Interaction of the primordial germ cell-specific protein C2EIP with PTCH2 directs differentiation of embryonic stem cells via HH signaling activation

Qisheng Zuo1, Kai Jin1, Jiuzhou Song2, Yani Zhang1, Guohong Chen1 and Bichun Li1

Abstract
Although many marker genes for germ cell differentiation have been identified, genes that specifically regulate primordial germ cell (PGC) generation are more difficult to determine. In the current study, we confirmed that C2EIP is a PGC marker gene that regulates differentiation by influencing the expression of pluripotency-associated genes such as Oct4 and Sox2. Knockout of C2EIP during embryonic development reduced PGC generation efficiency 1.5-fold, whereas C2EIP overexpression nearly doubled the generation efficiency both in vitro and in vivo. C2EIP encodes a cytoplasmic protein that interacted with PTCH2 at the intracellular membrane, promoted PTCH2 ubiquitination, activated the Hedgehog (HH) signaling pathway via competitive inhibition of the GPCR-like protein SMO, and positively regulated PGC generation. Activation and expression of C2EIP are regulated by the transcription factor STAT1, histone acetylation, and promoter methylation. Our data suggest that C2EIP is a novel, specific indicator of PGC generation whose gene product regulates embryonic stem cell differentiation by activating the HH signaling pathway via PTCH2 modification.

Introduction
Primordial germ cells (PGC) can be used for therapeutic purposes and genetic modification as an alternative to embryonic stem cells1,2. Unfortunately, harvesting plentiful PGC in vitro is often problematic because of the limited understanding of their ontogenetic mechanism, which greatly confines their clinical application3. Multiple gene products are specifically involved in PGC differentiation4. For example, hypoxia induces PGC generation by maintaining HIF1 expression and down-regulating Oct4 expression5,6, whereas Blimp1 and Prdm14 regulate PGC generation in mice by epigenetically modifying and re-programming them to differentiate into spermatogonial stem cells (SSC)7-10. Moreover, RNAi-mediated knockout of LIN28 downregulates expression of multiple germ cell surface markers during STELLA+ cell differentiation, including key PGC development genes such as Blimp1, Prdm14, and endogenous STELLA11,12. Although these genes can mark PGC and be used to track their migration and colonization, not all of them are uniquely expressed in PGC and their regulatory mechanisms are not conspicuous.

In the present study, we identified a novel PGC-specific marker gene, C2EIP13,14, and characterized its molecular function in detail. We previously reported that C2EIP was not noticeably expressed in embryonic stem cells (ESC), SSC, or chick embryo fibroblasts but are specifically expressed in PGC. Therefore, we speculated that C2EIP is likely a novel key genetic marker for PGC generation and differentiation. This hypothesis is supported by earlier...
studies showing that CRISPR/Cas9-mediated knockout of C2EIP in chick embryos significantly limited their development and reduced PGC generation efficiency, and C2EIP overexpression promoted PGC generation. The GST pull-down assay showed that C2EIP interacted with PTCH2, an important membrane protein of the Hedgehog (HH) signaling pathway, which can upregulate germ cell differentiation and PGC generation. However, whether C2EIP function is associated with HH signal activation remains unknown. Thus, we investigated the ubiquitination and dephosphorylation status of PTCH2 in response to different expression levels of C2EIP. We found that C2EIP overexpression induced ubiquitination and dephosphorylation of PTCH2 and upregulated PGC generation by releasing SMO protein and activating the HH signaling pathway. We also determined that C2EIP expression was regulated by transcription factor STAT1 mediated by the CG loci and affects promoter activity. Studies have shown that epigenetic factors (DNA methylation, histone acetylation, and transcription factors) affect promoter activity. We predicted, by bioinformatics analysis, that the cloned C2EIP promoter fragment does not contain CpG islands but does contain a small number of CG loci, suggesting that the C2EIP gene promoter activity is not likely to be affected by DNA methylation. To verify this hypothesis, we linked the C2EIP promoter fragment into the pGL3-Basic vector to construct the pGL3-787 recombinant vector, which we transfected into DF-1 cells that we treated with or without 10 µmol/L DNA methylation inhibitor (5-Zacd). The cells were collected for the dual luciferase reporter assay, which showed that the inhibition of methylation resulted in significant upregulation (37.24 ± 3.24) of C2EIP promoter activity compared to it without 5-Zacd treatment (23.41 ± 2.13), indicating that although the C2EIP promoter region does not contain a CpG island, DNA methylation can be mediated by the CG loci and affects promoter activity (Fig. 2b). C2EIP gene expression was detected in cultured PGC after treatment with 5-Zacd, which significantly increased C2EIP expression, confirming the luciferase assay results (Supplementary Figure 2b). While screening for other epigenetic factors affecting the activity of the C2EIP promoter, it was found that treatment of pGL3-787-transfected DF-1 cells with 10 µmol/L of the histone deacetylase inhibitor TSA significantly upregulated C2EIP promoter activity (68.47 ± 3.11) compared to it without TSA treatment (22.35 ± 1.48) (Fig. 2c), indicating that histone acetylation can increase C2EIP promoter activity. CHIP-qPCR results also show that histone acetylation was enriched in the C2EIP promoter region (Fig. 2d).

C2EIP is highly expressed in PGC via promoter and histone modification

Because we observed C2EIP more highly expressed in PGC than in ESC or SSC, we speculated that the activity of the C2EIP promoter changes during PGC generation, which leads to differential gene expression in PGC compared to ESC and SSC. Therefore, we identified the promoter region of the C2EIP gene based on the original promoter TATAbox and CAAATbox and cloned the C2EIP promoter fragment to replace the pEGFP-N1 promoter in a pcDNA3.0-EGFP recombinant vector. The constructed pcDNA3.0-EGFP vector (Fig. 2a) can express green fluorescent protein in DF-1 cells (Supplementary Figure 2a), confirming the C2EIP promoter activity.

Studies have shown that epigenetic factors (DNA methylation, histone acetylation, and transcription factors) affect promoter activity. We predicted, by bioinformatics analysis, that the cloned C2EIP promoter fragment does not contain CpG islands but does contain a small number of CG loci, suggesting that the C2EIP gene promoter activity is not likely to be affected by DNA methylation. To verify this hypothesis, we linked the C2EIP promoter fragment into the pGL3-Basic vector to construct the pGL3-787 recombinant vector, which we transfected into DF-1 cells that we treated with or without 10 µmol/L DNA methylation inhibitor (5-Zacd). The cells were collected for the dual luciferase reporter assay, which showed that the inhibition of methylation resulted in significant upregulation (37.24 ± 3.24) of C2EIP promoter activity compared to it without 5-Zacd treatment (23.41 ± 2.13), indicating that although the C2EIP promoter region does not contain a CpG island, DNA methylation can be mediated by the CG loci and affects promoter activity (Fig. 2b). C2EIP gene expression was detected in cultured PGC after treatment with 5-Zacd, which significantly increased C2EIP expression, confirming the luciferase assay results (Supplementary Figure 2b). While screening for other epigenetic factors affecting the activity of the C2EIP promoter, it was found that treatment of pGL3-787-transfected DF-1 cells with 10 µmol/L of the histone deacetylase inhibitor TSA significantly upregulated C2EIP promoter activity (68.47 ± 3.11) compared to it without TSA treatment (22.35 ± 1.48) (Fig. 2c), indicating that histone acetylation can increase C2EIP promoter activity. CHIP-qPCR results also show that histone acetylation was enriched in the C2EIP promoter region (Fig. 2d).

Results

C2EIP encodes a cytoplasmic protein that is differentially expressed in PGC

PGC are the ancestral cells of male germ cells and our research group has previously studied their migration, colonization, and differentiation. Bioinformatic analyses revealed that C2EIP is located on the second chromosome and contains a 648-bp coding sequence (CDS) that encodes 215 amino acids with a relative molecular weight of 24.26 kD. C2EIP is a hydrophilic, non-secretory protein situated mainly in the cytoplasm that does not include a signal peptide sequence or special structural domain. Previous RNA-seq results showed that expression of C2EIP (XM-001233327.1, Chr2) is PGC-specific (Fig. 1a, b) (Supplementary Figure 1a, b). High-throughput screening results showed that the expression of C2EIP in PGC was significantly higher than that in ESC and SSC, in which there was little to no C2EIP expression (Fig. 1c). Western blotting and qRT-PCR showed that expression of C2EIP in PGC was higher than in ESC and SSC (Figs. 1d, e), a finding supported by in vitro retinoic acid (RA) induction experiments that determined C2EIP expression was upregulated 5.5-fold compared to non-RA-induced cells (Fig. 1d). The stop codon of C2EIP was mutated, and a C2EIP-EGFP-N1 fusion expression vector was constructed to confirm that C2EIP successfully encoded protein in DF-1 cells (Fig. 1f) (Supplementary Figure 1c). To further investigate the subcellular localization of C2EIP proteins, we prepared a polyclonal antibody. We constructed a pcDNA3.0-C2EIP vector and injected it into mice intra-peritoneally. The antibody titer was 1:10 (Supplementary Figure 1d). pcDNA3.0-C2EIP vector was also transfected into well-cultured DF-1 cells and indirect immunofluorescence experiments were performed using the polyclonal antibody. The results showed that C2EIP was indeed expressed in the cytoplasm (Fig. 1g), and similar results were also found in PGC (Supplementary Figure 1e). Our results suggested that C2EIP was a PGC-specific gene that could encode protein.

C2EIP is highly expressed in PGC via promoter and histone modification

Because we observed C2EIP more highly expressed in PGC than in ESC or SSC, we speculated that the activity of the C2EIP promoter changes during PGC generation, which leads to differential gene expression in PGC compared to ESC and SSC. Therefore, we identified the promoter region of the C2EIP gene based on the original promoter TATAbox and CAAATbox and cloned the C2EIP promoter fragment to replace the pEGFP-N1 promoter in a pcDNA3.0-EGFP recombinant vector. The constructed pcDNA3.0-EGFP vector (Fig. 2a) can express green fluorescent protein in DF-1 cells (Supplementary Figure 2a), confirming the C2EIP promoter activity.

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**Fig. 1** C2EIP encodes a cytoplasmic protein that is differentially expressed in PGC. 

a) Transcriptome analysis of ESC, PGC, and SSC. The differential expression of genes in these cells can be divided into six clusters. Cluster1 represents genes with high expression in ESC but low expression in PGC and SSC; Cluster2 represents genes with high expression in ESC and PGC but low expression in SSC; Cluster3 represents genes with high expression in PGC but low expression in ESC and SSC; Cluster4 represents genes with high expression in PGC and SSC but low expression in ESC; Cluster5 represents genes with high expression in SSC but low expression in ESC and PGC; Cluster6 represents genes with high expression in ESC and SSC but low expression in PGC.

b) The distribution of differentially expressed genes in ESC vs. PGC and PGC vs. SSC indicate C2EIP is highly expressed in PGC. Red represents upregulated genes, green represents down-regulated genes.

c) The RPKM value of C2EIP, representing relative expression, in ESC, PGC, and SSC.

d) qRT-PCR analysis of C2EIP expression in indicated tissues, ESC, PGC, SSC, and RA-induced cells.

e) Western blot analysis of C2EIP protein levels in ESC, PGC, and SSC. β-actin was used as an internal reference.

f) Confirmation of C2EIP-EGFP fusion protein expression. Cells were transfected with EGFP-N1 vector as a positive control, or mock-transfected with double distilled water (DDW) as a negative control.

g) Indirect immunofluorescence showing cytoplasmic localization of C2EIP. Cells transfected with pcDNA3.0 vector were used as a negative control; Scale bar:20μm
The −264~-~-94 bp region was identified as the core region of the \textit{C2EIP} promoter (Fig. 2e) based on the luciferase activity of truncated fragments (Fig. 2a) (Supplementary Figure 2c), and this region includes binding sites for transcription factors such as STAT1, STAT10, Sox17, Sox2, and Klf5 (Fig. 2f). These transcription factors are involved in stem cell maintenance and differentiation. In this study, homologous recombination and seamless cloning were performed on five sites to make point mutations, and the promoter activity of the \textit{C2EIP} gene was tested after mutation at each transcription factor binding site. The results showed that increased STAT1 expression can increase the activation of normal PGL3.0/170 (142.54 ± 2.71), while STAT1 binding site mutation can significantly reduce the PGL3.0/170 promoter activity (68.34 ± 2.62) (Fig. 2g). The results for STAT10, Sox17, Sox2, and Klf5 and their binding sites were the opposite (Supplementary Figure 2d-f). Therefore, the transcription factor STAT1 appears to be a positive regulator of the \textit{C2EIP} promoter, whereas STAT10, Sox17, Sox2, and Klf5 are negative regulators. Together, our results indicate that STAT1 binding, histone acetylation, and promoter methylation regulate \textit{C2EIP} expression.

\textit{C2EIP} promotes PGC generation in vitro and in vivo

To study the role of \textit{C2EIP} in PGC generation, we implemented a model for inducing ESC differentiation into PGC by in vitro RA treatment (Fig. 3a) (Supplementary Figure 3a-f)\(^{20}\), in which we overexpressed \textit{C2EIP} or generated a \textit{C2EIP} knockout via CRISPR/Cas9.
These experiments showed that C2EIP overexpression promoted formation of PGC-like cells (Fig. 3d) (Supplementary Figure 3a, b). Expression levels of C-KIT and CVH, which encode gene products involved in PGC development, were markedly increased in the overexpression group (+2.3-fold; \( p < 0.01 \)), whereas their expression in the knockout group decreased 1.1-fold \( (p < 0.01) \) compared to controls (Fig. 3e). Flow cytometry analysis showed fewer CVH+ cells in the C2EIP knockout group (4.8 ± 0.16%; \( p < 0.01 \)) but increased numbers of CVH+ cells in the overexpression group (18.6 ± 0.13%; \( p < 0.01 \)) (Fig. 3f). Our results suggest that in vitro C2EIP overexpression positively regulates PGC generation.

To further investigate the function of C2EIP during in vivo PGC differentiation, we injected the overexpression knockout vector, mixed isovolumetrically with polyethylenimine (PEI), into chick embryos (Supplementary Figure 4). The body weight of 4.5-day-old chick embryos in the knockout group was significantly lower than that in the control and overexpression groups\(^{14} \), suggesting that C2EIP knockout negatively affects chick embryo development. qRT-PCR and Western blot results showed that expression of CVH in the knockout group was reduced compared to the control group (0.31 ± 0.04; \( p < 0.01 \)) (Figs. 4a, b), while CVH expression in the C2EIP overexpression group increased
compared to the C2EIP KO group (1.03 ± 0.34; p < 0.01) (Figs. 4a, b). Moreover, genital glands were observed in control and overexpression group embryos, while no complete genital glands were seen in embryos from the knockout group (Figs. 4c, d).

**Low C2EIP expression maintains ESC pluripotency and inhibits PGC generation**

We preliminarily analyzed the mechanism underlying C2EIP-mediated PGC generation by transfecting ESC with either the C2EIP overexpression construct or the CRISPR/Cas9-mediated knockout vector in in vitro RA induction assays. Overexpression of C2EIP down-regulated pluripotency-related genes Oct4 and Sox2 (Fig. 5a-d), reduced AKP activity (Fig. 5a), and reduced Nanog promoter-driven EGFP expression (1.82 ± 0.06; p < 0.01) (Fig. 5e), consistent with the increased PGC generation (Fig. 3e). In contrast, C2EIP knockout led to significant increases in Oct4 and Sox2 levels, AKP activity, and Nanog promoter-driven EGFP expression (2.41 ± 0.04; p < 0.01), respectively (Fig. 5e). With the effects of the overexpression and knockout constructs on expression levels of Oct4 and Sox2 in the genital ridge of 4.5-day-old chick embryos (data not shown) were similar to those in the in vitro model experiment. Thus, expression of C2EIP appears to limit ESC pluripotency and promote their differentiation into PGC.

**C2EIP and PTCH2 are co-expressed at the intracellular membrane**

We investigated the molecular mechanism underlying C2EIP-mediated PGC generation by characterizing the C2EIP-PTCH2 interaction with GST pull-down assays and mass spectrum analysis (Supplementary Table 1) (Supplementary Figure 5a-d). Co-immunoprecipitation (co-IP) experiments confirmed the protein-protein interaction (Fig. 6a). Our indirect immunofluorescence...
assays found that PTCH2 colocalized with known membrane protein SSEA-1, indicating that PTCH2 is a membrane protein, whereas C2EIP is primarily localized in the cytoplasm (Figs. 6b, c). C2EIP and PTCH2 form a complex at the intracellular membrane and likely function collectively at that location (Fig. 6c). Western blotting detected C2EIP in the GST-PTCH2 eluate from PGC lysate, suggesting that C2EIP and PTCH2 colocalize in the intracellular membrane of PGC (Figs. 6b-e).

**C2EIP regulates PGC generation by activating the HH signaling pathway**

HH signaling is a classical pathway associated with stem cell differentiation that relies on PTCH2 as a membrane surface receptor for regulation. When induction molecules are absent, PTCH2 binds to SMO and inhibits HH signaling. In the presence of signaling molecules, C2EIP can bind preferentially, but randomly, to PTCH2, thereby excluding SMO and activating HH signaling.
Here, we found that C2EIP interacts with PTCH2 at the intracellular membrane and thus speculated that C2EIP may regulate PGC differentiation by influencing HH signaling pathway activation. Therefore, we constructed lentiviral shRNA-IHH and shRNA-PTCH2 vectors to investigate the function of HH signaling in the process of PGC generation (Supplementary Figure 6a-c). ESC were transfected with shRNA-IHH or shRNA-PTCH2 in parallel with in vitro RA induction experiments. When ESC were routinely induced by RA, many PGC-like cells emerged 4–6 days after induction; numbers of PGC-like cells did not differ significantly between the IHH and PTCH2 knockdown groups. After IHH knockdown, PGC generation declined, and expression levels of C-KIT and CVH were significantly down-regulated compared to cells in the RA induction group. In contrast, PGC generation and expression of both C-KIT and CVH increased after PTCH2 knockdown (Supplementary Figure 6d-f), which was consistent with the results of our in vivo experiment (Supplementary Figure 7a, b). Thus, PGC generation appears to depend on C2EIP-PTCH2 interactions that lead to activation of the HH signaling pathway.

C2EIP regulates PGC generation by promoting ubiquitination and dephosphorylation of PTCH2

We previously confirmed that HH signaling participates in ESC differentiation into PGC. To explore the specific mechanism underlying the relationship between C2EIP-mediated activation of PTCH2 and HH signaling, we measured changes in epigenetic modification of PTCH2 during PGC generation. During the differentiation process, mRNA expression levels of C2EIP, PTCH2, and SMO increased, and the HH signaling pathway was activated, although transcription levels of C2EIP, PTCH2, and SMO were not consistent with their translational levels. Thus, PTCH2 protein might be post translationally modified, leading to reduced activity. Our ubiquitination and dephosphorylation experiments showed decreased dephosphorylation and increased ubiquitination of PTCH2 during ESC differentiation into PGC (Fig. 7a). We observed similar results in the in vitro RA induction experiments (Figs. 7c, d), leading us to conclude that the HH signaling pathway is activated following C2EIP-mediated modification of PTCH2 during PGC generation.

To further define the correlation between C2EIP and the ubiquitination and dephosphorylation of PTCH2, we overexpressed and knocked out C2EIP in PGC and examined the posttranslational modification of PTCH2. When C2EIP was overexpressed, PTCH2 phosphorylation was reduced (data not shown) and ubiquitination increased (Fig. 7e), and phosphorylation of SMO protein was upregulated (data not shown). When C2EIP was knocked out, constitutive phosphorylation of PTCH2 was restored (Fig. 7b) but de-ubiquitination occurred (Fig. 7e), and phosphorylation of SMO decreased (data not shown). Therefore, C2EIP might regulate PGC generation by...
promoting PTCH2 ubiquitination, excluding PTCH2-SMO interactions, and subsequently activating the HH signaling pathway.

**Discussion**

We determined that C2EIP is a genetic marker expressed specifically in PGC and can thus complement the use of CVH as a differentiation indicator, because CVH influences migration, aggregation, and localization of PGC but is not specifically expressed in these cells. C2EIP also plays an important role in the late stage of germ cell generation. Like CVH, C-KIT also contributes to PGC development but is widely expressed in germ cell lines. We utilized high-throughput sequencing and bioinformatics analysis to identify multiple PGC-specific markers, including Blimp1, Prdm14, and STELLA, but focused on the functional role of C2EIP in PGC generation and found that its gene product inhibits ESC pluripotency and directs their differentiation into PGC.

The process of ESC-to-PGC differentiation is regulated by both genetic and cell-signaling mechanisms. For example, BMP427,28 and NANOS229,30 can regulate the proliferation and differentiation of PGC by activating the TGF-β/Smad and RA signaling pathways, respectively. The HH signaling pathway, a classical stem cell regulation pathway, regulates cell differentiation by modulating downstream signals through epigenetic modifications. Specifically, IHH binds to PTCH, which leads to SMO release and activation of the HH signal. The PTCH membrane protein is composed of PTCH1 and PTCH2, which share similar ligand binding affinities but often display different expression patterns. PTCH1 functions mainly in somatic cells, while PTCH2 functions mainly in embryo development14. If both subunits are highly expressed, HH signaling is inhibited, but lack of expression leads to HH signal activation31. During male germ cell differentiation in chick embryos, we observed upregulation of PTCH1 and downregulation of PTCH2, thus highlighting the specific importance of PTCH2 in HH signal activation in ESC. Based on its functional characteristics, we questioned if C2EIP is an HH gene family homolog, although subsequent bioinformatic analysis revealed no conserved structural or functional domains between C2EIP and the HH gene family and identified different localization patterns. Thus, we began to explore the interaction between PTCH2 and C2EIP and its effect on HH signaling activation. During ESC differentiation, we determined that C2EIP induces the ubiquitination of PTCH2 protein and activates SMO and HH signaling, which together promote PGC generation.

Epigenetic modification is a common mechanism for signaling pathway regulation32. For example, ubiquitination and dephosphorylation of Smad proteins modulate TGF-β/BMP signals, while the HH signaling pathway enables phosphorylation of Smad5 and induces signals for downstream cell differentiation33,34. We demonstrated that C2EIP specifically activates the HH signal in ESC and promotes their differentiation into PGC. Expression of C2EIP and PTCH2 in ESC is very low initially but increases as numbers of PGC increase, thus perpetuating
the HH signal. Our co-IP and in vitro ubiquitination experiments revealed that during the PGC differentiation, phosphorylation of PTCH2 was constantly reduced, while its ubiquitination constantly increased, indicating activation of HH signaling pathway. Thus, these two types of epigenetic modification can reduce accumulated intracellular PTCH2, release SMO protein, transduce the HH signal, promote ESC differentiation, and positively regulate PGC generation.

C2EIP is differentially expressed in various cell types, and its promoter plays very important roles in expression modulation. Although C2EIP does not include a binding element of RNA polymerase II or a CpG island, the gene is regulated by promoter methylation, histone acetylation, and binding of the STAT1 transcriptional factor in the core area to enhance gene expression. Previous research determined that cell differentiation is negatively affected in STAT1-deficient mice and that cell differentiation of human promyelocytes is related to phosphorylation of STAT1. When we mutated STAT1, activity of the C2EIP promoter was significantly reduced but could be restored in ESC by RA induction in vitro.

In summary, we identified a PGC-specific marker gene in chick embryos and determined its role in a novel molecular mechanism responsible for directed ESC differentiation. At low methylation levels, STAT1 promotes expression of C2EIP, whose gene product binds to PTCH2, induces its ubiquitination, activates the HH signaling pathway, and positively regulates PGC generation.

Materials and methods

Cell culture and induction
ESC and PGC were cultured under previously described conditions. Well-grown second-generation ESC cells were used in the cell induction experiments. Induction medium was DMEM supplemented with 10 mM RA and 15% FBS. To investigate the effects of C2EIP gene expression on PGC generation in vitro, ESC were transfected with C2EIP overexpression and knockout vectors during induction.

Antibodies and reagents
Anti-C2EIP mouse antibody (previously used in) was used to detect the expression of C2EIP protein in different tissues. In the flow cytometry and immunohistochemical assays, antibodies against CVH (Abcam, San Francisco, USA; ab27591) and C-KIT (Abcam, San Francisco, USA; ab27591) were used to label PGC. Antibodies against GST (Abcam, San Francisco, USA; ab19256) and C2EIP were used to test the accuracy of the GST pull-down experiment. PTCH2 antibody (Abcam, San Francisco, USA; ab194574) was used to examine protein cellular localization. Antibodies against PTCH2 (Abcam, San Francisco, USA; ab151775) were used for the co-IP experiment. Ubiquitylation antibody (Abcam, San Francisco, USA; ab139467) was used to determine the ubiquitylation level of PTCH2 under different treatments. TSA (MCE, Monmouth, USA; HY-15144) and 5-Zacd (MCE, Monmouth, USA; HY-10586) were used to inhibit deacetylation and methylation, respectively. The dual fluorescein test kit (Solarbio, Beijing, China; D0010) was used to measure the activity of promoter fragments.

Fluorescence-activated cell sorting
The direct labeling method was used to label the cells. The 4–6 d cells were harvested by transferring suspension to 1.5-mL centrifugal tubes and centrifuging at 1000×g for 5 min, then rinsed with PBS 1–2 times and incubated at 4 °C for 1–2 h with 200 µL of 1:200 fluorescein-labeled antibody (CVH or C-KIT). Cells were rinsed with chilled PBS to remove the unbound antibody. Finally, cells were resuspended in 500 µL of chilled PBS. The resultant sample was subjected to flow cytometry analysis. Cells transfected by EGFP fluorescent plasmids were counted and 105 cells were subjected to flow cytometry analysis. For chick embryo cells, the genital ridge was removed after 4.5d incubation (n = 30 per group). Single-cell suspension was obtained by trypsin treatment and the cells were counted. Using CVH and C-KIT antibodies, 105 cells were labeled. The cells were then subjected to flow cytometry analysis.

Construction of overexpression and knockout vectors
Based on the CDS of C2EIP in the NCBI database, we designed specific primers to clone the full-length CDS and ligated it into the pCDNA3.0 vector (Supplementary Tables 2, 3). To construct the C2EIP gene knockout vector, we selected three specific targets at the APMLocus and ligated them into the VK100081 vector based on the sequence of each exon of C2EIP in the UCSC database. The knockout efficiency of three knockout vectors was tested by T7EI, SSA, and TA cloning and sequencing.

Chicken embryo injection and slice preparation
Plasmid was mixed with PEI at the equal volumes, and 100 µl of the mixture (12 µg) was injected into the chicken embryo via the top end. At 4.5 d after injection, frozen and paraffin-embedded slices were prepared. For frozen slices, the embryo was mixed with cryoprotectant and frozen in liquid nitrogen for 15 s. The frozen embryo was warmed up to −20 °C and then 0.7-µm slices were prepared. The slice was sealed with cover glass and real-time fluorescence was observed under an inverted microscope to observe protein expression. To prepare paraffin-embedded slices, embryos were fixed overnight by paraformaldehyde. After dehydration by alcohol, paraffin slices were prepared. The prepared slice was dewaxed, dehydrated, and...
used for immunohistochemical assays with CVH and C-KIT antibodies and PAS staining.

Construction of C2EIP promoter deletion fragments

Based on bioinformatics analysis of the C1EIP promoter in Rugao yellow chickens, we fixed the downstream primer, changed the upstream primer, and designed five pairs of primers to amplify different deletion fragments. We constructed PGL3 and N1-EGFP vector, cleaved the vectors and PCR products with dual enzymes, recovered the cleaved products, and ligated. After confirming enzymes by enzymatic cleavage and sequencing, these deletion fragments were named PGL3-Basic-P1 to PGL3-Basic-P4, and EGFP-N1-P0 (Supplementary Tables 4, 5).

Promoter activity assays

DF1 cells were transfected with promoter fragment plasmids using Lipofectamine TM 3000 transfection reagent. Control group cells were transfected by pEGFP-N1 plasmid, and blank control cells were not transfected. Well-grown DF1 cells were grown in 24-well plates at a starting density of 2 × 10^5 cells. The dual luciferase expression vectors or PGL3-Basic were mixed with PRL-Nl plasmid, and blank control cells were not transfected. Well-grown DF1 cells were grown in 24-well plates at a starting density of 2 × 10^5 cells. The dual luciferase expression vectors or PGL3-Basic were mixed with PRL-EV at a ratio of 1:30 and used to co-transfect the DF1 cells. After 24–48 h of co-transfection, the cells were harvested and the activity of each deletion fragment of the C2EIP promoter was tested by the dual luciferase reporter gene detection system. TSA or 5-Zac was added to the culture solution to test the effects of deacetylation or methylation, respectively, on promoter activity.

GSTPull-down and mass spectrum (MS) analysis

Specific primers were designed by Primer 5.0 software to clone the full-length CDS of C2EIP and introduce EcoRI and HindIII cleavage sites. After cleavage and ligation into the Pet-49(b)-GST plasmid, the experimental plasmids were transformed into the Escherichia coli BL21 expression strain. Expression was induced by IPTG at 18 °C, and the protein was isolated from cells, purified from the precipitate, concentrated, and used in GST pull-down experiments to identify interactions. To prepare samples for MS analysis, protein was alkylated and enzymatically cleaved after GST pull-down, and the sequence of interacting proteins was identified by LC-MS/MS.

Indirect immunofluorescence assay

Well-grown PGC after 6d induction were fixed by paraformaldehyde for 30 min, rinsed with PBS three times, permeabilized in 0.5% TritonX-100 for 10 min, and rinsed with PBS three times again. After blocking in PBS containing 10% BSA at room temperature for 30 min, PGC were incubated with C2EIP, PTCH2, C-KIT, or CVH antibodies overnight at 4 °C. These cells were then rinsed with PBST three times and incubated in the dark with secondary antibody at 37 °C for 1 h, rinsed by PBST three times again, and sealed by incubation in glycerol. Protein expression was observed under a fluorescence microscope.

Western blot analysis

ESC, PGC, or SSC from chicken embryo or 6-d-induced cells were collected and lysed in RIPA buffer. A total of 20 μg of protein in 5 μL loading buffer was boiled for 3–5 min to denature proteins and pipetted into each lane. After separation by 10% SDS-PAGE, the protein was transferred to a nitrocellulose membrane by the semi-dry method. The resulting membrane was blocked by TBST containing 5% fetal bovine serum at room temperature for 1 h. Primary antibodies against Smad2, Smad5, and Notch1 were added and the membrane was incubated overnight at 4 °C. The membrane was then incubated with secondary goat anti-rabbit antibody at 37 °C. The protein was visualized colorimetrically by DAB.

Co-IP test

The genital ridge of 4.5d chicken embryos was isolated under sterile conditions. The single-cell suspension was produced by trypsin treatment and labeled with the CVH and C-KIT antibodies. High-purity PGC were isolated by flow cytometry. The resulting cells were lysed in RIPA buffer and centrifuged at 13000 g at 4 °C for 15 min. The protein was stored at −70 °C. For co-IP, the extracted protein was incubated overnight with an optimal concentration of anti-PTCH2 antibody. The protein was then centrifuged at 1000×g at 4 °C for 10 s.

Data analysis

The promoter activity test, qRT-PCR, and FACS were replicated three times. The results were analyzed by t-test using SPSS19.0 software package. p < 0.05 was considered significant, and p < 0.01 highly significant. Charts were prepared in GraphPad Prism 6.

Accession Numbers

The SRA accession numbers for the RNA-seq data reported in this paper are: SRR3720923, SRR3720924, and SRR3720925.

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References
1. Matsui, Y., Ziebo, K. & Hogan, B. L. Derivation of pluripotential embryonic stem cells from murine primordial germ cells in culture. Cell 70, 841–847 (1992).
2. Hayashi, K. & Saitou, M. Generation of functional primordial germ cells from pluripotent stem cells. J. Marum. Dev. Res. 29, 2–10 (2012).
3. Resnick, J. L., Bixler, L. S., Cheng, L. & Donovan, P. J. Long-term proliferation of mouse primordial germ cells in culture. Nature 359, 550–551 (1992).
4. Godin, I. et al. Effects of the steel gene product on mouse primordial germ cells in culture. Nature 352, 807–809 (1991).
5. Lopez-Gonzales, P. et al. Hypoxia induces pluripotency in primordial germ cells by hif1α stabilization and oct4 deregulation. Antioxid. Redox Sign. 22, 205–223 (2015).
6. Liu, Y. et al. Hif-1alpha and hif-2alpha are critically involved in hypoxia-induced lipid accumulation in hepatocytes through reducing pgc-1alpha-mediated fatty acid beta-oxidation. Toxicol. Lett. 2, 116–123 (2014).
7. Nakamura, T. & Extavour, C. G. The transcriptional repressor blimp-1 acts downstream of bmp signaling to generate primordial germ cells in the cricket grilinus bimaculatus. Development 143, 255–263 (2016).
8. Vincent, S. D. et al. The zinc finger transcriptional repressor blimp1/prdm1 is dispensable for early axis formation but is required for specification of primordial germ cells in the mouse. Development 132, 1315–1325 (2005).
9. Yamaji, M. et al. Critical function of prdm14 for the establishment of the germ cell lineage in mice. Nat. Genet. 40, 1016–1622 (2008).
10. Kurimoto, K., Yamaji, M., Seki, Y. & Saitou, M. Specification of the germ cell lineage in mice: a process orchestrated by the pr-domain proteins, blimp1 and prdm14. Cell Cycle 7, 3514–3518 (2008).
11. Tong, M. H., Mitchell, D., Evanoff, R. & Griswold, M. D. Expression of miel7 family members in response to retinoic acid-induced spermatogonial differentiation in mice. Biol. Reprod. 85, 189–197 (2011).
12. Childs, A. J., Kinnell, H. L., He, J. & Anderson, R. A. Lin28 is selectively expressed by primordial and pre-meiotic germ cells in the human fetal ovary. Stem Cells Dev. 21, 2393–2394 (2012).
13. Zhang, Z. et al. Crucial genes and pathways in chicken germ stem cell differentiation. J. Biol. Chem. 290, 13605–13621 (2015).
14. Zuo, Q. et al. Site-directed genome knockout in chicken cell line and embryos can use crispr/cas gene editing technology. G3 Genom. Genet. 6, 1797–1792 (2016).
15. Chen, H. et al. Regulation of hedgehog signaling in chicken embryonic stem cells differentiation into male germ cells (gallus). J. Cell Biochem. 118, 1379–1386 (2016).
16. Lawson, K. A. & Hage, W. J. Clonal analysis of the origin of primordial germ cells in the mouse. Ciba. Found. Symp. 182, 68–84 (1994).