in animals [9], has been successfully applied to protect oocytes from aging in vitro [3, 4, 10, 11]. Previous reports show that oxidative stress is induced during postovulatory aging [12], which leads to impaired functions in oocytes including disrupted spindle assembly, chromosome alignment, actin polymerization, and mitochondrial integrity [13, 14]. Therefore, fertilization capability and the competence of early embryo development are reduced [4, 10, 13]. Being a free radical scavenger and potent antioxidant [15], melatonin vastly reduces reactive oxygen species (ROS) level and attenuates the defects of postovulatory aging oocytes. However, the molecular process by which melatonin delays oocyte postovulatory aging still needs to be further elucidated.

Epigenetic modifications are mechanisms that regulate gene expression independently of the DNA sequence. Genomic DNA methylation and histone modification are widely-studied epigenetic mechanisms. DNA methylation occurs on the 5-position cytosine, which suppresses gene expression, while demethylation activates gene expression. DNA methylation patterns are established during oogenesis to maintain subsequent development and offspring fitness [16]. Methylation not only takes place at the genomic DNA level but also at the loci of the histone lysine. In contrast to DNA methylation, histone methylation can switch gene expression on or off depending on which lysine loci are methylated. In general, methylation at H3K4 is associated with gene activation, but methylation at H3K27 is associated with gene repression. Both the modification of DNA and histones by methylation play important roles in oogenesis and early embryogenesis [17, 18]. Defects of epigenetic modifications may contribute to compromised oocyte quality during postovulatory aging [19, 20]. We and others have observed defective epigenetic modifications in postovulatory aging mouse and porcine oocytes [5, 21-23], whether these epigenetic abnormalities can be reversed needs to be further explored [19].

In the present study, we determined whether melatonin preserves epigenetic modifications in oocytes during in vitro aging. We observed that histone methylation, global genomic DNA methylation, and DNA methylation of individual gene were disrupted in porcine MII oocytes after prolonged culture. Moreover, gene expression and gene function was further altered and associated with hypermethylation level at the NNAT promoter. Conversely, melatonin could fully or partially reverse the altered epigenetic modification parameters in prolonged-culture oocytes. We report that melatonin could preserve epigenetic in oocytes after prolonged culture.

RESULTS

Melatonin attenuates the deterioration in the quality of prolonged-culture oocytes

A high-quality oocyte is essential for early development, therefore we tried to determine the protective role of melatonin on the quality of prolonged-culture oocytes via evaluating the blastocyst formation rate. To do this, fresh, prolonged culture (prolonged-culture) and prolonged culture + Mel (Melatonin, $10^{-3}$ or $10^{-5}$ M) oocytes were parthenogenetically activated and then cultured in vitro to check blastocyst formation. As shown in Figure 1A and B, the blastocyst formation rate of the prolonged-culture oocytes was lower than the fresh oocytes (Fresh, 37.0 ± 2.6%; prolonged culture, 22.5 ± 3.2%; $P<0.01$), indicating oocyte quality is impaired after prolonged culture. Conversely, melatonin supplement at $10^{-5}$ M and $10^{-3}$ M concentration increased blastocyst formation rate to 31.0 ± 2.9% and 37.4 ± 3.6%, respectively. Collectively, our data showed that melatonin enhances blastocyst formation of prolonged-culture oocytes, suggesting that melatonin prevents deterioration. A melatonin concentration of $10^{-3}$ M significantly improved blastocyst formation rate of the prolonged-culture oocytes (prolonged culture, 22.5 ± 3.2%; prolonged culture + Mel, 37.4 ± 3.6%; $P<0.05$), therefore this concentration was used in subsequent experiments.

Melatonin preserves the histone methylation of prolonged-culture oocytes

To test whether histone methylation in oocytes is influenced by prolonged culture and melatonin, oocytes were stained with antibody H3K4me2 and H3K27me2. As shown in Figure 2A and B, we found that the fluorescence intensity of H3K4me2 was higher in the prolonged-culture oocytes compared to the fresh oocytes (Fresh, 11.9 ± 0.9; prolonged culture 19.4 ± 1.1; $P<0.0001$), and the melatonin supplement reduced the fluorescence intensity of H3K4me2 in the prolonged-culture oocytes (prolonged culture, 19.4 ± 1.1; prolonged culture + Mel, 13.3 ± 0.9; $P<0.0001$). Similarly, as shown in Figure 3A and B, fluorescence intensity of H3K27me2 was higher in the prolonged-culture oocytes compared to the fresh oocytes (Fresh, 6.1 ± 0.3; prolonged culture, 26.9 ± 2.0, $P<0.0001$), but melatonin reduced the fluorescence intensity of H3K27me2 in the prolonged-culture oocytes (prolonged culture, 26.9 ± 2.0; prolonged culture + Mel, 12.4 ± 0.6; $P<0.0001$). Taken together, the above results suggest that histone methylation of the oocytes is disrupted by prolonged culture, however, melatonin protects against the defect.
Figure 1. Melatonin enhances the blastocyst formation rate of prolonged-culture oocytes. Porcine oocytes matured in vitro were continuously cultured in medium supplemented with or without melatonin (10⁻³ or 10⁻⁵ M) for 24 h. Oocytes were pathogenetically activated and then cultured for 7 days to examine the blastocyst formation rate. Views of blastocysts (A) and blastocyst formation rate (B) of fresh, prolonged-culture, and prolonged-culture + Mel oocytes. The data are presented as the mean ± SEM of at least three independent experiments. Bar=200 μm; PC, prolonged-culture; Mel, melatonin; *P<0.05, **P<0.01.

Figure 2. The effect of melatonin on histone methylation at H3K4me2 of prolonged-culture oocytes. Porcine oocytes matured in vitro were continuously cultured in medium supplemented with or without 10⁻³ M melatonin for 24 h. (A) Representative images of fresh, prolonged-culture and prolonged-culture + Mel oocytes stained with H3K4me2 antibody. (B) Quantitative analysis of H3K4me2 fluorescence intensity. The data are presented as mean ± SEM of at least three independent experiments. PC, prolonged-culture; Mel, melatonin; Scale bar=100 μm. Data are expressed as the mean ± SEM, ****P<0.0001.
Melatonin attenuates global genomic DNA methylation defects in prolonged-culture oocytes

DNA methylation is the most-studied epigenetic modification, which regulates gene expression independently of the gene sequence. We next tested whether DNA methylation in oocytes is influenced by prolonged culture and exposure to melatonin. The fresh, prolonged culture, and prolonged culture + Mel oocytes were stained with antibody 5mc. As shown in Figure 4A and B, the fluorescence intensity of 5mc in the prolonged-culture oocytes was significantly reduced compared to

Figure 3. Effect of melatonin on histone methylation at H3K27me2 of prolonged-culture oocytes. Porcine oocytes matured in vitro were continuously cultured in medium supplemented with or without $10^{-3} \text{M}$ melatonin for 24 h. (A) Representative images of fresh, prolonged-culture and prolonged-culture + Mel oocytes stained with H3K27me2 antibody. (B) Quantitative analysis of H3K27me2 fluorescence intensity. PC, prolonged-culture; Mel, melatonin. Scale bar=100 μm. The data are expressed as the mean ± SEM of at least three independent experiments, **** $P<0.0001$.

Figure 4. Effects of melatonin on global genomic DNA methylation and DNMT expression in prolonged-culture oocytes. Porcine oocytes matured in vitro were continuously cultured in medium supplemented with or without $10^{-3} \text{M}$ melatonin for 24 h. (A) Representative images of fresh, prolonged-culture, and prolonged-culture + Mel oocytes stained with 5mc antibody. (B) Quantitative analysis of 5mc fluorescence intensity. (C) Expression of $DNMT1$, $DNMT3A$, and $DNMT3B$ in fresh, PC, and PC + Mel oocytes. The data are presented as the mean ± SEM of at least three independent experiments. Scale bar=100 μm; PC, prolonged-culture; Mel, melatonin; **$P<0.01$, ***$P<0.001$, ****$P<0.0001$. 
the fresh oocytes (Fresh, 18.4 ± 0.6; prolonged culture, 10.5 ± 0.4; \( P<0.0001 \)). Interestingly, melatonin significantly increased the fluorescence intensity of 5mc in the prolonged-culture oocytes (prolonged-culture, 10.5 ± 0.4; prolonged-culture + Mel, 16.9 ± 0.7; \( P<0.0001 \)). Because DNA methylation is mediated by DNA methyltransferase (DNMTs), we next investigated DNMT expression. As shown in Figure 4C, the expression level of DNMT1, DNMT3A, and DNMT3B in the prolonged-culture oocytes were significantly reduced compared to the fresh oocytes (1.0 ± 0.1, 0.2 ± 0.1, \( P<0.01 \); 2.1 ± 0.1, 1.2 ± 0.1, \( P<0.001 \); and 3.9 ± 0.5, 2.3 ± 0.1, \( P<0.01 \); for DNMT1, DNMT3A, and DNMT3B, respectively). Notably, melatonin was able to significantly increase the expression level of DNMT1 compared to the prolonged-culture oocytes (prolonged culture, 0.2 ± 0.1; prolonged culture + Mel, 0.8 ± 0.2; \( P<0.01 \)), and DNMT3A and DNMT3B expression was also slightly increased (prolonged culture, 1.2 ± 0.1, 2.3 ± 0.1, prolonged culture + Mel, 1.4 ± 0.2, 2.8 ± 0.3 for DNMT3A and DNMT3B, respectively), suggesting it is possible that melatonin affects DNA methylation by regulating DNMT expression.

**Melatonin preserves DNA methylation, expression and function of NNAT in prolonged-culture oocytes**

A previous study has shown that promoter DNA methylation and the function of NNAT are disrupted in prolonged-culture porcine oocytes [5], therefore, we used NNAT as an indicator to examine whether melatonin could reverse these defects (Figure 5A). To examine DNA methylation level in fresh, prolonged culture, and prolonged culture + Mel oocytes we applied bisulfite sequencing. As shown in Figure 5B-D, the DNA methylation level of the NNAT promoter was 93% in fresh oocytes but reached 100% in the prolonged culture oocytes. Notably, DNA methylation at the NNAT promoter was reduced to 95.5% in the prolonged culture + Mel oocytes. Our results show that melatonin could attenuate a DNA methylation defect at a specific promoter.

Melatonin could preserve the DNA methylation of the NNAT promoter in prolonged culture oocytes, therefore, we further studied how melatonin impacts NNAT expression and its function. We first determined the NNAT expression level in fresh, prolonged culture, and the prolonged culture + Mel oocytes. As shown in Figure 6A, the mRNA level of NNAT in the prolonged culture oocytes was significantly lower than the fresh oocytes (Fresh, 0.7 ± 0.1; prolonged culture, 0.3 ± 0.1; \( P<0.05 \)), and slightly increased after treatment with melatonin (prolonged culture, 0.3 ± 0.1; prolonged culture + Mel, 0.6 ± 0.2; \( P=0.099 \)). We further stained the oocytes with NNAT antibody to examine protein level. Similar to mRNA expression, fluorescence intensity of NNAT was reduced in the prolonged culture oocytes (Fresh, 56.9 ± 3.6; prolonged culture, 37.6 ± 1.6; \( P<0.0001 \)) and increased in the prolonged culture +

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**Figure 5. The effect of melatonin on DNA methylation on the NNAT promoter of prolonged-culture oocytes.** Porcine oocytes matured in vitro were continuously cultured in medium supplemented with or without \( 10^{-3} \) M melatonin for 24 h. (A) Primer and CpG loci examined at the NNAT differentially methylated region (DMR) are shown. DNA methylation was examined by bisulfite sequencing in fresh (B), prolonged-culture (C) and prolonged-culture + Mel oocytes (D). Each line presents an individually cloned allele, with an open circle for a non-methylated CpG site and filled circles for methylated CpG. Ten clones successfully sequenced for each group are presented. Methylation level (%) = (Methylated CpG/Total CpG) \times 100. PC, prolonged-culture; Mel, melatonin.
Mel oocyte (prolonged culture, 37.6 ± 1.6; prolonged culture + Mel, 47.1 ± 2.1; P<0.05), further confirming that melatonin improved NNAT expression (Figure 6B and C). As previous studies revealed that down-regulation of NNAT inhibited glucose uptake in prolonged-culture porcine oocytes, we next examined the oocytes capability to uptake glucose after treatment with melatonin. To accomplish this, fresh, prolonged culture, and prolonged culture + Mel oocytes were incubated with 2-NBDG. As shown in Figure 6D and E, the glucose uptake level in the prolonged culture oocytes is lower than that of the fresh oocytes (Fresh, 42.2 ± 1.9; prolonged culture, 31.3 ± 1.0; P<0.0001), but melatonin significantly enhanced glucose uptake level (prolonged culture, 31.3 ± 1.0; prolonged culture+ Mel, 43.2 ± 1.3; P<0.0001). Collectively, our data indicate that melatonin preserves DNA methylation and gene function of prolonged-cultured oocytes.

**DNMT inhibitor 5-aza attenuates the protective role of melatonin on genomic 5mc in prolonged-culture oocytes**

Given that DNA methylation is mediated by DNMTs, we tried to explore whether melatonin affects DNA methylation via activating DNMT activity in prolonged-culture oocytes. To achieve this, DNMT inhibitor 5-aza was added to the medium during prolonged culture. If DNMT activity is inhibited, the protective role of melatonin on genomic DNA methylation will be attenuated. Expectedly, we first observed that the prolonged culture + Mel oocytes have higher DNMT1 protein level compared to that of the prolonged culture oocytes (prolonged culture + Mel, 27.8 ± 0.8; prolonged culture, 21.2 ± 0.8; P<0.0001), as shown in Figure 7A and B. Interestingly, 5-aza attenuated the upregulation of DNMT1 protein by melatonin (prolonged culture + Mel + 5-aza, 23.9 ± 0.9; prolonged culture + Mel, 27.8 ± 0.8; P<0.01). We next assayed whether 5-aza may weaken the action of melatonin to preserve genomic 5mc in prolonged-culture oocytes. We stained the prolonged culture, prolonged culture + Mel, and prolonged culture + Mel + 5-aza oocytes with 5mc antibody. Consistent with our above findings, prolonged culture + Mel oocytes have higher 5mc fluorescence intensity than the prolonged culture oocytes (prolonged culture + Mel, 35.5 ± 1.2; prolonged culture, 26.3 ± 1.4; P<0.0001), as shown in Figure 7C and D, but 5-aza significantly reduced the upregulation of 5mc fluorescence intensity induced by melatonin (prolonged culture + Mel + 5-aza, 23.9 ± 0.9; prolonged culture + Mel, 27.8 ± 0.8; P<0.01).

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**Figure 6. The effect of melatonin on NNAT expression and the glucose uptake of prolonged-culture oocytes.** Porcine oocytes matured in vitro were continuously cultured in medium supplemented with or without 10^{-3} M melatonin for 24 h. (A) NNAT mRNA levels in the fresh, prolonged-culture, and prolonged-culture + Mel oocytes. (B) Representative images of fresh, prolonged-culture and prolonged-culture + Mel oocytes stained with NNAT antibody. (C) Quantification of the NNAT protein level. (D) Representative images of fresh, prolonged-culture and prolonged-culture + Mel oocytes stained with 2-NBDG. (E) Quantification of the 2-NBDG level. PC, prolonged-culture; Mel, melatonin; Scale bar=100 μm. The data are presented as the mean ± SEM of at least three independent experiments. *P<0.05, **** P<0.0001.
culture + Mel + 5-aza, 31.8 ± 1.0; prolonged culture + Mel, 35.5 ± 1.2; P<0.05). Taken together, our results suggest that melatonin preserves genomic DNA methylation through the regulation of DNMT1 expression.

**DISCUSSION**

In the current study, we observed epigenetic defects, including disruption of histone methylation and DNA methylation, at overall and local levels in porcine oocytes subjected to prolonged culture. Along with previous findings, we demonstrate that the epigenetic abnormalities may contribute to the compromised quality of prolonged-culture oocytes [5, 21, 23, 24]. Though some compounds, such as melatonin have been found to effectively delay postovulatory oocyte aging and extend the window for optimal fertilization [21, 25, 26], the mechanisms behind this still need to be further explored. We revealed that melatonin preserved histone methylation and DNA methylation of prolonged-culture porcine oocytes. Notably, we observed that melatonin stabilizes the modification of DNA methylation via regulating DNMT1 activity. Our findings will help us to further understand the epigenetic defects in prolonged-culture oocytes, and whether the defects are reversible. Thus, it is valuable to preserve epigenetic modifications when delayed fertilization oocytes are applied in the in vitro production of animal embryos and in assisted reproductive technologies in humans [2].

Histone methylation regulates gene expression and plays a crucial role in regulating oocyte growth [27, 28]. In general, H3K4 methylation is related to gene activation while H3K27 methylation is associated with gene inactivation [29]. Therefore, we selected H3K4me2 and H3K27me2 as indicators to examine the impact of melatonin on histone methylation in prolonged-culture oocytes. Our results showed that both H3K4me2 and H3K27me2 gained methylation in prolonged-culture oocytes. Along with a recent study, which reported that H3K9me3 was disrupted in postovulatory aging mouse oocytes [21], our evidence demonstrates that a defect in histone methylation may be one of the molecular mechanisms contributing to the compromised quality of prolonged-culture oocyte. Exposure to toxic substances also alters histone methylation at H3K4me2 and H3K27me3 [30, 31]. The evidence implies that an abnormality in histone methylation may contribute to poor oocyte quality. Therefore, it is necessary to maintain histone methylation to prevent the deterioration of oocytes. Melatonin is widely applied in a protective role against aging in oocytes but the relative molecular processes need further study. We currently investigated the effect of melatonin on histone methylation in prolonged-culture oocytes and observed that melatonin reduced histone methylation at H3K4me2 and H3K27me2. Melatonin also could reverse the defect of histone methylation at H3K9me2 in mouse oocytes exposed to deoxynivalenol during IVM [32]. Although the underlying mechanisms...
still need to be further explored, it is confirmed that melatonin has a protective role on histone methylation in oocytes after prolonged culture.

DNA methylation patterns of MII oocytes are established during oogenesis and oocyte growth [33, 34]. Soon after fertilization, paternal pronuclei demethylates are followed by maternal pronuclei, but an imprinted gene maintains its DNA methylation pattern during this wave of demethylation [16]. If the DNA methylation patterns in oocytes or zygotes are disrupted, the subsequent development may be adversely impacted [35]. We observed that overall genomic DNA had a reduced level of methylation and that NNAT exhibited levels of hypermethylation at differentially methylated regions (DMRs) in prolonged-culture oocytes. We further found that hypermethylation at NNAT DMR inhibited glucose uptake. These findings show that not only DNA methylation but also gene function is disrupted in prolonged-culture oocytes. When cultures were supplemented with melatonin, DNA methylation at global and local levels was restored. Because melatonin may regulate gene expression by mediating epigenetic modification, we also found that NNAT expression is rescued in prolonged-culture oocytes [36]. Collectively, we revealed that epigenetic defects with histone and DNA methylation modifications are reversed by melatonin in prolonged-culture oocytes. However, the possible links between histone and DNA methylation and how they adversely affect oocyte quality still need to be further explored.

DNA methylation is one of the best studied epigenetic modifications [16]. We further tried to explore how melatonin restores genomic DNA methylation in prolonged-culture oocytes. The DNA methylation pattern is regulated by DNA methyltransferase, therefore, the protective role of melatonin on the modification of DNA methylation in the prolonged-culture oocyte could be mediated by DNMTs. We found that the DNMT1 protein was downregulated in the prolonged-culture oocytes, but it was reversed by melatonin. This is concomitant with the trend of global genomic DNA 5mc among fresh, prolonged culture, and prolonged culture + Mel oocytes. DNMT1 activity is majorly inhibited by 5-aza, leading to a hypomethylation status [37]. When DNMT1 activities were inhibited by 5-aza, the protective role of melatonin on genomic DNA methylation was attenuated. Moreover, a recent study demonstrates that melatonin could greatly enhance DNMT1 expression to promote porcine oocyte maturation in vitro [38], suggesting that the beneficial effects of melatonin on oocytes may be mediated by DNMTs.

Of note, we observed that global genomic DNA lost methylation but some specific loci gained methylation in prolonged-culture oocytes. It appears that both are inconsistent. This could be because only one gene was examined, we could not rule out whether some other genes also lost DNA methylation as in the global genomic DNA. This is supported by our previous observation that the imprinted gene Snrpn lost DNA methylation in postovulatory aging mouse oocytes [23]. Another possible interpretation is that the loss of genomic DNA methylation leads to overall hypomethylation and unstable genomic DNA. Thereby, some specific loci may gain or lose methylation. Interestingly, this phenomenon is observed in tumor tissue, global genomic DNA exhibits hypomethylation status while suppressed cancer genes are hypermethylated to promote tumor growth [39, 40]. In particular, melatonin treatment inhibits breast cancer tumor growth in mice through upregulating global DNA methylation [41].

The limitation of our study is that only histone and DNA methylation were investigated. There are still some other epigenetic modifications such as histone acetylation and non-coding RNA that need to be further explored in prolonged-culture and melatonin-exposed oocytes. However, our results at least provide indications that melatonin prevents oocytes from deterioration during postovulatory aging partially by preserving epigenetic modifications.

MATERIALS AND METHODS

All chemicals and reagents used in the present study were purchased from Sigma-Aldrich (St. Louis, MO, USA) except where otherwise stated.

In vitro maturation (IVM)

Porcine ovaries were obtained from a local slaughterhouse and transported to the laboratory in 0.9% saline at approximately 30ºC, as previously described [42]. Follicular fluid from antral follicles at 3–8 mm diameter was aspirated using an 18-gauge needle connected to a 10 ml disposal syringe. Cumulus-oocyte complexes (COCs) were collected and washed multiple times with polyvinyl alcohol (PVA)-TL-HEPES. COCs were matured in vitro in TCM-199 (Gibco) culture medium supplemented with 3.05 mM D-glucose, 0.91 mM sodium pyruvate, 0.57 mM cysteine, 0.01 μg/ml epidermal growth factor (EGF), 0.50 μg/ml follicle-stimulating hormone (FSH), 0.50 μg/ml luteinizing hormone (LH), 10% (v/v) porcine follicular fluid, 75.00 μg/ml penicillin G, and 50.00 μg/ml streptomycin. Approximately, fifty COCs were cultured per 200 μl IVM medium covered with mineral oil. COCs were cultured in an incubator with humidified air at 38.5ºC and 5% CO₂ for 44 h.
Prolonged culture and treatment

At 44 h post IVM, cumulus cells were removed from COCs in PVA-TL-HEPES containing 0.1% hyaluronidase. The oocytes with intact morphology and first polar body were selected and continuously cultured in PZM-3 medium supplemented with or without melatonin for 24 h. Melatonin was dissolved in absolute ethanol and the final concentration of ethanol in the aging medium did not exceed 0.1%. Based on previous studies and our preliminary test, \(10^{-5}\) M and \(10^{-3}\) M melatonin were added to the PZM-3 medium.

The DNMT inhibitor 5-aza was purchased from Aladdin (Shanghai, China). The 5-aza was dissolved in DMSO and the final concentration in the medium was \(1\) µM [37].

Parthenogenetic activation

The parthenogenetic activation was performed using electrical pulses (2 DC pulses of 1.2 kV/cm for 30 µs) under an ECM 2001 electro cell manipulator (BTX Inc., San Diego, CA, USA) in the activation medium (0.30 M Mannitol, 1.00 mM CaCl\(_2\)-2H\(_2\)O, 0.10 mM MgSO\(_4\), 0.50 mM HEPES plus 0.3% (w/v) BSA). Activated oocytes were immediately transferred to PZM-3 and cultured in a humidified air incubator at 38.5°C and 5% CO\(_2\). Blastocyst formation was examined at 168 h after activation.

Glucose uptake assay

A fluorescent glucose analog, 2-(N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl) amino)-2-deoxyglucose (2-NBDG, Beijing Chemsynlab Pharmaceutical Co. Ltd., Beijing, China; CAS 186689-07-6), was used to evaluate glucose uptake capability. Oocytes were cultured in 200 µM 2-NBDG dissolved in PBS for 30 min at 37°C. After washing with PBS, images were captured under a fluorescence microscope (Nikon, Japan). Fluorescence intensity was quantified by using NIH Image J software to compare glucose uptake capability.

RNA isolation and real-time PCR

Total RNA was extracted from 50 oocytes by using an RNaseq Pure Micro kit (Tiangen, Beijing, China) according to the manufacturer’s instructions. cDNA was synthesized by using a Fast Quant RT Kit (Tiangen). The real-time PCR reaction system was prepared as follows: 10 µl Super Real Premix Plus (2×) (Tiangen), 2 µl cDNA, and 0.2 µM primers. The PCRs were performed with a Bio-Rad CFX 96 (USA) unit using the following program: 1 cycle at 95°C for 15 min, 39 cycles at 95°C for 10 s, 60°C for 30 s. As previously described, eGFP was added to the oocyte sample before RNA isolation. The expression of eGFP was used for normalization and relative expression levels were determined by the \(2^{-\Delta\Delta C_T}\) method [5]. The primers used in this study are listed in Table 1.

Immunofluorescent staining

Oocytes were collected and washed with PBS containing 0.1% PVA then fixed with 4% paraformaldehyde (PFA) at room temperature (RT) for 30 min. The oocytes were then permeabilized with 1% Triton-100 in PBS at RT for 8–12 h.

| Gene     | Primer sequence 5’-3’ | Gene Access no./Reference | Length (bp) |
|----------|----------------------|---------------------------|-------------|
| Dnmt1    | F: GTGCGTTTGTGAGTTTGT R: CATCATCGTCTCCTCTTC | XM_021082064.1 | 150         |
| Dnmt3a   | F: GAAATGCCACCAATCAGCC R: GAACTTGCCGTCTCCGAAC | XM_021085534.1 | 193         |
| Dnmt3b   | F: ATTTGACGGGTACCGGAGAC R: TTCGGAACCGGTGGACTTT | XM_021077256.1 | 114         |
| Tet1     | F: TCTTCCGACCTTGTCTACC R: GCTCGTCTTCTCCACCA | NM_001315772.1 | 158         |
| NNAT     | F: CGACAATACACAGATCTCTT R: CTTGGTTCCAGAATGTTTG | Gao et. al. [5] | 138         |
| BSP-NNAT-out | F: ATAGTAGTTTGTGAGAGAG R: ATAATGCACAGATCTCTTCTT | Gao et. al. [5] | 748         |
| BSP-NNAT-in | F: ATAGTAGTTTGTGAGAGAG R: ATAATGCACAGATCTCTTCTT | Gao et. al. [5] | 224         |
For 5mc staining, oocytes were blocked with 1% BSA for 1 h and treated with 2 N HCl for 15 min. They were then neutralized with 100 mM Tris-HCl (PH=8.5) for 20 min at RT. After incubation in 0.05% Tween-20 for 1 h at RT, oocytes were incubated overnight at 4°C with 5mc antibody (Zymo Research, Irvine, CA, USA) at a dilution of 1:500. After washing with 0.05% Tween-20, oocytes were incubated with anti-mouse IgG/TRITC second antibody (ZSGB-BIO, Beijing, China) at a dilution of 1:300 for 1 h at RT.

For histone methylation and Dnmt1 staining, oocytes were blocked with 1% BSA for 1 h at RT. After washing with 0.05% Tween-20, oocytes were incubated with primary antibody H3K4me2 and H3K27me2 (Cell Signaling, Danvers, MA, USA) at a dilution of 1:500, and DNMT1 (Proteitech, Chicago, IL, USA) at a dilution of 1:200 overnight at 4°C. Then the oocytes were washed with 0.05% Tween-20 followed by incubation with secondary antibody Alexa Fluor 488 (ZSGB-BIO, Beijing, China) for 1 h at RT.

NNAT staining was performed according to a previous description [5]. Briefly, oocytes were washed three times with PBS, fixed with 4% PFA at RT for 30 min and then placed into 50% methanol for 5 min, 100% methanol for 5 min, and 100% acetone for 5 min to remove lipid droplets and then were immediately permeabilized in 1% Triton-100 overnight at RT. After blocking in 1% BSA for 1 h at RT, the oocytes were incubated with NNAT antibody (Bioss, Beijing, China) at a dilution 1:300 overnight at 4°C. Oocytes were washed three times with 0.05% Tween-20, stained with goat anti-rabbit Alexa Fluor 488 (ZSGB-BIO, Beijing, China) at a dilution of 1:200. Then they were washed with 0.05% Tween-20 and stained with 10 mg/ml hoechst33342 for 30 min at 37 °C. Oocytes were washed in PBS with 0.1% PVA and mounted on glass slides. Images were captured by a confocal laser scanning microscope (LAS - Leica TCS-SP8, Wetzlar, Germany). To compare the fluorescence intensity of the same antibody, the same scan settings were used in fresh, prolonged-culture, and prolonged-culture oocytes with melatonin (prolonged-culture + Mel).

Bisulfite sequencing (BS)

Oocyte genomic DNA was modified using the EZ DNA Methylation-Direct Kit (Zymo Research) according to the manufacturer’s instructions with minor modifications as previously described [43]. To obtain PCR products, nested PCR was carried out using the following reaction system: the first round comprised 2 µl bisulfite-converted DNA, 0.5 mM primers and 12.5 µl Zymo Taq™ Premix (Zymo Research) in a total volume of 25 µl. The second round of PCR was performed with 2 µl of the first-round product as template, 0.5 mM primers, and 10 µl Zymo Taq™ Premix in a total volume of 20 µl. The program for the first round was 1 cycle at 95°C for 10 min, 39 cycles of 95°C for 30 s, 53°C for 30 s, 72°C for 1 min; and 1 cycle at 72°C for 7 min. The second round PCR was performed as 1 cycle at 95°C for 10 min, 39 cycles of 95°C for 30 s, 56°C for 30 s, 72°C for 1 min; and 1 cycle at 72°C for 7 min. Products of the second round PCR were then recovered and gel-purified using the Universal DNA Purification Kit (Tiangen). Purified fragments were sub-cloned into a PMD18T-vector (Takara, Japan). Clones confirmed by PCR were selected for DNA sequencing. Snap Gene 2.3.2 software was used to blast the sequence.

Statistical analysis

Data were presented as mean ± SEM. Statistical analyses were made using Prism 5 software (GraphPad, San Diego, CA, USA) with analysis of variance (ANOVA) where appropriate. The fluorescent intensity was calculated by NIH Image J software. P < 0.05 was significantly different.

AUTHOR CONTRIBUTIONS

J.N. and X.L. conceived the study. J.N., P.X., X.W., X.Y., and H.X. performed the experiments. J.N., K.L., S.L., and X.L. analyzed the data and wrote the manuscript. All the authors reviewed the manuscript and approved the final version of the paper.

CONFLICTS OF INTEREST

All the authors declare that they have no conflicts of interest.

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