ESRG is critical to maintain the cell survival and self-renewal/pluripotency of hPSCs by collaborating with MCM2 to suppress p53 pathway

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Abstract

The mechanisms of self-renewal and pluripotency maintenance of human pluripotent stem cells (hPSCs) have not been fully elucidated, especially for the role of those poorly characterized long noncoding RNAs (lncRNAs). ESRG is a lncRNA highly expressed in hPSCs, and its functional roles are being extensively explored in the field. Here, we identified that the transcription of ESRG can be directly regulated by OCT4, a key self-renewal factor in hPSCs. Knockdown of ESRG induces hPSC differentiation, cell cycle arrest, and apoptosis. ESRG binds to MCM2, a replication-licensing factor, to sustain its steady-state level and nuclear location, safeguarding error-free DNA replication. Further study showed that ESRG knockdown leads to MCM2 abnormalities, resulting in DNA damage and activation of the p53 pathway, ultimately impairs hPSC self-renewal and pluripotency, and induces cell apoptosis. In summary, our study suggests that ESRG, as a novel target of OCT4, plays an essential role in maintaining the cell survival and self-renewal/pluripotency of hPSCs in collaboration with MCM2 to suppress p53 signaling. These findings provide critical insights into the mechanisms underlying the maintenance of self-renewal and pluripotency in hPSCs by lncRNAs.

Key words: ESRG, human pluripotent stem cells, pluripotency, cell survival, OCT4, MCM2, p53

Introduction

Human pluripotent stem cells (hPSCs), including human embryonic stem cells (hESCs) and induced pluripotent stem cells (hiPSCs), are pluripotent cells with self-renewal capability [1, 2]. In-depth study of their self-renewal and differentiation mechanisms is of great significance for understanding the mechanism of human embryonic development, directional induction differentiation and tissue engineering. Their pluripotent state and self-renewal ability are mainly maintained by transcription factors, signaling pathways and epigenetic modifications [3-5]. Among them, OCT4, NANOG and SOX2 have
been identified as key factors of the transcriptional regulatory network in hPSCs [6-9]. These core factors contribute to the hallmark characteristics of hPSCs by activating target genes that encode pluripotency and self-renewal associated proteins and repressing signaling pathways that promote differentiation [10]. In hPSCs, OCT4, NANOG and SOX2 co-occupy the promoters of hundreds of protein-encoding genes or long noncoding RNAs (lncRNAs) that are either expressed or repressed in the pluripotent state [11, 12]. This pattern suggests that there should be a complex regulatory circuit in which OCT4, NANOG and SOX2 collectively and uniquely regulate downstream targets to control PSC differentiation. However, the downstream effectors of these targets are largely unknown. Additionally, the mechanism by which these “master regulators” of pluripotency control the self-renewal and lineage-specific differentiation of hPSCs also remains to be elucidated.

ESRG (also known as HESRG), an hPSC-related lncRNA that was first cloned from an expressed sequence tag (EST, BF223023) in our previous study, is highly expressed in hPSCs and is significantly decreased or even completely silenced after differentiation [13]. Our previous analysis indicated that ESGR is located on human chromosome 3p14.3, which contains four exons and three introns with a full length of 3153 nucleotides. Homologous sequences could only be found in primates, such as Pan troglodytes and Macaca mulatta, but not in other species, implying that this gene may be a very specific gene only existing in primate cells [13]. No ESRG expression was detected in most human cell lines and tissues except in hESCs and several kinds of malignancies, including intracranial germinoma and embryonal carcinoma cells, colorectal cancer cells, and high-grade ovarian serous tumor cells [14-16]. In differentiated cells from RA-treated hPSCs and adult tissues, the expression of ESRG was tenuous or undetectable. As OCT4 regulates target gene expression by preferentially binding to an octamer sequence (ATTTCGAT), the presence of this octamer at -1,714 to -1,707 bp in the upstream region of the transcription start site of ESRG implies that ESRG may be directly regulated by OCT4 [13].

Recently, ESRG has been reported to be useful as a marker to identify early reprogramming cells [17] and as an excellent marker of undifferentiated hPSCs [18]. Wang et al. also [19] reported that ESRG is required for human-specific pluripotency. These studies suggested that ESRG is required for the maintenance of hPSC identity. However, a recent study controverted that ESRG is dispensable for human pluripotency [20].

In the current study, we investigated the roles and mechanisms of the regulation and action of ESRG in hPSCs and showed that ESRG is transcriptionally regulated by OCT4. ESRG is involved in the self-renewal and pluripotency maintenance of hPSCs by binding to minichromosome maintenance 2 (MCM2) to sustain its nuclear location and steady-state levels, safeguarding error-free DNA replication. Knockdown of ESRG results in DNA damage and activation of the p53 signaling pathway, subsequently promoting hPSC differentiation, cell cycle arrest, and apoptosis. Additionally, our results showed that hPSCs were extremely sensitive to p53 when ESRG was knocked down, and once the cells tolerated it, the cells would no longer be affected by p53. These data demonstrated that ESRG is an indispensable element in the maintenance of cell survival and self-renewal/pluripotency of hPSCs.

Materials and methods

Culture of hPSCs in knockout serum replacement (KSR)-based hPSC medium

Undifferentiated hPSCs [H9 and H1-hESCs (Madison, WI); RC1 and RC2-iPSCs (RC01001, Nuwacell)] were cultured on mitotically inactivated feeder cells (primary mouse embryonic fibroblasts, MEFs) in Dulbecco’s Modified Eagle Medium (DMEM/F12 [GIBCO] supplemented with 20% serum replacement (Knockout SR) [GIBCO], 1% MEM Non-Essential Amino Acids [GIBCO], 1% GlutaMAX-I [GIBCO], 4 ng/mL basic fibroblast growth factor [bFGF] [Peprotech] and 0.1 mM β-mercaptoethanol [Sigma]) [21]. The culture medium was refreshed every day, and the cells were conventionally passaged once a week. Cell colonies were loosened from the culture substrate with 1 mg/mL Collagenase Type IV solution (GIBCO, dissolved in DMEM/F12) and gently divided into small pieces with cell scrapers (Becton Dickinson). The hPSC clusters were passaged to a new feeder layer or growth factor reduced Matrigel-(Becton Dickinson)-coated plates at a splitting ratio between 1:6 and 1:10. The presence of the feeder layer complicates mRNA or protein level analysis, so the hPSCs were passaged to Matrigel-coated plates under feeder-free conditions in MEM-conditioned medium (CM) supplemented with 10 mM Y-27632 (Millipore) to maintain pluripotency, as described previously [22]. hPSCs cultured on Matrigel-coated plates were passaged as single cells using Accutase solution (Millipore). Cell morphology was recorded using an inverted fluorescence microscope (TE2000U, Nikon).
Culture of hPSCs in Essential 8 (E8) hPSC medium

hPSCs (H9, H1 RC1 and RC2-iPSCs) were cultured in Essential 8 (E8) medium on plates coated with Matrigel [23]. In brief, Matrigel was diluted in DMEM/F12, incubated in dishes for 1 h at room temperature, and then aspirated before replating cells. The hPSCs were passaged every 2-3 days using EDTA dissociation buffer. In brief, the cells were washed twice with DPBS and then incubated with EDTA dissociation buffer for 1-2 min at 37 °C. Then, the EDTA was aspirated and the cells were collected in E8 medium by gentle pipetting and replated at a 1:8-1:12 ratio.

Culture of other cell lines in DMEM medium

Other cell lines, including 293T cells, MEFs, HeLa cells and COS-7 cells, were grown in DMEM containing 10% FBS (GIBCO) and 1% MEM non-essential amino acids.

Design and screening of double-stranded siRNAs targeting ESRG

Double-stranded siRNAs targeting ESRG (siESRG) were designed by different companies or designed using online tools. The siRNAs targeting the ESRG RNA sequence could not share significant homology with other genes or sequences in the genome; therefore, homology analysis (www.ncbi.nlm.nih.gov/BLAST/) was performed to avoid off-target effects on other genes or sequences. Finally, 25 sets of siESRG sequences were selected and synthesized by RiboBio Co. (RiboBio). Each designed siESRG sequence corresponded to a different region in the target ESRG RNA sequence. To screen for the best-designed siRNA, we transfected the siESRG sequences into hESCs using Lipofectamine RNAiMAX (Invitrogen) [24].

Lentiviral vector construction, viral production and viral infection

ESRG siRNA (siESRG1, siESRG2 and siESRG3) and negative control siRNA (siN0000001, RiboBio) were transformed into shRNA sequences and cloned into the pGV118 vector (GeneChem). The sequences of the effective shRNAs were as follows: siESRG 1: 5’-GGATGTGGTCCGAGTGTGGT-, siESRG 2: 5’-GGATGGAGCCATAGAAGTT-, siESRG 3: 5’-CCATTAAAGGGAAGACATA-. The constructed vectors (pGV118-shESRG1, pGV118-shESRG2, pGV118-shESRG3 and pGV118-shControl) were transfected together with the lentivirus packaging vectors pMDLg/pRRE, pRSV-REV and pCMV-VSV-G (Addgene) into 293T cells using Lipofectamine 2000 (Invitrogen) transfection reagent according to the manufacturer’s instructions. The viral supernatant was harvested 48 h after transfection, centrifuged (10 min, 4000 g), filtered using a 0.45 µm Millex-GP filter (Millipore), and then concentrated using a Millipore Centricron-Plus-20 centrifuge filter unit (Millipore). A multiplicity of infection (MOI) of 50 was used to infect hESCs with the lentiviruses [25, 26]. Cells were dissociated with Accutase solution and seeded in CM supplemented with 10 mM Y-27632 on Matrigel-coated six-well plates, together with the generated lentiviruses and 8 µg/mL of polybrene (Sigma) for viral infection. To confirm the knockdown efficiency of ESRG, cells were harvested for qPCR analysis 5 days after viral infection.

Establishment of a tetracycline- (Tet-) inducible shRNA system

To establish a stable hESC line that constitutively expressed the TetR protein, H9 hESCs were transfected with the pCAG-TetRnls vector (a kind gift from Peter W. Andrews) using FuGENE HD Transfection Reagent (Promega) and then selected using 2 µg/mL puromycin (Sigma) 48 h after transfection. Cell clones (H9-TetR cells) that highly expressed TetR protein were selected, expanded, and used in subsequent studies.

According to the pSUPER RNAi system user manual (Oligoengine), OCT4 siRNA with the target sequence GGATGTGGTCCGAGTGTGGT, ESRG siRNA with the target sequence GGATGGAGCCATAGAAGTT and β2 M siRNA (negative control) with the target sequence GGACTGGTCTTTCTATC were designed as shRNAs using Oligoengine 2.0 and then cloned into the pSUPERregor. Puro vector (Oligoengine) [27]. Subsequently, H9-TetR cells were transfected with the constructed vectors pSUPERior-shOCT4, pSUPERior-shESRG or pSUPERior-shβ2 M using FuGENE HD and selected with 300 µg/mL G418 (Sigma). Once single, stable transfectants (H9-TetR-shOCT4/H9-TetR-shESRG cells) were generated, they were isolated, cultured on Matrigel-coated plates in CM, and assessed by qPCR for the efficiency of gene knockdown following treatment with doxycycline (Dox, Sigma).

Adenovirus vector construction and transfection

ESRG expression adenovirus and control adenovirus were synthesized by Vigenbio (Shandong, China). According to the actual situation and MOI value of the experiment, the adenovirus stock solution was diluted in the culture medium and then added to the target cells. After being mixed, the cells were incubated overnight in an incubator at 37 °C and 5% CO2. After 12 h of virus infection, the culture medium
was changed and tested 48 h later.

**Construction of the ESRG Pr-tdTomato reporter hPSC line**

We constructed an *ESRG* Pr-tdTomato reporter hPSC line using gene editing techniques to insert fusion sequences of the *ESRG* promoter and tdTomato fluorescent protein gene into the AAVS1 human safe harbor locus as previously described [28].

**Construction of truncated, antisense vector and domain vector**

The truncation and antisense sequences of *ESRG* were cloned into the pcDNA3.1(+) vector. The Flag-MCM2 domain (MCM2 full-length, MCM2-N, MCM2-C, MCM2-N1, MCM2-N2, and MCM2-N3) expression plasmids were cloned into the PB212 (1 BsiWI)-Puro vector.

**RNA isolation and qPCR**

Total cellular RNA was isolated with TRIzol reagent (Invitrogen) as described previously [29], followed by treatment with deoxyribonuclease I (DNase I, Promega) to remove any contaminating genomic DNA. First-strand cDNA was synthesized using 1 μg of total RNA and a RevertAid First Strand cDNA Synthesis Kit (Fermentas). qPCR analysis was then performed using FastStart Essential DNA Green Master Mix (Roche) and an MJ Mini Personal Thermal Cycler (Bio-Rad). The threshold cycle (Ct) values of each sample were used in the post-PCR data analysis and normalized against endogenous *GAPDH*. qPCR was conducted in triplicate for each sample. The primer sequences for the qPCR are listed in Table S3.

**Isolation of RNA and protein from the nucleus and cytoplasm**

The extraction of RNA and protein was performed according to the PARIS Kit (Invitrogen, AM1921). Briefly, 10^7 cells were collected and disrupted in 100-500 μL ice-cold Cell Fractionation Buffer and centrifuged for 5 min at 4 °C and 500 g after being incubated on ice for 5-10 min. Then, the cytoplasmic fraction was carefully aspirated away from the nuclear pellet, and the nuclear pellet was lysed in Cell Disruption Buffer. Finally, the lysate was divided for RNA isolation and protein analysis.

**AP staining**

AP staining was performed using a SIGMAFAST BCIP/NBT tablet (Sigma) following the manufacturer’s instructions. Briefly, cells were washed twice with PBS and stained with SIGMAFAST BCIP/NBT substrate solution in the dark for 5-10 min. Then, the substrate solution was removed when sufficient color had developed [22]. After staining, the cells were washed with PBS, and bright-field images were taken using an inverted fluorescence microscope (TE2000U, Nikon).

**Western blot analysis**

Western blot analysis was performed as previously described [30, 31]. Briefly, cells were washed twice with PBS and lysed in M-PER Mammalian Protein Extraction Reagent (Thermo) containing 1×Halt Protease Inhibitor Cocktail (Thermo) and PhosSTOP phosphatase inhibitors (Roche). Equal amounts (20-40 μg) of total protein were boiled, electrophoresed on 12% SDS-polyacrylamide gels and electroblotted onto PVDF membranes (Millipore). Membranes were blocked for 1 h with 5% fat-free milk solution or 5% BSA, probed with primary antibodies overnight at 4 °C, and then incubated with secondary antibodies at room temperature for 2 h. The blot was visualized using enhanced chemiluminescence (ECL, Thermo). Images were recorded using the Molecular Imager ChemiDoc XRS+ System (Bio-Rad) and analyzed with Quantity One 1-D analysis software (Bio-Rad). The antibodies used in this study are listed in Table S4.

**Cell cycle analysis**

hPSCs were carefully dissociated into single-cell suspensions using Accutase solution, washed twice with PBS, and then fixed overnight with cold 70% ethanol. The fixed cells were washed twice with PBS, followed by RNase (Sigma, 100 μg/mL) treatment and propidium iodide (Sigma, 50 μg/mL) staining for 30 min at 37 °C. Approximately 1×10^6 cells were analyzed using a FACSCanto II (Becton Dickinson) to determine the cell cycle distribution pattern. The percentages of cells in G1, S and G2/M phases of the cell cycle were analyzed using ModFit 4.0 software (Verity Software House).

**Cell proliferation assays**

Cell proliferation was assessed using a Cell Counting Kit-8 (CCK-8, Vazyme). Briefly, cells were seeded as single cells in the wells of a 96-well plate coated with Matrigel and transfected with si*ESRG* or control siRNA. At different time points after siRNA treatment, the cells were incubated for 4 h with CCK-8 at 37 °C. The absorbance of the formazan dye generated by the dehydrogenase activity in living cells was measured using a PARADIGM Detection Platform (Beckman Coulter) at a wavelength of 450 nm (reference, 650 nm).

**EdU assay**

For EdU assay, cells were pre-cultured with EdU for 2 h using a Mixture Reagent Kit (C10338-3, RiboBio) following the manufacturer’s protocol. The
cells were collected in a 15 mL centrifuge tube, then fixed with 4% paraformaldehyde for 30 min, and incubated with 0.5% Triton-X-100 in PBS for 20 min. After incubation with Apollo staining solution for 10 min, the cells were immediately detected by flow cytometry.

**Apoptosis assay**

Cells undergoing apoptosis were assessed using a FITC Annexin V Apoptosis Detection Kit I (BD Biosciences). Cells were carefully dissociated into a single-cell suspension using Accutase solution, washed twice with ice-cold PBS, and resuspended in 100 µL of 1× binding buffer at a density of 1×10^6 cells/mL. Subsequently, Annexin-V (5 µL) and PI (5 µL) solution, which label early and late apoptotic cells, respectively, were added to the cells. The mixture was then vortexed gently and incubated for 15 min at room temperature in the dark. After staining, 400 µL of 1× binding buffer was added, and the cells were analyzed using a FACSCanto II (Becton Dickinson).

**Microarray analysis**

Total RNA was extracted from the two independent samples (siControl versus siESRG) using the RNAiso reagent (Takara). Following the manufacturer’s instructions, RNA for analysis was labeled and hybridized onto the Illumina Human HT-12 v4 array by GenomeScan.

**PCR array analysis of the human p53 signaling pathway**

Gene expression profiles related to the p53 signaling pathway were analyzed using a Human p53 signaling pathway RT Profiler PCR Array (SABiosciences) according to the manufacturer’s instructions (QIAGEN). Total RNA was isolated using the RNeasy Mini Kit (QIAGEN) and purified with the RNase-Free DNase Set (QIAGEN), and cDNAs were synthesized using the RT First Strand Kit (QIAGEN) on a thermocycler PCR machine (Bio-Rad). Subsequently, RT SYBR Green Mastermix (QIAGEN) was used to evaluate the expression profiles of 84 key genes involved in apoptosis, cell cycle, cell growth, proliferation, differentiation, and DNA repair on a CFX96 qPCR machine (Bio-Rad). Data analysis was performed using PCR Array Data Analysis web portal (www.SABiosciences.com/pcarraydataanalysis.php). Gene expression changes with a fold change greater than 2 were considered biologically significant.

**Immunofluorescence-FISH (IF-FISH)**

The cells were fixed with 4% paraformaldehyde, digested with protease K and incubated at 37 °C for 1 h. Then, a FISH probe was added, and the mixture was incubated at 4 °C overnight followed by nuclear staining the next day. A FISH/IF assay is a combination of RNA FISH and protein IF as described previously [32, 33].

**Northern Blot analysis**

An ESRG-specific DNA probe (ESRG-F: 5’ ATGAAAGGGAAGACATACAAA 3’, ESRG-R: 5’ TGAACATAGCAAGGGAAA 3’) with a length of 314 bp was obtained by PCR. Total RNA was extracted from cell lines by TRIzol method and detected by northern blotting. The RNA samples were subjected to agarose gel electrophoresis under denaturation conditions, and then the RNA on the gel was transferred to the solid phase support in situ. After dry roasting, the samples were hybridized with an ESRG probe and detected by specific autoradiography.

**RNA pulldown and mass spectrometry analysis**

In vitro biotin-labeled RNAs (ESRG, its antisense RNA, and four truncated RNAs from ESRG) were transcribed with a biotin RNA labeling mix (Roche) and T7 RNA polymerase (Roche), and then purified with an RNeasy Mini Kit (QIAGEN) after treatment with RNase-free DNase I (Promega). Three micrograms of biotinylated RNA was heated to 95 °C for 2 min, and then left at room temperature (RT) for 20 min to form the proper secondary structure before use. Biotinylated RNA was incubated with cellular protein extracts and streptavidin agarose beads (Invitrogen), and then the pulled down proteins were separated on SDS-PAGE gels followed by mass spectrometry.

**MMP (mitochondrial membrane potential) analysis**

MMP was analyzed by fluorescence microscopy using the mitochondrial membrane potential assay kit with JC-1 (Beyotime) following the manufacturer’s instructions. Briefly, cells were rinsed with PBS, incubated with a mixture of 1 mL of culture medium and 1 mL of JC-1 working solution for 20 min at 37°C, washed twice with JC-1 buffer solution, and analyzed using a Nikon Eclipse TE2000-S inverted microscope (Nikon). At high MMPs, JC-1 can form aggregates in the mitochondrial matrix and fluoresces red, whereas at low MMP, JC-1 exists in the monomeric form outside the mitochondrial matrix and fluoresces green fluorescence. The fluorescence intensity shift from red (JC-1 aggregates) to green (JC-1 monomers) indicates a decrease in the MMP and early stages of apoptosis.

**RIP (RNA immunoprecipitation)**

The RIP assay was performed with the
EZ-Magna RIP Kit (Millipore, USA) according to the manufacturer’s instructions. Anti-MCM2 antibody was purchased from Proteintech. The coprecipitated RNAs associated with MCM2 were extracted with the TRIzol reagent, and ESRG enrichment was examined using qPCR. Enrichment associated with normal rabbit IgG served as the controls.

**Co-IP (Co-immunoprecipitation)**

For the co-immunoprecipitation (Co-IP) assay, proteins were harvested in lysis buffer (Beyotime Biotechnology) and supplemented with protease inhibitor (Selleck). After culturing with primary antibody as indicated in the figure legends, or mouse IgG for 4 h, protein A/G PLUS beads (sc-2003, Santa Cruz Biotechnology) were added and incubated overnight. The precipitants were washed at least three times with lysis buffer.

**Luciferase reporter construction and luciferase activity assay**

DNA fragments (approximately 2000 bp) upstream of the transcription starting site of the ESRG gene were PCR-amplified using genomic DNA from hESCs as a template. The primers used were 5’-CGGGGTACCCCATCCCACCAAAACTTACT-3’ (forwards, -2049 to -2029; the underlined part indicates a KpnI restriction site) and 5’-TCCCCGGGAGAGAGTCACGAAGGGAGATAA-3’ (reverse, -12 to +11; the underlined part indicates a SmaI restriction site). PCR fragments were digested by KpnI and SmaI, inserted into a pGL3-basic vector and sequenced. To construct the pESRG-OCT4B-luciferase reporter, DNA fragments encompassing OCT4-binding elements upstream of the ESRG transcription start sites were PCR-amplified using hESC genomic DNA. The primers used were as follows: wild-type, 5’-CGGGGTACCCCATCCCACCAAAACTTACT-3’; forward primer, underlined part indicates a KpnI restriction site, italicized bases represent the OCT4-binding site [-1713 to -1721]) and 5’-TCCCCGGGAGAGAGTCACGAAGGGAGATAA-3’; reverse primer, underlined part indicates a SmaI restriction site). PCR fragments were digested by KpnI and SmaI, inserted into a pGL3-promoter vector, and sequenced. The same procedure was followed to obtain the reporter constructs pESRG-OCT4B-luciferase with mutations in the OCT4-binding elements of the ESRG promoter regions; however, the following was used as the forwards primer: (5’-CGGGGTACCCCATCCCACCAAAACTTACT-3’, in which the dashed line indicates the element substitutions).

hPSCs were plated in a 6-well plate at a density of 2×10^5 cells/well. After 24 h, pGL3 reporter plasmids were introduced into hESCs using FuGENE HD transfection reagent. Forty-eight hours later, the cells were washed twice and suspended in 500 μL of reporter lysis buffer (Promega), and firefly luciferase activity was measured using the Dual-Luciferase reporter assay system (Promega) and a GloMax 20/20 luminometer (Promega) according to the manufacturer’s protocol. The Renilla luciferase plasmid plRl-TK (Promega) was cotransfected as an internal control. The corresponding target mRNA sequence for the siOCT4 was GCTTCAAGAACCATGTGTAAA. Furthermore, the ORF of OCT4 was cloned from cDNA from H9 hESCs and inserted into the BamHI/XbaI restriction site of the pcDNA3.1 vector to obtain the OCT4 expression vector (pcDNA3.1-OCT4), which was then transfected into 293T cells using FuGENE HD according to the manufacturer’s protocol.

**ChIP assay**

ChIP analysis was performed according to the instructions provided in the EZ-ChIP Kit (Millipore). Chromatin was incubated overnight with anti-OCT4 antibody (Santa Cruz) or control rabbit IgG (Millipore). Precipitated DNA was further purified and amplified with PCR primers (Forwards: 5’-TGG CATTGCACTGAGAA-3’; Reverse: 5’-CTGACTGG CAGTTGGTG-3’) surrounding the OCT4 binding site upstream of the ESRG gene.

**Mice and teratoma assay**

NOD-PrkdscidIIrgem2/SMOC (NSG) mice (male, 4-6 weeks) were purchased from the Shanghai Model Organisms. Then, the mice were bred in specific pathogen free (SPF) conditions, and the mouse experiments were approved by the ethics committee of the Central South University. According to the methods of a previous study [34], H9-TetR-shESRG cells were pretreated with Y27632 for 3 h prior to injection, and mice received Dox in their drinking water 1 week before injection and during the whole period of teratoma formation (H9-TetR-shESRG+Dox). A 200 μL cell suspension mixture containing 50% Matrigel and 1×10^7 Y27632-treated cells was injected into the hindlimb muscles of the NSG mice. The teratoma tissues were removed 6-8 weeks after injection and fixed with 10% neutral formalin buffer. Then, tissue sections were prepared and stained with hematoxylin and eosin.

**Comet assay**

H9 hESCs were transfected with siESRG and siControl for 48 h and then harvested in PBS. The
comet assay was performed according to manufacturer’s protocol (KGA240, KeyGEN BioTECH). In detail, three layers of gels were prepared on slides with agarose at different dissolution points, in which the cell suspension was mixed with low melting agarose (LMA) as the second layer of gel. The slides were lysed by placing them in a dish with 20 mL of lysis buffer and then placed in the electrophoretic solution for unwinding. Finally, the samples were stained with PI and observed under a fluorescence microscope for capture image.

Statistical analyses

All analyses were performed using SPSS Statistics 19.0 software. The error bars represent the standard deviation (SD) of three independent experiments, and statistically significant differences by Student’s t test are indicated by *, **, and *** (or #, ##, and ###) indicating $P < 0.05$, $P < 0.01$, and $P < 0.001$, respectively. $P$ values $< 0.05$ were considered significant.

Results

ESRG maintains pluripotency of hPSCs

ESRG is located on human chromosome 3p14.3 and consists of four exons and three introns with a full length of 3153 nt [13]. Its high and exclusive expression in hPSCs was verified by northern blotting, which showed high expression of ESRG in hPSC lines such as H9, H1, RC1 and RC2-iPSCs but not in HFF and 293T cells (Fig. 1A). To demonstrate the dynamic changes in ESRG expression during hPSC differentiation, we induced hESC differentiation in vitro with retinoic acid (RA). As shown in Fig. 1B, along with the differentiation of H9 hESCs, the expression of ESRG dramatically decreased to undetectable levels, and the change trend of its expression level was similar to those of OCT4, SOX2 and NANOG. By in situ hybridization (ISH), we found that ESRG was mainly located in the nucleus of hPSCs, with a small part in the cytoplasm (Fig. S1A), which was further confirmed by cellular fractionation assays (Fig. S1B). Thus, ESRG may primarily exert its biological function in the nucleus.

To assess the functional significance of ESRG in hPSCs, twenty-five sets of siRNAs targeting different regions of the ESRG sequence were designed and transfected into H9 hESCs. Cy3-labeled negative control siRNA (5’-Cy3-siControl) was used to evaluate the transfection efficiency. As shown in Fig. S1C, nearly 100% of the cells were Cy3 positive, indicating that siRNA could be effectively delivered into hPSCs. Then, three siRNA sequences (si2364, si2243, and si2621) with the highest knockdown efficiency matching the ESRG sequence starting at nucleotides 2364, 2243 and 2621, respectively (Fig. S1D-F), were selected, renamed as siESRG1, siESRG2 and siESRG3, and used in the subsequent experiments. The qPCR results showed that the expression level of ESRG decreased by more than 75% at 48 h after transfection with any of the three sets of siRNAs (Fig. 1C). In addition, a panel of shESRG lentiviruses was also used to silence ESRG gene expression (with an interference efficiency >70%) in hPSCs (Fig. S1G).

Then, we knocked down ESRG expression in hPSCs (herein siESRG1, the same below). Forty-eight hours after the transfection, the hPSCs with reduced ESRG expression appeared dispersed and slender, and did not form compact colonies, whereas the cells transfected with siControl retained the typical cell and colony morphology of undifferentiated hPSCs (Fig. 1D and Fig. S1I), and these differences were even more obvious after 72 h (Fig. 1D). More importantly, targeting ESRG with three effective siRNA sequences caused identical phenotypic changes in hPSCs (Fig. 1D and Fig. S1J-K), confirming that this phenotype was not a result of off-target effects of the siRNAs. Furthermore, specific knockdown of ESRG in hPSCs by lentivirus carrying shESRG caused phenotypic changes similar to those caused by the highly efficient transfection of siESRG (Fig. S1H).

Differentiation was marked by the loss of alkaline phosphatase (AP) activity. AP staining analysis showed an almost complete loss of AP activity in the siESRG group compared with the apparent AP activity in the siControl group (Fig. 1E and Fig. S2A), indicating that knockdown of ESRG caused the differentiation of hPSCs. In addition, we observed that knockdown of ESRG in hPSCs resulted in significant downregulation of pluripotency markers such as OCT3/4, Cripto, FOXD3, Tert1, LIN28 and JMJD5 (Fig. 1F and Fig. S2B). Subsequently, we examined the protein expression levels of OCT4 and NANOG. As shown in Fig. 1G, the fluorescence signals of OCT4, SSEA4 and TRA-1-60 in the cells transfected with siESRG were weaker than those in the control cells. As expected, knockdown of ESRG caused significant downregulation of the steady-state protein levels of OCT4 and NANOG compared with control hPSCs, as detected by western blot (Fig. 1H and Fig. S2C). The altered differentiation state of the hPSCs after knockdown of ESRG in H9 was further identified by qPCR based on the expression of triploblastic marker genes, including endoderm-specific, mesoderm-specific and ectoderm-specific markers. As shown in Fig. 1I and Fig. S2D, these germ layer markers were all significantly upregulated, with the endoderm-specific markers increasing the most, suggesting that ESRG plays a role in repressing hPSC
Figure 1. ESRG maintains pluripotency of hPSCs. (A) Northern blot analysis of ESRG with DIG Probe in various cell lines. Lanes 1 to 6 were H9, H1, RC1-iPSC, RC2-iPSC, HFF and 293T cell lines, respectively. (B) After the differentiation of H9 hESCs was induced by retinoic acid (RA) in vitro, the expression of ESRG, OCT4, SOX2 and NANOG was detected by qPCR. (C) The RNA level of ESRG was analyzed by qPCR after transfecting each of the three effective siRNA sequences (siESRG1, siESRG2, siESRG3) for 48 h. (D) Brightfield images of H9 hESCs transfected with siESRG (herein siESRG1, the same below). Scale bar, 100 μm. (E) AP staining was performed in H9 hESCs transfected with siESRG or siControl, and the dark blue color indicated undifferentiated, AP-positive cells. Scale bar, 100 μm. (F) The expression of pluripotency marker genes was analyzed by qPCR in H9 hESCs transfected with siESRG or siControl. (G) H9 hESCs transfected with siESRG or siControl were stained with anti-OCT4, anti-SSEA4 and anti-TRA-1-60 antibodies (upper panel). Hoechst 33342 was used to label cell nuclei (lower panel). Scale bar, 100 μm. (H) Protein levels of OCT4, NANOG and LIN28A were detected by Western blot analysis in ESRG knockdown and control H9 hESCs. (I) The expression of endoderm, mesoderm, ectoderm and trophectoderm marker genes was analyzed by qPCR in H9 hESCs transfected with siESRG. (J) Teratomas were derived from H9-TetR-shESRG cells inoculated NSG mice with or without treatment of Dox. Data are presented as mean ± SD. *P < 0.05, **P < 0.01, ***P < 0.001 (two-tailed Student’s t test).
Furthermore, by transfecting pCAG-TetRnls and pSUPERIOR-shOCT4/-shESRG vectors into H9 hESCs (Fig. S2E-F), we established doxycycline (Dox)-inducible OCT4/ESRG RNAi systems (H9-TetR-shOCT4/H9-TetR-shESRG cells) allowing knockdown of OCT4 (Fig. S2G-H) or ESRG (Fig. S2I-J) selectively in hPSCs. Inducible knockdown of ESRG in hESCs resulted in obvious phenotypic changes and cell differentiation, as expected. To examine the role of ESRG in modulating the differentiation of hPSCs in vivo, a teratoma formation assay was performed, and H9-TetR-shESRG cells with or without Dox were subcutaneously injected into NOD-Prkdc<sup>cs</sup>/IL2rg<sup>em2</sup>/Smoc (NSG) mice. After approximately 6-8 weeks, the teratomas formed by injection of H9-TetR-shESRG cells with Dox were postponed and much smaller than those formed in the control group without Dox, and the higher the Dox concentration was, the smaller the tumors formed (Fig. 1J and Fig. S2K). When the H9-TetR-shESRG cells were pretreated with Dox for 5 days prior to injection, no teratoma developed (data not shown). Together, these in vitro and in vivo results suggest that ESRG functionally maintains pluripotency of hPSCs.

**ESRG is essential for hPSC self-renewal**

In addition to the differentiation induced by knockdown of ESRG, cell proliferation was decreased, suggesting that ESRG plays a critical role in maintaining hPSC proliferation and self-renewal. To investigate the effects of ESRG knockdown on hPSC proliferation and survival, we employed an EdU (5-ethynyl-2'-deoxyuridine) incorporation assay to quantitatively assess cell proliferation. As expected, ESRG knockdown cells showed less EdU incorporation than control cells (Fig. 2A-B and Fig. S3A), which was further confirmed via CCK-8 assay, and the cell growth rate of ESRG knockdown hPSCs decreased significantly compared with the control cells (Fig. 2C).

To further characterize the role of ESRG in hPSC survival, cell cycle analysis was performed using PI staining. The percentage of ESRG knockdown cells in the G2/M phase was significantly higher than that of the controls, showing that hPSCs subjected to ESRG silencing accumulated in G2/M phase of the cell cycle, with a concomitant decrease in the fraction of cells in G0/G1 or S phases (Fig. 2D-E and Fig. S3B). In addition, EdU/PI staining was used to analyze cell cycle distribution, and the results were consistent with those of PI staining alone (Fig. S3D).

Moreover, we found that the number of floating dead cells reminiscent of apoptotic cells in ESRG-silenced hPSC cultures was increased compared with that in cultures of control cells. Therefore, changes in apoptosis were detected using a FITC Annexin V apoptosis assay. The Annexin V/PI apoptotic cell population in ESRG-silenced cells was significantly increased at 48 h and 72 h after transfection (Fig. 2F-G and Fig. S3C). Next, analysis of the mitochondrial membrane potential (MMP) showed that ESRG knockdown cells exhibited a remarkable decrease in MMP compared with control cells (Fig. 2H), suggesting that ESRG knockdown resulted in hPSC apoptosis. Together, these results indicated the essential role of ESRG in sustaining hPSC cell survival and self-renewal. Similar results were obtained after the transfection with the three ESRG siRNAs (Fig. 2 and Fig. S3E-J).

To further confirm that these phenotypes and functions caused by ESRG knockdown are not the result of RNAi off-targets effects, we constructed an ESRG adenovirus vector and performed a rescue analysis. The results showed that overexpression of ESRG rescued the morphological changes caused by ESRG knockdown and reversed the loss of pluripotency and the inhibition of cell proliferation (Fig. S4). These results confirmed that this series of biological effects are indeed caused by ESRG knockdown, indicating that ESRG plays an important role in the maintenance of the self-renewal and pluripotency of hPSCs.

**ESRG is transcriptionally regulated by OCT4**

The core promoter region of ESRG was predicted to be -1,346 to -1,297 bp upstream of the transcription start site [13]. To identify the activity of the predicted promoter region, we amplified a fragment spanning from position -2049 bp to +11 bp (-2049/+11) of ESRG, cloned it into a pGL3 basic vector (pGL3-2049/+11), and transfected the vector into hPSCs. The results from the luciferase reporter assay showed that this fragment exhibited strong promoter activity compared with a blank control, which suggested that the -2049/+11 fragment plays an important role in triggering the transcription of ESRG in hPSCs (Fig. 3A).

Next, we used the online tool TESS (Neural Network Promoter Prediction) (http://fruitfly.org/seq_tools/promoter.html) to analyze the transcription factors that might regulate ESRG expression and found that the OCT4 binding site (ATTTCGAT) was -1714 to -1707 bp upstream of ESRG. Chromatin immunoprecipitation (ChIP) experiments verified that OCT4 directly bound to the OCT4 binding site within ESRG promoter (Fig. 3B). To explore the possible role of OCT4 in regulating ESRG transcription, siRNA targeting OCT4 mRNA was cotransfected into hPSCs with a luciferase reporter vector containing the ESRG promoter that includes the OCT4 binding site.
Downregulating OCT4 decreased the luciferase activity of the reporter vector group as compared with the control (Fig. 3C). Furthermore, pcDNA 3.1-OCT4, the plasmid expressing OCT4, was cotransfected with the pESRG-OCT4B-luciferase plasmid into 293T cells without endogenous OCT4 expression. A luciferase activity assay showed that ectopic OCT4 expression significantly increased the luciferase activity of the reporter vector group compared with that of negative control (Fig. 3D). These results suggested that OCT4 positively regulates ESRG transcription.

To further confirm that OCT4 regulates the activity of the ESRG promoter, we constructed a mutated luciferase reporter vector containing a mutated OCT4 binding consensus sequence with two nucleotides replaced (mutant pESRG-OCT4B-luciferase). The wild-type and mutant reporter vectors

![Figure 2. ESRG is essential for hPSC self-renewal.](https://www.ijbs.com)

(A) EdU assay was analyzed by flow cytometry in H9 hESCs transfected with siESRG or siControl at 24 h (first column), 48 h (second column) and 72 h (third column) after transfection. (B) The quantified analysis of EdU positive cells. (C) Data from the CCK-8 assay of ESRG knockdown and control H9 hESCs. (D) Cell cycle distribution was analyzed by flow cytometry in H9 hESCs transfected with siESRG or siControl at 24 h (first column), 48 h (second column) and 72 h (third column) after transfection. (E) The quantified analysis of cell cycle distribution. (F) Apoptosis was analyzed using a FITC/Annexin V apoptosis assay in H9 hESCs transfected with siESRG or siControl at 24 h (left column), 48 h (middle column) and 72 h (right column) after transfection. (G) The quantified analysis of the apoptosis assay. (H) H9 hESCs transfected with siESRG or siControl were stained with JC-1 to evaluate the MMP. Scale bar, 100 μm. All representative examples of the data from at least three independent experiments are shown. Data are presented as mean ± SD. *P < 0.05, **P < 0.01, ***P < 0.001 (two-tailed Student’s t test).
were transfected into hPSCs respectively, and the luciferase activity was analyzed at 48 h after transfection. As shown in Fig. 3E, the luciferase activity in the mutant reporter vector group was significantly lower than that in the wild-type reporter vector group, suggesting that the predicted OCT4 binding site upstream of ESRG acts as a positive cis-element in regulating ESRG transcription. qPCR analysis also indicated that silencing OCT4 downregulated ESRG in both siOCT4-transfected cells (Fig. 3F) and Tet-inducible shOCT4 hPSCs (H9-TetR-shOCT4) (Fig. 3G). Moreover, the level of ESRG downregulation was positively correlated with the silencing efficiency of OCT4, indicating that ESRG is a downstream target of OCT4 (Fig. 3F-G).

Figure 3. ESRG is transcriptionally regulated by OCT4. (A) The relative luciferase activities in hPSCs (H9 and RC1) transfected with a luciferase vector containing the -2049/+11 fragment of ESRG are similar to that of the pGL3 control vector. (B) A ChIP assay was performed in H9 hESCs lysates. (C) The relative luciferase activity in H9 hESCs co-transfected with either siOCT4 or siControl and either a pESRG-OCT4B-luciferase reporter plasmid or a pGL3 control plasmid. (D) The relative luciferase activity in 293T cells co-transfected with either pcDNA3.1-OCT4 or pcDNA3.1 (negative control) and either pESRG-OCT4B-luciferase reporter plasmid or pGL3 control plasmid. (E) Detection of luciferase activity in H9 hESCs transfected with a mutant pESRG-OCT4B-luciferase reporter plasmid harboring a mutated OCT4 binding site and a wild-type pESRG-OCT4B-luciferase reporter plasmid. The relative luciferase activity represents the ratio of firefly luciferase activity to Renilla luciferase activity (internal control). (F) QPCR analysis was performed for OCT4 and ESRG expression in H9 hESCs transfected with siOCT4 or siControl. (G) QPCR analysis of OCT4 and ESRG expression in two Dox inducible H9-TetR-shOCT4 cell clones (C1 and C4) treated with or without Dox. (H) Fluorescence intensity of ESRG-tdTomato reporter H9 hESCs treated with RA for 5 days. Red fluorescence represents ESRG expression. Scale bar, 100 μm. (I) Phase images of ESRG Pr-tdTomato reporter H9 hESCs treated with RA for 5 days. Red fluorescence represents ESRG expression. Scale bar, 100 μm. Data are presented as mean ± SD. **P < 0.01, and ***P < 0.001 by two-tailed Student’s t test.
ESRG functions in hPSCs by directly binding MCM2

ESRG functions in hPSCs by directly binding MCM2

We assessed the binding proteins of ESRG using RNA pull-down assays and mass spectrometry analysis (Fig. S5A). Using the antisense sequence of ESRG as a negative control, we obtained seventy-two proteins specifically binding to ESRG. Among them, we found that the DNA replication licensing factor MCM2 was specifically associated with ESRG (Table S1). The expression level of MCMs is positively correlated with cell proliferation and regeneration [35, 36]. As a component of the MCM2-7 complex, MCM2 is a putative replicative helicase that is essential for the initiation and elongation of “once per cell cycle” DNA replication in eukaryotic cells [37]. As we demonstrated in this study, knockdown of ESRG resulted in a serious impact on DNA replication and cell proliferation of hPSCs. Therefore, we hypothesized that ESRG might function in hPSCs by interacting with MCM2.

To prove this hypothesis, we first verified the association of ESRG with MCM2 through RNA pull-down. MCM2 was enriched in the sense strand groups compared with the negative control antisense groups in hPSCs (Fig. 4A and Fig. S5B). RNA immunoprecipitation (RIP) assays also confirmed an enrichment of ESRG in the complexes precipitated with the antibody against MCM2 (Fig. 4B and Fig.
SSC). In addition, immunofluorescence combined with RNA-FISH assays demonstrated that ESRG and MCM2 were mainly colocalized in the nuclei of hPSCs, further confirming their binding relationship (Fig. 4C and Fig. S5D).

Next, a series of ESRG deletion mutants were constructed to determine the nucleotides in ESRG that are responsible for binding to MCM2. As shown in Fig. 4D, further mapping of the region with RNA pull-down indicated that the 2,001-3,153 nt domain was required for the ESRG-MCM2 interaction in hPSCs. In the same way, to map the region or regions within MCM2 that interact with ESRG, we tested the binding of MCM2 truncation variants to ESRG. First, we constructed two MCM2 domains, namely, MCM2-N and MCM2-C (Fig. 4E). After pull-down assays and RIP experiments in H9 hESCs and 293T cells, we found that MCM2-N, but not the MCM2-C domain of MCM2, was required for ESRG binding (Fig. 4E-F). Thus, the MCM2-N domain of MCM2 was necessary and sufficient for ESRG binding. We further decomposed MCM2-N into MCM2-N1, MCM2-N2 and MCM2-N3 domains and found that ESRG specifically binds to the MCM2-N1 domain (Fig. 4G-H). Together, these data revealed that residues 1-256 of MCM2 and residues 2,001-3,153 of ESRG are critical for the ESRG-MCM2 interaction.

**ESRG sustains the steady-state level and nuclear location of MCM2**

We found that MCM2 protein levels were significantly reduced when ESRG was knocked down in both hESCs and hiPSCs (Fig. 5A). To further determine whether ESRG affects MCM2 stability, we treated hESCs with cycloheximide (CHX), a translation inhibitor, and found that ESRG knockdown markedly shortened the half-life of MCM2 protein in hESCs (Fig. 5B). Moreover, the ubiquitination of MCM2 was enhanced by ESRG knockdown (Fig. 5D). Then, we treated the cells with the proteasome inhibitor MG132 to prevent MCM2 degradation. As shown in Fig. 5C, the ESRG siRNA-induced MCM2 reduction was abolished by the proteasome inhibitor MG132. Similar results were obtained in hiPSCs after CHX and MG132 treatment (Fig. S5E-G).

Next, we investigated the mechanisms of ESRG-mediated inhibition of MCM2 ubiquitination and identified the factors targeting MCM2 ubiquitination. Previous studies have reported that two ubiquitin ligases, TRAIP and CRL2LRR1, can bind to the MCM2-7 complex to regulate DNA replication [38-40], so we speculated that they might be involved in ESRG-mediated MCM2 degradation. Therefore, we used siRNAs to interfere with the expression of ESRG and TRAIP or CRL2LRR1 at the same time and found that inhibition of TRAIP rather than CRL2LRR1 reversed MCM2 downregulation by ESRG siRNA (Fig. 5E and Fig. S5H). These results suggest that TRAIP can bind to MCM2 and mediate the posttranscriptional regulation of MCM2 by ESRG. Then, the binding of TRAIP to MCM2 was confirmed by a Co-IP assay (Fig. 5F). Importantly, ESRG knockdown enhanced the TRAIP/MCM2 interaction in hPSCs (Fig. 5F). Together, these data indicate that ESRG stabilizes MCM2 protein by interfering with the TRAIP/MCM2 interaction and inhibiting the ubiquitination of MCM2.

Based on the interaction of ESRG and MCM2, we examined whether ESRG affects the cellular localization of MCM2 as well as its expression level. Therefore, the changes in MCM2 localization were detected by assessing the nuclear and cytoplasmic fractions of hPSCs. Our results showed that knockdown of ESRG sequestered more MCM2 in the cytoplasm (Fig. 5G), which was also confirmed by an immunofluorescence assay (Fig. 5H and Fig. S5I). Taken together, these findings implied that ESRG affects the DNA replication and self-renewal of hPSCs by influencing the cellular localization of MCM2.

We next tested whether MCM2 is essential for ESRG function. We examined the effect of MCM2 overexpression in ESRG knockdown hPSCs and found that overexpression of MCM2 partially rescued the morphological changes and the reduction in clone formation ability caused by ESRG knockdown (Fig. 5J). Additionally, MCM2 overexpression also partially rescued ESRG knockdown-induced proliferation slowing and apoptosis enhancement (Fig. 5K-N). Interestingly, MCM2 not only affected the maintenance of self-renewal by ESRG, but also partially restored the expression of the pluripotency markers OCT4, SOX2 and NANOG (Fig. 5J). These data suggest that MCM2 is essential for ESRG to maintain hPSC self-renewal and pluripotency.

**ESRG-MCM2 maintains the self-renewal and pluripotency of hPSCs by suppressing the p53 signaling pathway**

MCM2 is involved in the initiation of DNA replication through the formation of the MCM2-7 complex, and the abnormalities in MCM2 and MCM2 depletion can activate a DNA damage response [38, 41, 42]. To evaluate the DNA damage in hPSCs after ESRG knockdown, a single-cell gel electrophoresis assay (comet assay) was performed. The migration of DNA fragments released from the nucleus forms the tail, with a longer tail length and tail olive moment representing severe damage. As shown in Fig. 6A-B, the olive tail moment was significantly increased after ESRG knockdown compared with the control group.
In addition, we found that after ESRG knockdown in hPSCs, γ-H2AX protein levels and foci formation increased (Fig. 6C-D and Fig. S6A-C), further confirming the induction of double strand breaks (DSBs). Importantly, this phenomenon could be partially reversed by MCM2 overexpression (Fig. 6C-D), revealing that siESRG caused DNA damage through MCM2. Then, DNA damage could trigger G1/S or G2/M arrest as a protective response, giving the cells time to repair before division [43], which is consistent with our previous results (Fig. 2D-E).

Figure 5. ESRG sustains the steady-state levels and nuclear location of MCM2. (A) The expression of MCM2 was detected by Western blot after transfection of H9 cells and RC1-iPSCs with siESRG. (B) The expression of MCM2 in H9 cells (siControl vs siESRG) was detected by Western blot after treatment with CHX (20 μg/mL) for various
time periods respectively. (C) H9 cells were transfected with ESRG siRNA and pre-incubated with MG-132 (20 μM) for 4 h. Cell lysate was immunoblotted by anti-MCM2. (D) H9 cells were pre-incubated with MG-132 (20 μM) for 4 h. Ub was immunoprecipitated (IP) by anti-MCM2 and immunoblotted (IB) by anti-Ub. The ubiquitination of MCM2 protein was detected after ESRG knockdown. (E) H9 cells were transfected with TRAIP siRNA and pre-incubated with MG-132 (20 μM) for 4 h. Cell lysate was immunoblotted by anti-MCM2. (F) H9 cells were pre-incubated with MG-132 (20 μM) for 4 h. Ub was immunoprecipitated (IP) by anti-MCM2 and immunoblotted (IB) by anti-Ub. The ubiquitination of MCM2 protein was detected after ESRG knockdown. (G) H9 cells were transfected with TRAIP siRNA followed by ESRG knockdown. MCM2 protein expression was detected by Western blot. (H) MCM2 overexpression partially rescued morphological changes (I) and the protein levels of OCT4 and NANOG (J) induced by ESRG knockdown. The quantified analyses of EdU and apoptosis assay are shown in (L) and (N). All representative examples of the data from at least three independent experiments are shown. Data are presented as mean ± SD. **P < 0.01 by two-tailed Student’s t test.

Figure 6. ESRG-MCM2 maintains the cell survival and self-renewal/pluripotency of hPSCs by suppressing the p53 signaling pathway. (A) Representative pictures of comet assay performed 48 h in H9 hESCs after treatment with siControl and siESRG. Scale bar, 200 μm. (B) The quantified analysis of comet assay. (C and D) H9 hESCs were transfected with MCM2 vector followed by ESRG knockdown. γ-H2AX protein expression was detected by Western blot (C) and immunofluorescence staining (D). Scale bar, 100 μm. (E) Western blot was performed to analyze the protein levels of p53, Pho-p53, ATM and Pho-ATM in H9 cells treated with siControl and siESRG. (F and G) The expression of p53 signaling pathway genes was analyzed by qPCR (F) and Western blot (G) in H9 hESCs transfected with siESRG or siControl. (H) Western blot was performed to analyze the protein levels of apoptosis-related proteins in H9 cells treated with siControl and siESRG. All representative examples of the data from at least three independent experiments are shown. Data are presented as mean ± SD. *P < 0.05, **P < 0.01 and ***P < 0.001 by two-tailed Student’s t test.

It has been reported that ATM kinase is activated after DNA damage, and ATM rapidly phosphorylates p53, which then activates the p53 pathway [44, 45]. Here, we detected the phosphorylation levels of ATM kinase and p53 in hPSCs after ESRG knockdown, and the results confirmed this hypothesis (Fig. 6E). Pathway information generated by KEGG showed that ESRG knockdown led to the changes in multiple signaling pathways, including the p53 pathway, DNA replication pathway, cell cycle pathway, PI3K-Akt pathway, apoptosis pathway and pathways regulating the pluripotency of stem cells (Fig. S6D and Table S2). Significant changes in the expression of many important components (p21, DR5, 14-3-3σ, GADD45A, Wec1, etc.) of the p53 signaling pathway were observed between ESRG knockdown and control cells, which was also confirmed by qPCR and PCR Array of Human p53 Signaling Pathway (Fig. 6F and Fig. S6E-F). Furthermore, the protein levels of p53, p21, DR5, GADD45A and 14-3-3σ were also significantly increased in ESRG knockdown hPSCs (Fig. 6G).
The protein levels of the G2/M transition components of the cell cycle, including CDK1, CCNB1 and CCNA2, were significantly downregulated, and the expression levels of proapoptotic proteins, including Cyto C, BAX, Caspase3, cleaved (c)-Caspase3, Caspase7, c-Caspase7, PARP and c-PARP, were upregulated in ESRG knockdown hPSCs compared with the control cells (Fig. 6H and Fig. S6G). The results of ESRG knockdown are consistent with previously reported studies indicating that MCM2 knockdown leads to DNA damage and reduced cell proliferation [41, 42], and our further experiments showed that inhibition of p53 could partially rescue these phenomena (Fig. S6H-I). Collectively, these results suggested that ESRG knockdown leads to MCM2 degradation and nuclear export, resulting in DNA damage and activation of the p53 signaling pathway, thus impairing pluripotency and self-renewal of hPSCs (Fig. 6I).

However, a recent study by Takahashi, et al., demonstrated ESRG is dispensable when ESRG-knockout hPSC lines are generated under p53 siRNA treatment [20]. To explain the seemingly contradictory observations, we hypothesize that temporarily knocking down p53 during the construction of ESRG knockout cell lines may have led to the different results. We tested the hypothesis with H9-TetR-shESRG inducible cell line (Fig. 7). When ESRG silencing was induced with Dox without p53 inhibition, the number of dead cells increased significantly, and almost all the cells died by day 7 (Fig. 7A and 7D). However, Dox-induced cells grew normally under continuous treatment with p53 inhibitor PFTα (Fig. 7B and 7D). After p53 inhibition was stopped on Day 3, severe cell death was observed again, however, some cells survived with normal stem cell morphology (Fig. 7C and 7D). We cultured and passaged the surviving cells and found that the cell morphology was consistent with normal hPSCs (Fig. 7E). Then we collected cells for a series of tests. qPCR and western blot results showed that compared with normal cells, the expression of ESRG was extremely low while p53 expression was normal (Fig. 7F-H). Immunofluorescence assay also showed the normal expression of OCT4 and NANOG (Fig. 7I). These results suggest that hPSCs can survive when ESRG is knocked down with p53 inhibition, which agrees with the report by Takahashi, et al. [20].

**Discussion**

ESRG, which was identified in our previous studies, exists only in human and some primate cells and is highly expressed only in hPSCs, suggesting that ESRG has unique significance in hPSCs. Wang, et
al., found that ESRG is required for the maintenance of the self-renewal and pluripotency of hESCs, as knockdown of ESRG leads to the differentiation of H9 hESCs [19]. Furthermore, the studies of Rand et al. and Sekine et al. also support our results [17, 18]. However, Takahashi, et al., showed that ESRG is dispensable for human pluripotency, which causes a great discrepancy [20]. This paper has addressed the controversy from a variety of aspects.

We demonstrated that a decrease in ESRG expression caused a drastic change in cell and colony morphology, with the loss of several stem cell markers, including OCT4, SOX2 and NANOG. hESCs and hiPSCs changed into scattered, long and thin differentiated cells with a slow proliferation rate when ESRG expression was downregulated. Recent studies have clearly shown that the self-renewal and differentiation of stem cells are closely connected to cell cycle progression [46, 47]. OCT4, SOX2, NANOG, JMJD5, p53 and the RB family [48-50] control the self-renewal and/or pluripotent status of hPSCs by cell cycle regulation. Our data showed that ESRG also plays an important role in regulating the cell cycle progression of hPSCs. Specific repression of ESRG in hPSCs caused the accumulation of cells in the G2/M phase and resulted in increased apoptosis and decreased cell proliferation. We used different siRNA sequences and obtained consistent results. Rescue experiments also showed that there were no off-target effects. Then, we used BLAST and other tools to compare siRNA sequences, and all genes that might be affected were detected by qPCR to exclude possible off-target effects. In addition, several attempts were made to construct ESRG knockout cell line by using CRISPR technology without p53 knockdown, but none of them were successful, which further demonstrates the importance of ESRG for hPSCs. In fact, studies have shown that ESRG can serve as a pluripotency marker, which is turned on as early as day 3 after reprogramming induced by OCT3/4, SOX2 and KLF4 (OSK), and can be used as a marker for early reprogramming of cells and undifferentiated hPSCs [17, 18]. These studies also strongly support our results regarding the essential role of ESRG in hPSCs.

OCT4, SOX2 and NANOG form a regulatory feedback circuit to maintain pluripotency in PSCs [3, 51]. In this circuit, all three transcription factors regulate themselves and each other. This regulatory circuit is essential to PSC identity [11, 52]. Previous bioinformatics analysis predicted an OCT4 binding site upstream of the ESRG gene, suggesting that ESRG may be regulated by OCT4 [13]. Our results showed that ESRG is a downstream target of OCT4 in hPSCs. We thus concluded that the interaction of OCT4 with the regulatory region of ESRG serves to enhance the transcription of ESRG. ESRG is involved in the OCT4, SOX2 and NANOG transcriptional regulation network.

LncRNAs usually function in stem cells by binding to proteins [53-56]. Here, seventy-two proteins specifically bound to ESRG were screened out by RNA pull-down and mass spectrometry. After further experimental verification, we identified that ESRG bound to MCM2, a replication-licensing factor, to sustain its steady-state level and nuclear location. MCM2, as a member of the MCM family of proteins, encodes a protein of 904 amino acids containing a presumptive zinc-finger domain [57]. It forms a double hexamer with MCM3-7 and loads onto chromatin with the origin recognition complex, Cdc6 and Cdt1, to form replication complexes and initiate DNA replication [58]. The importance of MCM2 in DNA replication and cell proliferation makes it a highly specific and sensitive marker, which has diagnostic and prognostic significance in various human malignant tumors [59-63]. As an indispensable member of MCM2-7, MCM2 plays a significant role in DNA replication, and the loss of MCM2 can inhibit cell proliferation [64-66] and lead to cell death due to DNA damage [41, 67, 68].

In response to DNA damage, p53 maintains genomic stability by regulating the expression of its downstream target genes (p21, GADD45A, 14-3-3σ and DR5), which are involved in apoptosis, the cell cycle, cell proliferation, cell differentiation, and DNA repair [69-70]. p53 expression is induced during hPSC differentiation, and a highly undifferentiated state and regenerative potential of hPSCs can be obtained by suppressing p53 [8]. In this study, we showed that ESRG knockdown activated the p53 signaling pathway, leading to cell cycle arrest, reduced proliferation, and increased apoptosis and differentiation of hPSCs, and these results are similar to the effects of knockdown of OCT4, Rem2 and Aurka in hPSCs [71, 72]. Thus, our results suggested that ESRG knockdown leads to MCM2 degradation and nuclear export, resulting in DNA damage and the activation of the p53 signaling pathway, leading to cell proliferation slowing, cell cycle arrest and apoptosis, thus impairing the cell survival and self-renewal/pluripotency of hPSCs (Fig. 6f).

Our findings of ESRG’s essential roles are consistent with a few previous reports. However, a recent study by Takahashi, et al., demonstrated ESRG is dispensable when ESRG-knockout hPSC lines are generated under p53 siRNA treatment [20]. Through our experiments, we found that hPSCs can survive when ESRG is knocked down with p53 inhibition, which agrees with the report by Takahashi, et al. Based on these results, we concluded that the hPSCs
are extremely sensitive to p53 when ESRG is knocked down, and once the cells tolerate it, the cells will no longer be affected by p53. Although p53 inhibition is temporary, the effect is critical to prevent acute cell death caused by the loss of ESRG. As for why p53 is so important for ESRG knockdown effects and how the cells adapt after a few days, it awaits further investigation.

In addition to MCM2, there are other proteins that bind to ESRG with high affinity, such as PSMD11, PODXL and NPM1. Although these proteins have not been shown to play a role in hPSCs by interacting with ESRG, several studies have shown that they are important for self-renewal and maintenance of pluripotency in stem cells. For example, hPSCs have higher proteasome activity than differentiated cells, which is induced by PSMD11/RPN6, a scaffold subunit that promotes proteasome assembly [73, 74]. Undifferentiated hPSCs can be isolated using a specific recognition antibody against PODXL, which can also be used in stem cell therapy and has been patented. NPM1 is an essential gene for embryonic development and the proliferation of embryonic stem cells [75], and is closely related to the proliferation and apoptosis of neural stem cells [76]. These studies suggest that in addition to maintaining the proliferation and self-renewal of hPSCs by binding to MCM2, ESRG combines with other proteins to participate in the vital activities of hPSCs.

Overall, this study identified a clear role of ESRG in maintaining cell survival of hPSCs. We expanded on the significance of ESRG as a target of the stem cell regulatory factor OCT4 in hPSCs and demonstrated that in hPSCs, ESRG binds with MCM2 to sustain the steady-state level and nuclear location of MCM2, inhibit cell differentiation, promote cell proliferation and inhibit cell apoptosis by suppressing p53 signaling. These findings contribute to our understanding of the molecular mechanisms of self-renewal and pluripotency maintenance of hPSCs and provide insights for further biological study and the development of promising clinical applications of ESRG in stem cell therapy.

**Abbreviations**

AP: alkaline phosphatase; bFGF: basic fibroblast growth factor; ChIP: chromatin immunoprecipitation; CHX: cyclohexamide; CM: conditioned medium; Co-IP: co-immunoprecipitation; Dox: doxycycline; DSBs: double strand breaks; E8 medium: essential 8 medium; hESCs: human embryonic stem cells; hiPSCs: induced pluripotent stem cells; hPSCs: human pluripotent stem cells; IF-FISH: immunofluorescence-FISH; ISH: in situ hybridization; IncRNAs: long non-coding RNAs; MCM2: minichromosome maintenance 2; NSG: NOD-PrkdcscidI L2rgem2/Smoc1; OCT3/4, SOX2 and KLF4; RIP: RNA immunoprecipitation; RT: room temperature; SD: standard deviation; SPF: specific pathogen free.

**Supplementary Material**

Supplementary figures and tables.
https://www.ijbs.com/v19p0916s1.pdf

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**Author contributions**


**Data availability**

The data that support this study are available from the corresponding author upon reasonable request. Microarray data have been deposited in the NCBI GEO under accession GSE190957. Mendeley datasets are accessible at the website (http://dx.doi.org/10.17632/ yd24ws6s83.1).

**Competing Interests**

The authors have declared that no competing interest exists.

**References**


