LIN28A inhibits DUSP family phosphatases and activates MAPK signaling pathway to maintain pluripotency in porcine induced pluripotent stem cells

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ABSTRACT
LIN28A, an RNA-binding protein, plays an important role in porcine induced pluripotent stem cells (piPSCs). However, the molecular mechanism underlying the function of LIN28A in the maintenance of pluripotency in piPSCs remains unclear. Here, we explored the function of LIN28A in piPSCs based on its overexpression and knockdown. We performed total RNA sequencing (RNA-seq) of piPSCs and detected the expression levels of relevant genes by quantitative real-time polymerase chain reaction (qRT-PCR), western blot analysis, and immunofluorescence staining. Results indicated that piPSC proliferation ability decreased following LIN28A knockdown. Furthermore, when LIN28A expression in the shLIN28A2 group was lower (by 20%) than that in the negative control knockdown group (shNC), the pluripotency of piPSCs disappeared and they differentiated into neuroectoderm cells. Results also showed that LIN28A overexpression inhibited the expression of DUSP (dual-specificity phosphatases) family phosphatases and activated the mitogen-activated protein kinase (MAPK) signaling pathway. Thus, LIN28A appears to activate the MAPK signaling pathway to maintain the pluripotency and proliferation ability of piPSCs. Our study provides a new resource for exploring the functions of LIN28A in piPSCs.

Keywords: LIN28A; MAPK; Pluripotency; piPSCs; DUSP

INTRODUCTION
LIN28A has two nucleic acid-binding domains: i.e., cold shock domain and zinc-knuckle domain (Moss et al., 1997), which bind specific sequences and play vital roles in physiology (Balzer et al., 2010; Hafner et al., 2013; Polesskaya et al., 2007; Shyh-Chang & Daley, 2013; Xu et al., 2009; Zhu et al., 2011). In addition to RNA binding, LIN28A also functions as a transcription factor during reprogramming (Liao et al., 2008; 2011). In addition, LIN28A activates MAPK signaling pathway to maintain the pluripotency and proliferation ability of piPSCs. Therefore, we explored the function of LIN28A in piPSCs based on its overexpression and knockdown.
Yu et al., 2007). LIN28A can regulate the naïve to primed state conversion by regulating stem cell metabolism in induced pluripotent stem cells (iPSCs) (Zhang et al., 2016). Knockdown of LIN28A promotes the transformation of embryonic stem cells (ESCs) into the naïve state (Chandrasekaran et al., 2017; Kumar et al., 2014; Marks et al., 2012), and LIN28A regulates pluripotent state transformation in mouse ESCs by inhibiting DPPA3 expression (Sang et al., 2019). However, few studies have explored LIN28A in porcine iPSCs (piPSCs) and the mechanism underlying LIN28A functions in piPSCs remains unclear. Previous research has indicated that inhibiting LIN28A expression via miR-370 may reduce piPSC proliferation ability and alkaline phosphatase (AP) activity, and up-regulate the expression of differentiation-relevant genes (Zhang et al., 2017). This finding differs from studies on LIN28A in human and mouse PSCs (Zhang et al., 2017). Thus, LIN28A may play a different role in piPSCs.

The mitogen-activated protein kinase (MAPK) signaling pathway plays important roles in controlling cell cycle, differentiation, proliferation, and apoptosis (Pearson et al., 2001; Shaul & Seger, 2007). Activation of the mitogen-activated protein kinase kinase (MEK)/extracellular signal-regulated kinase (ERK) signaling pathway can promote the differentiation of mouse ESCs (mESCs), while its suppression can prevent mESC differentiation (Burdon et al., 1999; Deathridge et al., 2019). MESC can be cultured in the 2i (CHIR99021 and PD0325901) system using MEK1 and glycogen synthase kinase-3 (GSK3) inhibitors (Ying et al., 2008). In vitro activation of MAPK signaling helps maintain the primed state, whereas repression of MAPK signaling through ERK inhibition reverts PSCs to the naïve state (Chen et al., 2015; Hackett & Surani, 2014; Kalkan et al., 2019; Ying et al., 2008). The dual-specificity phosphatase (DUSP) family dephosphorylates MAPK signaling and plays an important role in regulating the duration, magnitude, and spatiotemporal profiles of MAPK activity (Caunt & Keyse, 2013; Chen et al., 2019). According to previous reports, the pluripotency of piPSCs is rapidly lost following treatment with 1.0 μmol/L MEK1 inhibitor PD0325901 (Gao et al., 2019). Thus, piPSCs may differ from mESCs in MEK/ERK signaling requirements and inhibiting MAPK may impair their pluripotency.

Our laboratory previously reported that the doxycycline (DOX)-inducible porcine PSC line (DOX-piPSC) can be cultured with cytokines (leukemia inhibitory factor (LIF), basic fibroblast growth factor (bFGF)), signaling inhibitors (CHIR99021, SB431542), feeder cells, and serum (Ma et al., 2018; Zhu et al., 2021). PiPSCs can be maintained in the pluripotent state with the addition of DOX, but differentiate after its withdrawal (Ma et al., 2018; Zhang et al., 2017; Zhu et al., 2021). In this study, we explored the function of LIN28A by knockdown and overexpression. Results showed that proliferation ability and colony size decreased significantly when LIN28A was knocked down. Furthermore, piPSCs overexpressing LIN28A maintained colonies after DOX withdrawal. We also performed total RNA sequencing (RNA-seq) of negative control knockdown in the OEOT4-shNC group (OEOT4-shNC) and LIN28A knockdown in the OEOT4-piPSC group (OEOT4-shLIN28A) after withdrawal of DOX. Based on RNA-seq analysis, a reduction in LIN28A expression up-regulated differentiation-relevant gene expression and promoted neuroectoderm differentiation. LIN28A also inhibited the expression of DUSP family members and activated the MAPK signaling pathway to maintain the pluripotency of piPSCs.

**MATERIALS AND METHODS**

**Cell culture**

HEK293T cells were cultured in 6-well plates (140675, Thermo Fisher Scientific, USA) using Dulbecco's Modified Eagles Medium (DMEM) (HyClone, USA) with 10% fetal bovine serum (FBS) (V IS, New Zealand). Mouse embryonic fibroblasts (MEFs) were cultured in 100 mm vessels (140675, Thermo Fisher Scientific, USA) using DMEM (10% FBS) and treated with mitomycin for 2.5 h. The mitomycin-treated MEFs were then passaged at 1×10^5 cells/well into 12-well plates (140675, Thermo Fisher Scientific, USA) as the culture matrix for piPSCs. DOX-piPSCs were cultured in the LB2i system, which included 15% FBS, 0.1 mmol/L nonessential amino acids (NEAA) (Gibco, USA), 1 mmol/L L-glutamine (Gibco, USA), 10 ng/mL LIF (14890-HNAE, Sino Biological, China), 10 ng/mL bFGF (10014-HNAE, Sino Biological, China), 0.1 mmol/L β-mercaptoethanol (M3148, Sigma-Aldrich, USA), 3 μmol/L CHIR99021 (HY-10182, MCE, USA), 2 μmol/L SB431542 (S1067, Selleck, USA), and 4 μg/mL DOX (D9891, Sigma-Aldrich, USA). The piPSCs were passaged using TrypLE™ Select (Invitrogen, USA) into single cells at 2×10^4 cells/well in a 12-well plate every 5–6 days (Ma et al., 2018).

**Cell growth curve**

To obtain the cell growth curves for the negative control knockdown group (shNC), LIN28A knockdown group (shLIN28A1/2), negative control overexpression group (OEC), and LIN28A-overexpression group (OE LIN28A), cells were cultured in 24-well plates at an initial density of 1×10^5 cells/well. The piPSCs of the shNC, shLIN28A1/2, OEC, and OELIN28A groups were cultured in the LB2i system for 5 days and cell number in each group was counted daily using a blood counting chamber.

**Vector construction and cloning**

All lentivirus backbone vectors were derived from pCDH-CMV-MCS-EF1-GFP-T2A-PURO (www.zoores.ac.cn). shRNA of porcine LIN28A was designed using BLOCK-iT™ RNAi Designer (https://rnaidesigner.thermofisher.com/mairexpress/design.do) (Supplementary Table S1) and the relevant genes (Zhang et al., 2017). This finding differs from studies on LIN28A in human and mouse PSCs (Zhang et al., 2016). However, few studies have explored LIN28A expression up-regulated differentiation-relevant gene expression and promoted neuroectoderm differentiation. LIN28A also inhibited the expression of DUSP family members and activated the MAPK signaling pathway to maintain the pluripotency of piPSCs.

**Construction of short hairpin RNA (shRNA) vector**

Here, shRNA of porcine LIN28A was designed using BLOCK-iT™ RNAi Designer (https://maidesigner.thermofisher.com/mairexpress/design.do) (Supplementary Table S1) and the interference fragments were synthesized (enzymatic cleavage sites were BamH I and EcoRI). The double-stranded fragment was then connected to pCDH-U6-MCS-EF1-GFP-T2A-PURO linearized by BamH I and EcoRI via the T4 DNA ligase. All interference vectors were transfected into DOX-piPSCs and their interference efficiencies were verified, i.e., 80.78% (shLIN28A1) and 85.14% (shLIN28A2).
Construction of overexpression vector

Porcine testis cDNA was used as a template, and the porcine LIN28A fragment was successfully obtained using Prime Star Max DNA Polymerase (R045B, Takara, Japan). The fragment was then connected to pCDH-EF1-MCS-T2A-PURO linearized by BamHI and EcoRI via the T4 DNA ligase. The overexpression vector of porcine LIN28A was transduced to DOX-piPSCs and overexpression efficiency was detected.

Lentiviral packaging

HEK293T cells were cultured in a 6-well plate (140.675, Thermo Fisher Scientific, USA) at a density of 80%–90%. Lentiviral plasmids and packaged viral vectors (pVSV-G and psPAX2) were transfected into the HEK293T cells using polyethylenimine (PEI, Sigma-Aldrich, USA). In total, 1 μg of pVSV-G, 1 μg of psPAX2, and 2 μg of the lentiviral vectors were mixed in 12 μL of PEI (1 mg/mL). The plasmid mixture was then rested for 1 min at room temperature, after which 200 μL of optiMEM was added. After 12 h, the culture medium was replaced with DMEM. The HEK293T cells were then cultured for 48–72 h to produce lentiviral particles. The lentiviral particles (culture supernatant) were filtered through a 0.45 μm filter to remove cell debris.

Lentiviral particle transduction

The piPSCs were cultured in a 12-well plate covered with MEF at 37 °C for 12 h. The lentiviral particles (supernatant) and piPSC medium were mixed at a 1:1 ratio and 4 μg/mL polybrene was added to the mixture. The piPSCs were cultured with mixed medium at 37 °C for 8–12 h, which was then replaced by new iPSC medium. Fluorescence was observed after 2–3 days and puromycin was used to select puro-positive cells after the piPSCs were cultured for 1 week.

Total RNA extraction, reverse-transcription polymerase chain reaction (PCR), and quantitative real-time PCR (qRT-PCR)

Total RNA was extracted using the RNeasy Plus reagent (9108, Takara, Japan) via a three-step process. The total RNA was transcribed to obtain cDNA using a FastKing RT Kit (with gDNase) (KR116, day roots). Quantitative RT-PCR was performed using a SuperReal PreMix Plus (SYBR Green) (FP215, Tiangen, China) via a three-step process. The primers used for qRT-PCR are shown in Supplementary Table S2.

Western blot analysis

The piPSCs were cultured for 5 days were digested by TrypLE™ Select (Invitrogen, USA), and the same volume of DMEM was added to neutralize the reaction. The mixture was then transferred to a 1.5 mL tube and centrifuged at 5 000 g at 4 °C for 3 min. The supernatant of the culture medium was discarded and RIPA lysate (P0013B, Beyotime, China) with 10 mmol/L protease inhibitor FMSF (Sigma-Aldrich, USA) and phosphatase inhibitor was used to lyse the piPSCs for 30 min. Then, 5×SDS-PAGE loading buffer (JC-PE007, GENSHARE G, China) was added, followed by heating at 100 °C for 5 min. The protein samples were added to 8%–12% SDS-PAGE gel and run at 100 V for 1.5 h, then transferred onto polyvinylidenefluoride (PVDF) membranes at 15 V for 45 min using a Trans-Blot SD Semi-Dry Electrophoretic Transfer Cell (BioRad, USA). The membranes were then blocked using 8% skim milk (5% bovine serum albumin (BSA) in phospho-p44/42 MAPK and p44/42 MAPK) at room temperature for 2 h. Primary antibodies, including LIN28A (1 : 400; Santa Cruz Biotechnology, USA), proliferating cell nuclear antigen (PCNA) (1 : 500; Boster, China), FLAG (1 : 1 000; Sigma-Aldrich, USA), β-actin (1 : 4 000; Sungene Biotech, China), phospho-p44/42 MAPK (1 : 1 000; Cell Signaling Technology, USA), and p44/42 MAPK (1 : 1 000; Cell Signaling Technology, USA), were diluted in TBS-T buffer (20 mmol/L Tris HCl (pH 8.0), 150 mmol/L NaCl, 0.05% Tween 20) according to the instructions and then incubated at 4 °C for 12 h.

The PVDF membranes were washed using TBS-T buffer at room temperature for 15 min. AffiniPure Goat Anti-Mouse/Rabbit IgG (H+L) was then used to combine the antibody at 37 °C for 1 h (Hu et al., 2020). The membranes were washed using TBS-T buffer at room temperature for 15 min and a Thanon-5200 automatic chemiluminescence image analysis system (Tanom, China) was used to detect the horseradish peroxidase (HRP) signal. The relative gray is the western blots were analyzed by ImageJ.

Immunofluorescent staining

After twice washing with PBS, the piPSCs (cultured for 5 days) were fixed in 4% paraformaldehyde (pH 7.4) at room temperature for 15 min. We used 0.1% Triton-100 to perforate the membranes at room temperature for 10 min. Then, 10% FBS was used to block the membranes at room temperature for 1 h. The membranes were incubated with primary antibodies, including LIN28A (1 : 200; Santa Cruz Biotechnology, USA) and FLAG (1 : 1 000; Sigma-Aldrich, USA), for 12 h at 4 °C and then washed three times with PBS. The membranes were then incubated with goat anti-mouse IgG (H+L) secondary antibody Alexa Fluor 488 conjugate (1 : 500; ZSGB-BIO, China) at room temperature for 1 h and nuclei were stained with Hoechst33342 (1 : 1 000) at room temperature for 5 min (Ma et al., 2019; Wei et al., 2021).

AP staining

The piPSCs (cultured for 5 days) were fixed in 4% paraformaldehyde (pH 7.4) at room temperature for 15 min, with AST Fast Red TR and α-naphthol AS MX phosphate (Sigma-Aldrich, USA) then used to stain the cells according to the manufacturer’s instructions. The piPSCs were incubated in 1.0 mg/mL Fast Red TR, 0.4 mg/mL α-naphthol AS-MX, and 0.1 mmol/L Tris-HCL 8.8 buffer at room temperature for 20 min. The AP-positive piPSCs clones showed red (Zhang et al., 2017). Images were obtained using a Nikon phase difference microscope.

5-Ethynyl-2’-deoxyuridine (EdU) staining

EdU detection was performed on 3 d cultured piPSCs according to the Cell-Light EdU Apollo567 In-Vitro Kit instructions (RiboBio, China). The piPSCs were exposed to 50 μmol/L EdU medium at 37 °C for 20 min, and then fixed in
4% paraformaldehyde (pH 7.4) at 37 °C for 15 min. After this, 2 mg/mL glycine was added to neutralize excess aldehyde. The piPSCs were then exposed to 1×Apollo staining solution at 37 °C for 30 min and washed three times with PBS. We then used 0.1% Triton-100 to perforate the membranes for 10 min at 37 °C and the nuclei were stained with Hoechst33342 (1 : 1 000).

RNA-seq

To explore the molecular mechanism of LIN28A in piPSCs, the s/NC and shLIN28A2 groups with two biological replicates underwent RNA-seq. Total RNA was extracted using RNAiso Plus reagent (9108, Takara, Japan) with the guanidine isothiocyanate phenol-chloroform method (Chomczynski & Sacchi, 2006). Extracted RNA quality was detected using a NanoDrop™ spectrophotometer (Thermo Fisher Scientific, USA) and agarose gel electrophoresis. Total RNA was treated using Oligo dT-enriched mRNA and purification. Fragmented RNA was then reverse-transcribed using random N6 primers to form double-stranded DNA (second-strand cDNA synthesis with dUTP instead of dTTP). Next, 3'adenylated and adaptor ligation was added to the synthesized double-stranded DNA, which was amplified using PCR with specific primers. The PCR products were thermally denatured into a single strand and a bridging primer was used to form a loop to obtain a single-stranded circular DNA library. Quality control (QC) was conducted on the DNA library, and sequencing was then performed on the DNBSEQ platform. Raw reads were filtered through QC using SOAPnuke. Finally, Bowtie2 was used to align the clean reads. Heatmaps were plotted through heatmap (v1.0.12) to represent specific gene expression levels. Both Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway analyses were conducted using clusterProfiler (v3.12.0). An adjusted P-value of <0.05 and Q-value of <0.05 were used to define the working threshold for statistical significance.

Statistical analysis

Two-tailed t-tests were used to determine significant differences between two groups and one-way or two-way analysis of variance (ANOVA) was used to determine significant differences between three groups. All data are shown as mean±standard error of the mean (SEM). Differences were considered significant when P<0.05.

RESULTS

Effects of LIN28A on piPSC proliferation ability

LIN28A mRNA expression fluctuated under different concentrations of DOX (Supplementary Figure S1A), reaching a maximum level with the addition of 2 μg/mL DOX and then gradually decreasing with increasing concentrations of DOX (4 μg/mL to 16 μg/mL). Interestingly, colony size decreased with LIN28A expression after the administration of increasing concentrations of DOX (4 μg/mL to 16 μg/mL) (Supplementary Figure S1B).

Therefore, we next explored the relationship between the expression level of LIN28A and colony size. We designed two pairs of shRNA and constructed a LIN28A interference vector. The interference efficiency of the vector was detected by qRT-PCR. Results showed that the mRNA and protein expression levels of LIN28A decreased significantly in the shLIN28A1/2 groups (Figure 1A, B). Compared with the shNC group, colony size and AP activity decreased following LIN28A knockdown (Figure 1C). The cell growth curve showed that proliferation ability and protein expression of proliferating cell nuclear antigen (PCNA) decreased significantly when LIN28A was knocked down (Figure 1B, D). These results indicate that the proliferation ability of piPSCs decreased after LIN28A knockdown.

To further explore its function, porcine LIN28A was overexpressed in piPSCs (OELIN28A group) and its expression level was detected by qRT-PCR, western blotting, and immunofluorescent staining (Supplementary Figure S2A, B). Compared to the negative control overexpression group with DOX (OENC+DOX+), colony size and AP activity in the OELIN28A+DOX+ group did not change significantly (Supplementary Figure S2C). Furthermore, compared to the OENC+DOX+ group, the percentage of Edu-positive cells in the OELIN28A+DOX+ group did not change significantly (Supplementary Figure S2D, E). The cell growth curve also showed that there was no significant change in proliferation ability after overexpression of LIN28A (Figure 1D).

The DOX-piPSCs were maintained in the pluripotent state with DOX but differentiated after its withdrawal (Figure 1E, first column), consistent with previous studies (Ma et al., 2018; Zhang et al., 2017). The colonies in the negative control overexpression group without DOX (OENC+DOX−) gradually disappeared (Figure 1E, first column). However, colonies in the OELIN28A group without DOX (OELIN28A+DOX−) were still observed, although their size and number decreased significantly compared with the OENC+DOX+ and OELIN28A+DOX+ groups (Figure 1E, F). These findings indicate that LIN28A can maintain typical colonies after withdrawal of DOX. Pluripotent-relevant genes were detected by qRT-PCR. Results showed that the expression levels of SALL4 and NANOG increased and decreased, respectively, in the OELIN28A groups (Figure 1G). Thus, the knockdown and overexpression experiments showed that LIN28A plays a vital role in maintaining the proliferation ability of piPSCs.

Effects of LIN28A on piPSC pluripotency

LIN28A mRNA expression levels in the piPSCs fluctuated with DOX concentrations (Supplementary Figure S1A). Therefore, we explored the function of LIN28A after excluding the influence of DOX. Results showed that OEOCT4-piPSCs maintained colonies and proliferation ability in the absence of DOX (Supplementary Figure S3A). Therefore, experiments were performed on piPSCs overexpressing OCT4 (OEOCT4-piPSCs).

The interference efficiency of shLIN28A1 and shLIN28A2 was similar, but the proliferation rate of shLIN28A2 decreased more obviously according to the cell growth curve and expression of PCNA, so subsequent experiments were performed on the shLIN28A2 groups. Compared with the OEOCT4-shNC group without DOX, colony size decreased in the OEOCT4-shLIN28A2 group without DOX (Figure 2A, B), consistent with the shLIN28A2 group results (Figure 1C). The
Figure 1  Effects of LIN28A on piPSC proliferation ability
A: LIN28A mRNA expression levels in shNC, shLIN28A1, and shLIN28A2 groups with DOX were detected by qRT-PCR. Data are mean±SEM. *: P<0.01, n=3. B: Left: PCNA and LIN28A protein expression levels in shNC, shLIN28A1, and shLIN28A2 groups with DOX were detected by western blotting. Data are mean±SEM. n=2. Right: Quantitative analysis of PCNA and LIN28A protein expression levels in shNC, shLIN28A1, and shLIN28A2 groups with DOX is shown in histogram. Data are mean±SEM. **: P<0.01, n=3. C: Left: Morphology and AP staining of shNC, shLIN28A1, and shLIN28A2 groups with DOX. Scale bar: 100 μm for phase and 100 μm for AP. Right: Area ratio of colonies in shNC, shLIN28A1, and shLIN28A2 groups with DOX. **: P<0.01. D: Growth curve in shNC, shLIN28A1, shLIN28A2, OENC, and OELIN28A groups with DOX. Data are mean±SEM. *: P<0.05, *: P<0.01, n=3. E: Morphology and AP staining of the negative control overexpression group without DOX (OENC+DOX−), negative control overexpression group with DOX (OENC+DOX+), OELIN28A group without DOX (OELIN28A+DOX−), and OELIN28A group with DOX (OELIN28A+DOX+). Scale bar: 100 μm. F: Number of colonies in OENC+DOX−, OENC+DOX+, OELIN28A+DOX−, and OELIN28A+DOX+ groups. Data are mean±SEM. *: P<0.05, **: P<0.01, n=3. G: mRNA expression levels of pluripotent genes in OENC+DOX−, OENC+DOX+, OELIN28A+DOX−, and OELIN28A+DOX+ groups were detected by qRT-PCR. Data are mean±SEM. *: P<0.05, **: P<0.01, n=3. ns: No significance.
LIN28A mRNA expression level was detected again and was significantly decreased (Figure 2C). The percentage of EdU-positive cells also decreased significantly in the OEOCT4-shLN28A2 group (Figure 2D, E). The AP staining assays showed that AP activity and colony size decreased significantly in the OEOCT4-shLN28A2 group without DOX (Figure 2F). Based on AP staining assays (Sang et al., 2019), colonies can be classified into three shapes: i.e., typical primed diffuse-shape, compact dome-shape, and mixed-shape. Compact dome-shaped colonies in the OEOCT4-shNC group accounted for 60% of total colonies in this group, while compact dome-shaped colonies in the OEOCT4-shLN28A2 group accounted for ~50%; however, the total number of colonies in the OEOCT4-shLN28A2 group decreased significantly and colonies in the OEOCT4-shLN28A2 group basically disappeared (Figure 2G).

The results obtained for the OEOCT4-shLN28A2 group with DOX were similar to that of the OEOCT4-shLN28A2 group without DOX.

**Figure 2 Effects of LIN28A on piPSC pluripotency**

A: Left: Morphology of OEOCT4-shNC and OEOCT4-shLN28A2 groups without DOX. Scale bar: 100 μm. Right: Area ratio of colonies in OEOCT4-shNC and OEOCT4-shLN28A2 groups without DOX. **: P<0.01.  B: Morphology of OEOCT4-shNC and OEOCT4-shLN28A2 groups without DOX using fluorescence microscopy. Scale bar: 400 μm.  C: LIN28A mRNA expression levels in OEOCT4-shNC and OEOCT4-shLN28A2 groups without DOX were detected by qRT-PCR. Data are mean±SEM. *: P<0.05, **: P<0.01. n=3. D: EdU staining of OEOCT4-shNC and OEOCT4-shLN28A2 groups cultured on third day without DOX. Scale bar: 400 μm.  E: Percentage of EdU-positive cells in OEOCT4-shNC and OEOCT4-shLN28A2 groups cultured on third day without DOX. Scale bar: 400 μm. Data are mean±SEM. *: P<0.05. n=3.  F: AP staining of OEOCT4-shNC and OEOCT4-shLN28A2 groups without DOX. Scale bar: 100 μm for AP.  G: Quantification of AP-positive colonies in OEOCT4-shNC and OEOCT4-shLN28A2 groups without DOX. Data are mean±SEM. *: P<0.05, **: P<0.01. n=3.
Figure 3  LIN28A inhibited expression of differentiation-related genes
A: Number of up- and down-regulated DEGs in OEOCT4-shNC and OEOCT4-shLIN28A2 groups without DOX. B: Heat map shows expression of pluripotent genes in OEOCT4-shNC and OEOCT4-shLIN28A2 groups without DOX based on RNA-seq. C: Gene Ontology enrichment analysis of up-regulated DEGs, highlighting positive regulation of cell differentiation, neuronal differentiation, and negative regulation of cell proliferation. D: Heat map showing expression of genes involved in negative regulation of cell proliferation based on RNA-seq. E: Heat map showing expression of genes involved in positive regulation of cell differentiation and neuronal differentiation based on RNA-seq. F: mRNA expression levels of genes involved in positive regulation of cell differentiation, neuronal differentiation, and negative regulation of cell proliferation were detected by qRT-PCR. Data are mean±SEM. *, P<0.05; **, P<0.01. n=3.
group without DOX. Compared to the OEOCT4-shNC group with DOX, colony size (Supplementary Figure S3B, E) and AP activity in the OEOCT4-shLIN28A2 group with DOX decreased significantly (Supplementary Figure S3E). The percentage of EdU-positive cells also decreased significantly in the OEOCT4-shLIN28A2 group with DOX (Supplementary Figure S3C, D). The number of compact dome-shaped colonies in the OEOCT4-shLIN28A2 group with DOX decreased significantly compared with that in the OEOCT4-shNC group (Supplementary Figure S3F). These results indicate that pluripotency decreased after LIN28A knockdown and LIN28A plays a vital role in maintaining the pluripotency of piPSCs.

**LIN28A inhibited expression of differentiation-related genes**

We performed total RNA-seq on two samples in the OEOCT4-shNC and OEOCT4-shLIN28A2 groups after withdrawal of DOX. The table in Figure 3A shows the number of up- and down-regulated differentially expressed genes (DEGs) in the OEOCT4-shNC and OEOCT4-shLIN28A2 groups after DOX withdrawal. Among them, the expression levels of OCT4/SOX2/LIN28B showed no significant changes, but the expression level of LIN28A decreased in the OEOCT4-shLIN28A2 group (Figure 3B). After analyzing the RNA-seq results, GO analysis showed enrichment in positive regulation of cell differentiation, neuronal differentiation, negative regulation of cell proliferation, and DNA-binding transcription factor activity (Figure 3C). The expression of genes involved in the negative regulation of cell proliferation increased significantly in the OEOCT4-shLIN28A2 group based on heat map and qRT-PCR analyses that showed the expression levels of genes involved in neuronal differentiation and positive regulation of cell differentiation also increased significantly in the OEOCT4-shLIN28A2 group (Figure 3D, F), which may explain the significant decrease in AP activity after LIN28A knockdown. These data indicate that LIN28A can inhibit the expression of differentiation-related genes in piPSCs and maintain the proliferation ability and pluripotency of piPSCs.

**LIN28A inhibited expression of DUSP family and activated MAPK signaling pathway**

Based on KEGG analysis, the primary enriched pathways included Axon guidance, Notch signaling pathway, and MAPK signaling pathway (Figure 4A). The MAPK signaling pathway plays an important role in pigs, and its inhibition can result in loss of pluripotency in porcine PSCs (Gao et al., 2019). Both heat map and qRT-PCR analyses showed that the mRNA expression levels of DUSP-family members increased in the OEOCT4-shLIN28A2 group (Figure 4B, C), and the protein expression levels of ERK and phospho-ERK (p-ERK) in the OEOCT4-shLIN28A2 group significantly increased and decreased, respectively (Figure 4D). These findings indicate that the MAPK signaling pathway is inactivated following LIN28A knockdown. The DUSP mRNA expression levels were significantly decreased (Figure 4E) and the ERK and p-ERK protein expression levels were significantly decreased and increased, respectively, when LIN28A was overexpressed (Figure 4F). These results indicate that the MAPK signaling pathway was activated when LIN28A was overexpressed and LIN28A activated the MAPK signaling pathway by inhibiting the DUSP-family phosphatases. Results showed that cell proliferation ability and AP activity decreased when the MEK1 inhibitor (PD0325901) was used, consistent with the phenomena in the OEOCT4-shLIN28A2 cells (Figure 4G). Cell proliferation ability and AP activity decreased with the addition of 1 μmol/L PD0325901, but this decrease was rescued by the overexpression of LIN28A (Figure 4H). These results suggest that LIN28A can maintain the pluripotency and proliferation ability of piPSCs by activating the MAPK signaling pathway.

**DISCUSSION**

PiPSCs can be generated using human OCT4, SOX2, KLF4, and c-MYC lentiviruses in porcine fetal fibroblasts (Esteban et al., 2009; Ezashi et al., 2009). All piPSCs can exhibit terminal differentiation and generation of teratoma in vivo, but none can produce chimeric offspring and germline transmission, suggesting that their pluripotency may be defective (Cheng et al., 2012; Esteban et al., 2009; Ezashi et al., 2009; Xue et al., 2016; Zhang et al., 2015, 2017). In the past several years, various laboratories have obtained piPSCs and modified the culture system (Cheng et al., 2012; Haraguchi et al., 2012; Hou et al., 2016; Xu et al., 2020; Xue et al., 2016; Zhang et al., 2015, 2017). Here, we attempted to obtain naive piPSCs by modifying the patterns of gene expression.

Lin28a is the on-off switch between naïve and primed states. PSCs convert to the naïve state when Lin28a decreases, but to the primed state when Lin28a increases (Marks et al., 2012). Interestingly, in our study, piPSC proliferation, colony size, and AP activity all decreased following LIN28A knockdown (Figure 1B–D). We also found that the LIN28A mRNA expression level in piPSCs fluctuated with DOX concentration as LIN28A is regulated by OCT4/SOX2 (Supplementary Figure S1A) (Buganim et al., 2012). Therefore, we further explored the function of LIN28A excluding the influence of DOX. Results showed that the DOX-piPSCs started to differentiate after the withdrawal of DOX (Figure 1E, first column). By exploring single pluripotency gene function using a lentiviral overexpression system, we found that piPSCs maintained typical colonies after withdrawal of DOX when OCT4 was overexpressed (Supplementary Figure S3A) (Zhu et al., 2021). Compared with piPSCs, the cell proliferation and AP activities of the OEOCT4-piPSCs after DOX withdrawal showed no obvious differences. Therefore, experiments were performed on OEOCT4-piPSCs, with similar results as for piPSCs. Based on RNA-seq, we demonstrated that the pluripotency of the piPSCs disappeared and piPSCs differentiated into neuroectoderm cells when LIN28A was knocked down. Lin28a is also highly expressed in ESCs and is downregulated in response to differentiation (Balzer et al., 2010; Richards et al., 2004; Yang & Moss, 2003). Moreover, the expression levels of
LIN28B showed no significant changes, indicating that knockdown of LIN28A did not affect LIN28B expression. LIN28A also plays an important role in nervous system development (Faunes, 2020; Romer-Seibert et al., 2019; Yermalovich et al., 2020). The expression of genes involved in the negative regulation of cell proliferation, neuronal differentiation, and positive regulation of cell differentiation also increased in the OECOT4-shLIN28A2 group (Figure 3D–F). Thus, after LIN28A knockdown, the pluripotency of pIPSCs disappeared, the proliferation ability of pIPSCs decreased, and the pIPSCs differentiated into neuroectoderm cells. These results are consistent with previous study, which found that miR-370 can inhibit the expression of LIN28A (Zhang et al., 2017). To further explore its function, porcine LIN28A was overexpressed in the pIPSCs. However, colony size, proliferation ability, and AP activity demonstrated no significant change after LIN28A overexpression. This was not obvious with the addition of DOX but was observed after the withdrawal of DOX. The OELIN28A+DOX− group maintained typical colonies, whereas the OENC+DOX− group did not (Figure 1E), suggesting that LIN28A plays an important role in maintaining the proliferation ability of pIPSCs.

Previous immunohistochemical analysis reported that ERK phosphorylation is up-regulated in Lin28A transgenic mice (Kobayashi & Kozlova, 2018), which, in turn, promotes the phosphorylation and protein stability of LIN28A (Tsao & Daley, 2017; Tsao et al., 2017). We found that LIN28A inhibited the expression of DUSP-family phosphatases, which activated ERK signaling (Figure 4C–F). The mRNA expression levels of DUSP6/8/10 were markedly up-regulated after LIN28A knockdown. As a member of the DUSP family, DUSP6/8/10 inhibits ERK activity to regulate MAPK signaling in ovarian epithelial cancer (Gao et al., 2020), pancreatic cancer (Liu et al., 2021), and human epidermal stem cells (Hiratsuka et al., 2020). The increase in DUSP6/8/10 expression indicates inactivation of MAPK signaling (Comacchia et al., 2019). The weakly positive AP activity in the shLIN28A1/2 and OECOT-shLIN28A2 groups was consistent with the results obtained when DOX-pIPSCs were supplemented 1.0 μmol/L MEK1 inhibitor PD0325901 (Figures 1C, 2F, 4G). In vitro, MAPK is the key for maintaining the primed state, whereas PSCs transform into the naïve state with MAPK signaling repression (Chen et al., 2015; Hackett & Surani, 2014; Ying et al., 2008). However, the pluripotency of pIPSCs is rapidly lost with 1.0 μmol/L MEK1 inhibitor PD0325901 (Gao et al., 2019). This indicated that inactivation of MAPK signaling impaired the pluripotency of pIPSCs. We demonstrated that LIN28A maintained the pluripotency of pIPSCs by activating the MAPK signaling pathway. Therefore, further investigations on the molecular mechanisms underlying how LIN28A regulates its downstream genes and interacts with other transcription factors in pIPSCs are warranted.

**SUPPLEMENTARY DATA**

Supplementary data to this article can be found online.

**COMPETING INTERESTS**

The authors declare that they have no competing interests.

**AUTHOR CONTRIBUTIONS**

X.L.W., Z.S.Z., and J.L.H. designed the research. X.L.W., Z.S.Z., and S.Y. performed the research. X.L.W., X.X. and J.L.H. wrote the paper. X.X., O.Y.S., and M.Z.L analyzed the data. X.L.W., Z.S.Z., J.Q.Z, W.Y., R.Z., X.H., S.P., S.Q.Z., N.L., M.Z.L, and J.L.H. modified the manuscript. All authors read and approved the final version of the manuscript.

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