Nucleosome remodeling and deacetylation complex and MBD3 influence mouse embryonic stem cell naïve pluripotency under inhibition of protein kinase C

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The pluripotency of naïve mouse embryonic stem cells (mES) is regulated by multiple signaling pathways, with inhibition of protein kinase C (PKCi) playing a particularly important role in maintaining naive mES. However, the regulatory function of nucleosome remodeling and deacetylase (NuRD) complex in mES cultured in a PKCi system is unknown. We found that, compared with 2iL-derived mES, PKCi-derived mES showed low mRNA expression of NuRD complex subunits, including MBD3, HDAC1/HDAC2, MTA1, and RbAP46/RbAP48. Western blot showed that PKCi-derived mES expressed lower protein levels of MBD3 and HDAC2 at passage 3, as well as MBD3, HDAC2, and MTA1 at passage 10, indicating that PKC suppressed NuRD complex expression. Knockdown of MBD3 increased PKCi-derived mES pluripotency by increasing NANOG and OCT4 expression and colony formation. By contrast, overexpression of MBD3 or removal of PKC inhibitor-induced differentiation of mES, results in reduced NANOG, OCT4, and REX1 expression and colony formation, increased differentiation-related gene expression, and differentiation into flat cells. Knockdown of MBD3 in mES upon PKC inhibitor removal partially reversed cell differentiation. Our results show that the regulatory NuRD complex and its MBD3 subunit influence the naïve pluripotency of mES cultured in a PKCi system.

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INTRODUCTION

The nuclear remodeling and deacetylation (NuRD) complex is an abundant and conserved regulator of chromatin structure remodeling and transcriptional repression [1, 2]. The NuRD complex contains several subunits (e.g., methyl-CpG-binding domain protein MBD2/3, histone deacetylase core proteins HDAC1/2, metastasis-associated protein MTA1/2/3, ATP-dependent nucleosome remodeling enzyme CHD3/4, histone-binding/chaperone proteins RbAP46/48, zinc-finger proteins p66o/B, and DOC1), mediates two major biological functions: nucleosome remodeling in chromatin formation and histone deacyetilation, resulting in the silencing of gene transcription [3]. NuRD was recently found to modulate chromatin structure at regulatory elements of active transcription sites, thereby regulating gene expression in a finely tuned manner [1].

Embryonic stem cells (ES), first derived by Evans and Kaufman in 1981 [4], possess the characteristics of self-renewal, indefinite proliferation in vitro, multi-lineage differentiation, and germline transmission. Mouse ES (mES) can exist in naïve or primed states [5]. Two different culture systems can maintain mES naïve pluripotency: 2iL and inhibition of the PKC signaling pathway (PKCi). The 2iL system includes leukemia inhibitory factor (LIF), which activates transcription factor signal transducers and activators of transcription 3 (STAT3) and the small-molecule inhibitors PD0325901 and CHIR99021, which in turn inhibit mitogen-activated protein kinase (MAPK) and glycogen synthase kinase 3 (GSK3) pathways, respectively [6, 7]. 2iL-derived mES are frequently used in stem cell studies. The PKCi system also maintains the naïve state of mES and rat ESs [8, 9]. Members of the PKC family are intracellular mediators of several hormones, neurotransmitters, phorbol esters, and tumor promoters that play essential roles in growth regulation, ES self-renewal, cell differentiation, neurotransmission, and cell death [10]. In particular, inhibition of the PKC–NF-κB–microRNA-21/microRNA-29 axis is key to maintaining ES self-renewal and naïve pluripotency [8, 9]. However, the role of NuRD subunits in PKCi-derived naïve mES, as well as differences in NuRD complex expression between 2iL- and PKCi-derived mES, are unknown.

Previous studies revealed the mechanisms by which NuRD maintains stem cell pluripotency via the regulation of gene expression related to cell plasticity, self-renewal, and differentiation during development [1, 11]. MBD3, a specific ES NuRD subunit, is essential for early embryogenesis and the development of pluripotent stem cells [12, 13]. As a methylated DNA binding protein that scaffolds other NuRD subunits, MBD3 is crucial for recruiting other subunits and assembling the NuRD complex [14, 15] and modulates transcriptional heterogeneity, and maintains ES lineage commitment by directly regulating the expression...
of pluripotency genes in ESs [16]. MBD3 is required for commitment to a full spectrum of embryonic lineages [15, 17, 18]. MBD3 \(-/-\) ES maintains the expression of pluripotency genes such as OCT4 and NANOG but fails to form stable NuRD complexes, exhibit severe defects in differentiation [12], and shows silencing of Oct4 expression upon withdrawal of LIF [15]. However, the role of MBD3 in the growth and self-renewal of mES colonies in the PKCi system is unclear. In particular, identifying differences in target gene regulatory loci specifically recognized by MBD3 between 2iL- and PKCi-derived mES will provide a fuller
understanding of how different signaling pathways induce naïve mES.

In this study, we examined differences in NuRD complex RNA and protein expression in naïve mES derived in 2iL versus PKCi culture systems. We found that NuRD subunits are present in both types of mES, suggesting that NuRD is necessary for maintaining ES self-renewal and pluripotency. However, the RNA expression of most NuRD subunits, including HDAC2, MBD3, and MTA1, was lower in PKCi-derived mES than in 2iL-derived mES. Furthermore, using overexpression and knock-down of MBD3, we demonstrate that NuRD and MBD3 play important roles in PKCi-derived mES self-renewal and naive pluripotency by modulating the expression of pluripotency and differentiation genes.

RESULTS

PKCi downregulated expression of the NuRD complex in mES

To investigate how the PKCi culture system affects the expression of NuRD complex components, we first measured the mRNA expression of the NuRD complex in PKCi-derived mES at passage 3. Quantitative polymerase chain reaction (qPCR) showed that compared with 2iL-derived mES, PKCi-derived mES showed lower mRNA levels of MBD3 (from 1.0 to 0.2), HDAC2 (from 1.0 to 0.4), HDAC1 (from 1.0 to 0.3), MTA1 (from 1.0 to 0.1), RbAp46 (from 1.0 to 0.02), RbAp48 (from 1.0 to 0.05), and p66α (from 1.0 to 0.8) (P < 0.05; Fig. 1A), whereas CHD3, p66β, and DOC1 levels were similar between two mES types. Western blot further revealed that PKCi-derived mES showed lower protein levels of MBD3 (from 1.0 to 0.5) and HDAC2 (from 1.0 to 0.9, P < 0.05; Fig. 1B). These changes in RNA and protein expression were maintained (Fig. 1C, D), except for further reduced protein MTA1 (Fig. 1D) at passage 10. When we examined the expression of genes downstream and targets of NuRD [1], we found significantly lower expression of carbonyl reductase 3 (CBR3) and high-temperature requirement serine protease A1 (HTRA1) mRNA in PKCi-derived mES (Fig. 1E).

PKCi-derived mES expressed naive pluripotency genes and possessed germline transmission

Compared with mouse embryonic fibroblasts, PKCi- and 2iL-derived mES showed higher mRNA expression of the pluripotency markers such as NANOG, OCT4, C-MYC, and SOX2 (Fig. S1A) and naive-state markers FGFR4, NROB1, REX1, and KLFC4 (Fig. S1B) but not primed-state markers FGFR5 and T (P < 0.05; Fig. S1C).

Signaling pathways involved in PKCi-derived mES self-renewal

As WNT, ERK, and AKT signaling pathways are involved in the self-renewal of 2iL-derived mES [6, 7], we examined key proteins in these pathways in both 2iL- and PKCi-derived mES. Western blot showed that the ratio of phosphorylated (p)-β-catenin/β-catenin was increased in PKCi-derived mES (P < 0.01), whereas p-ERK/ERK and p-AKT/AKT ratios were similar between two mES types (Fig. 2A, B). In addition, levels of HDAC5, a key protein in the PKCε signaling pathway [19], were reduced in PKCi-derived mES (P < 0.01; Fig. 2).

MBD3 knockdown promoted PKCi-derived mES self-renewal

To understand the role of MBD3 in mES self-renewal, we knocked down MBD3 in PKCi-derived mES. Compared with PKCi and shNC control groups, MBD3 mRNA (from 1.0 to 0.3) (Fig. S3A) and protein (from 1.0 to 0.4) (Fig. S3B) expression were reduced in MBD3 knocked-down mES (P < 0.01). MBD3 knockdown increased the mRNA levels of pluripotency markers NANOG (from 1.0 to 3.7) and OCT4 (from 1.0 to 2.6) (P < 0.01), whereas the expression of other naive pluripotency, primed-state, and differentiation genes was unchanged (Fig. S3C). As expected, Western blot revealed that the protein levels of NANOG (from 1.0 to 2.0) and OCT4 (from 1.0 to 3.0) were also increased (P < 0.01; Fig. 3D). Immunostaining indicated that MBD3 knockdown did not affect the morphology of mES (Supporting Information Fig. S1D). Although AP staining indicated that compared with PKCi and control groups, MBD3 knockdown did not affect the total number of AP-positive colonies, the percentage of mixed colonies increased (from 42.1% to 46.4%) and differentiated colonies decreased (from 9.1% to 3.8%), respectively (P < 0.05; Fig. 3E).

Overexpression of MBD3-induced mES differentiation

Compared with PKCi and FUW-M2rtTA control groups, MBD3 overexpression increased mRNA (from 1.0 to 2.0; Fig. 4A) and protein (from 1.0 to 1.4, P < 0.05; Fig. 4B) levels of MBD3 in PKCi-derived mES. qPCR showed that MBD3 overexpression significantly decreased mRNA levels of the pluripotency markers NANOG (from 1.0 to 0.5) and OCT4 (from 1.0 to 0.8) and the naive-state marker REX1 (from 1.0 to 0.2) and increased mRNAs levels of the primed-state marker FGFR5 (from 1.0 to 4.2). mRNA levels of the differentiation markers CK8 (from 1.0 to 5.6), cTnT (from 1.0 to 3.6), BMP4 (from 1.0 to 1.9), DESMIN (from 1.0 to 1.4), PAX6 (from 1.0 to 4.4), and SOX17 (from 1.0 to 1.8) were increased in MBD3 overexpressing mES (P < 0.05; Fig. 4C). Also, MBD3 overexpression decreased protein levels of NANOG (from 1.0 to 0.3) and OCT4 (from 1.0 to 0.2) and increased protein levels of cTnT (from 1.0 to 3.0, P < 0.05; Fig. 4D).

MBD3 overexpression induced mES differentiation into flat cells that did not express NANOG (Supporting Information Fig. S2C). Compared with the PKCi control group, MBD3 overexpression reduced the total number of AP-positive colonies (from 178 to 50) and the percentage of undifferentiated (from 45.4% to 6.1%) and mixed (from 46.0% to 13.4%) colonies, but increased the percentage of differentiated colonies (from 8.7% to 80.5%, P < 0.05; Fig. 4E).

Removal of PKC inhibitor increased MBD3 expression and mES differentiation, which was partially rescued by MBD3 knockdown

When the PKC inhibitor Gö6983 was removed from the PKCi culture medium for 48 h, MBD3 mRNA (from 1.0 to 2.1) and protein (from 1.0 to 1.7) levels increased in mES (P < 0.05; Fig. 5A, B). This change was associated with decreased mRNA levels of pluripotency markers NANOG (from 1.0 to 0.2), OCT4 (from 1.0 to 0.2), and SOX2 (from 1.0 to 0.3), decreased levels of naïve-state markers KLFC4 (from 1.0 to 0.4), FGFR4 (from 1.0 to 0.2), NROB1 (from 1.0 to 0.2), and REX1 (from 1.0 to 0.2); and increased mRNA levels of
PKC inhibitor removal induced mES differentiation, and a few cell colonies existed that were negative for AP staining (Fig. 5E). Quantitative analysis showed that PKC inhibitor removal significantly reduced the number of AP-positive colonies (from 171 to 26), and the percentage of mixed (from 42.5% to 27.4%) and undifferentiated (from 51.3% to 0%) colonies and increased the percentage of differentiated colonies (from 7.2% to 72.6%; Fig. 5E). However, 

**DISCUSSION**

We demonstrated that inhibition of the PKC-signaling pathway by G06983 reduced the expression of NuRD components at both RNA (MBD3, HDAC1, HDAC2, MTA1, RbAP46/RbAP48, and p66α) and protein (MBD3 and HDAC2 at passage 3, further MTA1 at passage 10) levels in mES. The PKC-signaling pathway is a complex signal transduction network that participates in other signaling pathways, such as MEK/ERK [20], CREB [21], PKCζ-NF-κB [8, 9], PKCµ [19], and GSK3β [22]. Small molecules, such as PD0325901 and CHIR99021 inhibit MEK/ERK and GSK3β pathways, which together with activation of JAK-STAT3 maintain mES naïve pluripotency [5, 7]. In our study, we found comparable ratios of p-ERK/ERK and p-AKT/AKT between 2iL- and PKCi-derived mES, suggesting that inhibition of PKC by the small molecule G06983 plays a similar role as PD0325901 in suppressing MEK/ERK pathway. On the other hand, we observed an elevated ratio of p-β-catenin/β-catenin in PKCi-derived mES, implying that G06983 inhibits GSK3β more effectively than CHIR99021. A previous study [8] reported that
GSK3, ERK1/2, AKT, and their respective downstream target genes β-catenin, RSK1, and STAT3 were not phosphorylated in the PKCi system, suggesting that PKCi regulates mES self-renewal independently of traditional signaling pathways. However, we found that the p-ERK/ERK ratio was lower in the PKCi system compared with that after PKC inhibitor removal, indicating that the ERK-signaling pathway is involved in PKCi-derived mES self-renewal.

Interestingly, in human ES [23], FGF2 activates PI3K/AKT, MEK/ERK1/2, and PKC isoforms (i.e., PKCδ/ε/ζ), resulting in phosphorylation of GSK3β, and that activation of AKT signaling promotes self-renewal whereas activation of GSK3β and ERK1/2 induces differentiation. In addition, PKCi system is involved in the downregulation of HDAC5, a key protein in the PKCµ pathway, which maintains mES self-renewal and prevents cell lineage...
Fig. 4 Overexpression of MBD3-induced PKCi-derived mES differentiation. A qPCR showed that mRNA levels of MBD3 increased after PKCi-derived mES at passage 5 were transfected with FUW-MBD3, with the PKCi and FUW-M2rtTA plasmid groups used as controls. B MBD3 and β-actin protein levels were evaluated by Western blot after PKCi-derived mES were transfected with FUW-MBD3 (upper panel). Quantitative density analysis showed that protein levels of MBD3 increased after transfection with FUW-MBD3 (lower panel). C qPCR showed decreased mRNA levels of pluripotency genes NANOG and OCT4 and naïve-state marker REX1, but increased mRNA levels of primed-state marker FGF5 after MBD3 was overexpressed (left panel). MBD3 overexpression also increased mRNA levels of the endoderm marker CK8, mesoderm markers cTnT, BMP4, and DESMIN, and ectoderm markers Pax6 and Sox17 (right panel). D Western blot detection of NANOG, OCT4, REX1, FGF5, cTnT, and β-actin protein expression after MBD3 overexpression (left panel). Quantitative density analysis showed that MBD3 overexpression decreased protein levels of NANOG, OCT4, and REX1, but increased protein levels of FGF5 and cTnT (right panel). E MBD3 overexpression induced PKCi-derived mES differentiation and resulted in a loss of AP staining. Scale bar, 200 μm (upper panel). Overexpression of MBD3 reduced the total number of AP-positive colonies (left panel), decreased the percentage of undifferentiated and mixed colonies, and increased the percentage of differentiated colonies (right panel). Data were shown as mean ± SD (n = 3). The letters a and b indicated significant differences among groups (P < 0.05).
commitment and differentiation, moreover, the inhibition of PKCµ suppresses myoblast differentiation by inhibiting MYOD and myocyte-specific enhancer factor 2C [19]. Key transcription factors are pivotal for maintaining gene expression patterns, chromatin structure, and the epigenetic landscape allowing ES self-renewal [5, 7, 8]. These transcription factors affect the expression of other repressive regulatory factors, including the NuRD complex. Although transcription levels of HDAC1, MTA1, CHD3, RbAP46, and RbAP48 were changed, protein levels were not affected, probably due to the regulation of translation and the lower
threshold of RNA required for translation. The mechanism that PKCI down-regulates the expression of MBD3 and other genes is unknown. In the present study, a reduction in NurD expression was confirmed by decreased expression of NurD targeted CBP3 and HTRA1 mRNA [1] in PKC-i derived mES. It is assumed that NurD targeted gene may play a role in regulating the naive pluripotency.

Previous studies using G60983, a selective inhibitor of PKC isoforms, show that the PKC-β-NF-kB-miRNA-21/miRNA-29 regulatory signaling axis plays a critical role in inducing mES lineage commitment and is capable of maintaining mES and rat ES-specific epigenetic modifications and self-renewal [8, 9]. Our study confirms this finding by showing that knockdown of PKCβ partially rescued the amount of AP-positive mES colonies upon the removal of PKC inhibitor (Fig. 5A,5B), indicating that inhibition of the NF-kB signaling pathway plays an important role in maintaining PKC-i derived mES pluripotency.

The NurD complex is an abundant and conserved remodeling complex repressor that increases nucleosome density and fine-tunes differential gene expression, even at active transcription sites [1]. MBD3, a key NurD subunit, is necessary for the development of pluripotent cells, as MBD3-depleted mES are viable but fail to form stable NurD complexes, which severely affect their cell commitment and differentiation capacities [12]. In the present study, we found that MBD3 plays an important role in mES self-renewal and differentiation. In the PKCI system, knockdown of MBD3 increased the expression of the pluripotency genes NANOG and OCT4, thereby promoting mES self-renewal. Interestingly, knockdown of MBD3 did not affect other pluripotency genes such as KLF4, SOX2, REX1, NROB1, or FGF4. In particular, previous study reported that KLF4, a naive-state marker and direct downstream target of LIF/STAT3 [24, 25], mediates the self-renewal-promoting effects of GBX2 in ESs [26]. Therefore, we assume that the LIF/STAT3-signaling pathway does not play a dominant role in the PKCI system. In fact, a previous study found mES devoid of STAT3, suggesting that LIF/STAT3 signaling does not instruct ES self-renewal but may instead act in unrefined culture conditions [7]. By contrast, we observed that MBD3 overexpression and/or PKC inhibitor removal increased MBD3 mRNA and protein levels, downregulated mRNA levels of pluripotency genes such as NANOG and OCT4 and naive-state genes such as REX1, and upregulated the expression of differentiation genes such as the endodermal marker CK6, mesoderm markers cTnT, BMP4, and DESMIN, and ectoderm markers PAX6 and SOX17. Overexpression of MBD3 also reduced protein levels of NANOG, OCT4, and REX1 but increased protein levels of the primed-state marker FGF5 and differentiation marker cTnT. As a result, the number of ES colonies decreased due to cell differentiation. Furthermore, MBD3 knockdown by RNA interference partially reversed the cell differentiation induced by PKC inhibitor removal. Therefore, our findings support previous findings that the repression of genes related to lineage commitment and cell differentiation is necessary for maintaining mES self-renewal [7, 27]. MBD3 directly regulates pluripotency gene transcription in ESs [16], with one study reporting that MBD3 and Brg1 antagonistically regulate a common set of genes by regulating promoter nucleosome occupancy to maintain pluripotency of embryonic stem cells in mouse [28]. The balance between repressive NurD and activating chromatin remodeling complex BAF may finely tune gene expression specific to stem cell pluripotency [1]. As a result, mES exhibit a transient cell state poised for cell lineage differentiation during development [5, 29].

In summary, we found that the PKCI culture system reduced the expression of the repressive and epigenetic regulatory NurD complex and its MBD3, HDAC2, and MTA1 subunits in mES. This reduction in NurD promoted the expression of pluripotency genes that maintain the naive state of mES. Functional knockdown and overexpression experiments revealed that MBD3, a key NurD subunit, plays an important role in regulating the expression of genes responsible for mES pluripotency. This demonstration of the transcriptional/translational regulatory effects of the NurD complex on a wide spectrum of developmental genes increases our understanding of ES pluripotency, lineage commitment, and differentiation.

MATERIALS AND METHODS

Chemicals and reagents

Unless otherwise stated, all chemicals and reagents were purchased from Sigma-Aldrich (St. Louis, MO, USA).

Animal maintenance, hormone-induced superovulation, and blastocyst collection

Animal experimental protocols were approved by the Animal Care and Use Committee of Nanjing Normal University (NSD-2013-30) and were performed according to guidelines from the US National Institutes of Health. At 6–8 weeks of age, female C57BL/6J mice were intrauterinally injected with 7.5 IU pregnant mare serum gonadotropin (Ningbo Second Hormone Factory, China). Forty-eight hours later, mice were intrauterinically injected with 7.5 IU human chorionic gonadotropin (Ningbo Second Hormone Factory) and mated with males (1:1). Blastocysts were flushed from the uterus with M2 medium on day 3.5.

De novo derivation of mES

Blastocysts were seeded on Mitomycin C-treated mouse embryonic fibroblasts on 0.1% gelatin-coated plates (ES-006-B, Millipore, USA) with PKC inhibitor (5 μM G60983, 133053-19-7, Selleck, USA) in basic culture medium supplemented with Dulbecco modified Eagle’s medium (DMEM; 10829016, Gibco, USA) containing 15% knockout serum replacement (10828028, Gibco), 1% penicillin/streptomycin (S3V0010, HyClone, USA), 2 mM glutamine (35050061, Gibco), 1 mM sodium pyruvate (11360088, Cell Death Discovery (2022) 8:344
| Gene symbol | Forward primer (5′–3′) | Reverse primer (5′–3′) | PCR condition | Size (bp) |
|-------------|------------------------|------------------------|---------------|----------|
| **NANOG**   | AGAAGTACCTCACGTCCACAGC| AGATGCGTTCACCAGATAGCC | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 224      |
| **OCT4**    | GAGGAGCCGCAAACTGAGG   | TGGTAGTGATCTGTCGAGAGG | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 163      |
| **C-MYC**   | GAGCCCCACCAACATCCTCT  | GGGGAACAGATCTGGCAGT   | 95 °C 1 s, annealing/extension 60 °C 60 s, 40 cycles | 185      |
| **SOX2**    | GTGATCCCTCTCCACACTCCAGG | TGGTGCTTAATGGGCAGTGCC | 95 °C 1 s, annealing/extension 60 °C 60 s, 40 cycles | 172      |
| **FGF4**    | GTGTTGAGCATCTCCGAGTGG | GCGTAGGATTGATGACGTTGT | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 146      |
| **NROB1**   | TTGACACCAAGATGCCTCTAC | AAGGGCAGCTCGATCAGCC   | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 182      |
| **REX1**    | GCATTGCCACACCCATCATC  | CTGGTGAGCAAGACATCCAG | 95 °C 1 s, annealing/extension 60 °C 60 s, 40 cycles | 143      |
| **KL4**     | ACTGTCACCCTGCGCCTCTCT | CCCTCTTTTGCTGCGCCTCT | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 165      |
| **FGF5**    | GCTCGGAACATAGCAGTTTTT | CCGTAATTGCTGTTACACAC | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 151      |
| **T**       | ACCTATGCGGACAATTCATC  | CAGACACAGAGACTGGGATAC | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 155      |
| **Ctb**     | AGATGGAAGCGCTCAGGCTCT | AGCTTAGGATCGTTCAAAGCT | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 187      |
| **BMP4**    | ATCCGGAAAGCTCGATTGGA | GAGATCACCCTATTCTGCTGG | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 100      |
| **DESMIN**  | AGAAAGTGCAAGGAGGGAG  | CTCAAGATGTTTTTACGCC | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 156      |
| **PAX6**    | CGGAAAGCTCAGGAAATAG  | CCTTCTTTGCTGCTTCAAGTT | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 145      |
| **SOX17**   | GCGTGGAACAGAGCGGCCTTCTTTT | GGACACTCATGACTGAGAGACTG | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 101      |
| **cTNT**    | AGACTGGAGTGAAGAGAGGAG | CTGGGGCTTGGGTTGGTTGTC | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 186      |
| **HDAC1**   | ACAAGCCCATGCTGGGAGA | TCAACACACCTATCAACACC | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 154      |
| **HDAC2**   | CGTGTGTGATTGACCTTGTTT | AGAACCTGATGCTCTTGACT | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 150      |
| **MBD3**    | CAGGCTATGCGTCTCTTACT | CTGTCACAGATAGCCTTGGCC | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 129      |
| **MTA1**    | ACAACAGACACACAGGAAATG | CAACTGCGGAGACAGGAAACAG | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 200      |
| **CHD3**    | AGACTGGAGGAGCCAGGATAG | TCAGAAGGAAAGTTGCGGATG | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 156      |
| **RbAP46**  | TGACTGATGTCCTGGATTGG | CAGATGAACTGCTGGTTAGG | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 158      |
| **RbAP48**  | CATACGACAGATGAGGAGGAG | TGTGAGTCAACCGAGTGGC | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 186      |
| **DOC1**    | GGAAGCTGGCTAACACTCCCTACT | ACTCTGAGGACCCGCTTCATT | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 126      |
| **P66α**    | TTGGCAAGACCTACTCCGACC | GACTTTGCCAGACCGATTA | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 284      |
| **P66β**    | CTCGCGATGCTCTGCTTACAAGA | ATGGCTGACAAATGACGCTT | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 105      |
| **GAPDH**   | GTGGCGAAAGTGGAAGATGG | CTCCGTGAAAGATGGTGATG | 95 °C 15 s, annealing/extension 60 °C 60 s, 40 cycles | 164      |
Gibco), and 0.1 mM 2-mercaptoethanol (ES-007-E, Millipore). After culture for 7 days, outgrowths were collected and digested into single-cell suspensions with accutase (A1110501, Gibco) and re-seeded in new plates coated with feeder cells. mES passages was performed by incubating colonies with accutase, followed by plating at a density of 1 x 10^5 cells/cm^2 into a new 24-well plate coated with new feeder cells at 3- to 4-day intervals. Collected mES were frozen in a cryopreservation medium with 90% fetal bovine serum (FBS; SH300070.03, HyClone) and 10% dimethyl sulfoxide (DMSO) and stored in liquid nitrogen.

**MBD3 overexpression in mES**

FUW-MBD3 (#52356) and control FUW-M2rtTA (#20342) were purchased from Addgene. Lentiviral infection and overexpression were performed as previously described [30]. Briefly, 293T cells were cultured for 2 days in DMEM (C11995500BT, Gibco) supplemented with 10% FBS (v/v) at 37 °C in 5% CO2. After 24 h of incubation, the medium was changed. After transfection for 48 h, viral supernatants were collected and filtered (0.45-µm pore size; Millipore) to infect mES. For lentiviral transfection, mES at 70–80% confluency were infected with filtered viral supernatants (FUW-MBD3 or FUW-M2rtTA) supplemented with an equal volume of fresh PKCi medium. Twelve hours later, mES were repeated infected with viral supernatants up to four times within 48 h as needed.

**RNA interference in mES**

Lentiviral supernatants containing short hairpin RNA (shRNA) targeting mouse MBD3 mRNA (shMBD3), PKÇζ1, PKÇζ2, or shRNA negative control (shNC) were purchased from GenePharma (Shanghai, China). The shPKÇζ1 KD1 target sequence was GGGACGAAGTGCTCAGG, shPKÇζ1 KD2 was TTCTCCGAACGTGTCACGT. After transfection, colonies with accutase, followed by plating at a density of 1 × 10^3 cells/cm^2 for 7 days, outgrowths were collected and digested into single-cell suspensions with accutase (A1110501, Gibco), and 0.1 mM 2-mercaptoethanol (ES-007-E, Millipore). After culture for 80% confluence, mES were transfected with FUW-MBD3 or control FUW-M2rtTA along with the viral packaging plasmids pSPAX and PMD2.G (5:3:2) with Lipofectamine 2000 reagent (1947415, Invitrogen, USA) at a 1:2 ratio of DNA (g) to Lipofectamine 2000 medium. After 6 h, the medium was changed. After transfection for 48 h, viral supernatants were collected and filtered (0.45-µm pore size; Millipore) to infect mES. For lentiviral transfection, mES at 70–80% confluency were transfected with FUW-MBD3 or control FUW-M2rtTA supplemented with an equal volume of fresh PKCi medium. Twelve hours later, mES were repeated infected with viral supernatants up to four times within 48 h as needed.

**qPCR**

Total RNA was extracted from mES with Trizol reagent (T9424). Reverse transcription reactions were performed with 1 µg RNA using HiScript II Reverse Transcriptase (R223-01, Vazyme, China). Complementary DNA was used as a template, and 2xSYBR Green Fast qPCR Mix with High Rox substrates (REF 11745832910, Roche, Switzerland) for 20–30 min at room temperature, washed with DPBS, and observed under an inverted fluorescence microscope.

**Alkaline phosphatase staining**

mES were washed with DPBS three times, fixed with 4% paraformaldehyde for 10 min at room temperature, washed with DPBS three times, incubated with 0.2% Triton-X 100 (Solarbio, China) for 5 min, washed with DPBS three times, incubated with 2% FBS (HyClone) for 30 min at room temperature to block nonspecific binding, and incubated with primary antibodies at 4 °C overnight. The next day, cells were washed with DPBS three times, incubated with secondary antibody conjugated with FITC (goat anti-rabbit IgG (H + L), 1:300, AS011, Abclonal) for 2 h at room temperature, washed with DPBS, and stained with 100 ng/ml DAPI (SunShine, China) for 10 min at room temperature in the dark. Finally, after washing with DPBS, cells were observed under an inverted fluorescence microscope.

**Immunocytochemical staining for pluripotency markers**

Immunostaining experiments were performed with the pluripotency marker NANOG (1:200, Lannuo Biotechnologies). Briefly, mES were washed with Dulbecco’s phosphate-buffered saline (DBPS) three times, fixed with 4% paraformaldehyde for 10 min at room temperature, washed with DBPS three times, incubated with 0.2% Triton-X 100 (Solarbio, China) for 5 min, washed with DPBS three times, incubated with 2% FBS (HyClone) for 30 min at room temperature to block nonspecific binding, and incubated with primary antibodies at 4 °C overnight. The next day, cells were washed with DPBS three times, incubated with secondary antibody conjugated with FITC (goat anti-rabbit IgG (H + L), 1:300, AS011, Abclonal) for 2 h at room temperature, washed with DBPS, and stained with 100 ng/ml DAPI (SunShine, China) for 10 min at room temperature in the dark. After washing with DPBS, cells were observed under an inverted fluorescence microscope.

**Western blot**

mES total protein was collected with lysis buffer and quantified using a BCA Kit (GK5012, Beyotime Biotechnology, China). Western blot analysis was performed as previously described [31]. Briefly, 6 µg total protein was loaded into each well of a 12% gel for SDS–PAGE. After electrophoresis, separated proteins were transferred to a polyvinylidene fluoride membrane (0301004001, Roche, Basel, Switzerland) by electrotransfer. Membranes were blocked with 5% non-fat powdered milk (A600669, Sangon Biotech, China) for 1 h, after which membranes were washed with tris-buffered saline containing Tween 20 (TBST; 9005-645, Sangon Biotech) and incubated with anti-β-actin (1:1000, 9561T, Sangon Biotech) and incubated with anti-β-actin (1:1000, A19657, Abclonal, China), anti-p-ERK (1:1000, 3510, CST), anti-ERK (1:1000, 4695, CST), anti-p-AKT (1:1000, AP1214, Abclonal), anti-AKT (1:1000, 9602, Abclonal), anti-HDAC3 (1:1000, A0238, Abclonal), anti-HDAC1 (1:1000, A0238, Abclonal), anti-HDAC2 (1:1000, A0238, Abclonal), anti-MTA1 (1:1000, A16085, Abclonal), anti-MBD3 (1:1000, A2251, Lot 00046180101, Abclonal), anti-CHD3 (1:1000, A2221, Abclonal), anti-RbAP46 (1:1000, A6967, Abclonal), anti-RbAP48 (1:1000, A13934, Abclonal), anti-NANOG (1:1000, Lannuo Biotechnology, China), anti-OCT4 (1:1000, Lannuo Biotechnology), anti-REX1 (1:1000, Lannuo Biotechnology), anti-FGF5 (1:1000, ab88118, Abcam), or anti-cTNT (1:1000, A4914, Abclonal) antibodies overnight at 4 °C. The next day, membranes were washed with TBST and incubated with horseradish peroxidase-conjugated goat anti-rabbit secondary antibody (1:5000, B513278, BioWorld) for 1 h. After washing, membranes were processed using an enhanced chemiluminescence reagent (E411-04, Vazyme), and protein bands were visualized using a LAS-4000 imager (Tanon, Shanghai, China). β-actin (1:1000, AC026, Abclonal) was used as an internal control. Target protein expression levels were normalized to those of β-actin. Values from the PKCi group were defined as 1.0 and were used for comparisons with other treatments.

**Statistical analysis**

All experiments were repeated at least three times. mES derived from either PKCi or LIF-2i were from different blastocysts, but mES after derivation in two conditions were the same strains to exclude the possibility of variations. Data were analyzed using SPSS version 18.0 (SPSS, Inc., Chicago, IL, USA) and were shown as mean ± standard deviation (SD). Statistical comparisons were performed using analysis of variance (ANOVA). The letters a, b, and c indicated significant differences among groups (P < 0.05).

**DATA AVAILABILITY**

All data related to this paper may be requested from the corresponding author.

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AUTHOR CONTRIBUTIONS

FD, CH, LA, and YD designed the experiments; YD performed the experiments; JS, NH, and CH analyzed the experimental data; FD and YD wrote the manuscript; CH and FD discussed the manuscript arrangement.

COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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