MicroRNA-32 silences WWP2 expression to maintain the pluripotency of human amniotic epithelial stem cells and β islet-like cell differentiation

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Abstract. Human amniotic epithelial stem cells (HuAECs) exhibit pluripotent characteristics, which are similar to those of embryonic stem cells, and can differentiate into various adult tissues and cells through directed induction. However, in culture, HuAECs tend to lose their pluripotency, and their directed differentiation capability declines with increasing passage number. The stem cell pluripotency factor octamer-binding protein 4 (Oct4) is an important transcription factor that promotes stem cell self-proliferation and maintains their pluripotency. Previous studies have demonstrated that WW domain containing E3 ubiquitin protein ligase 2 (WWP2) negatively regulates Oct4 expression and stem cell pluripotency. Therefore, the present study aimed to investigate the regulation of WWP2 by microRNAs (miRs), and to evaluate the expression of the downstream factor Oct4 and the maintenance of HuAEC pluripotency. Bioinformatics analysis identified a complementary binding site for miR-32 in the 3' untranslated region of the WWP2 gene, thus suggesting that it may be a target gene of miR-32. Post-infection of HuAECs with a vector overexpressing miR-32, the endogenous expression of WWP2 was significantly decreased, whereas Oct4 expression was significantly increased. Furthermore, miR-32-infected cells differentiated into β islet-like cells by directed induction. The results indicated that after induction, HuAECs overexpressing miR-32 also overexpressed the biomarkers of β islet-like cells. In addition, the ability to secrete insulin was markedly enhanced in response to glucose stimulation, in cells overexpressing miR-32. In conclusion, the present study suggested that miR-32 may effectively inhibit WWP2 expression in HuAECs and promote Oct4 overexpression to maintain their pluripotency.

Introduction

Originally derived from trophoblasts, human amniotic epithelial stem cells (HuAECs) are a specific type of cell that possess the pluripotency of embryonic stem cells (ESCs) (1-3). HuAECs can be induced to differentiate into various human tissues and cells, and have specific therapeutic effects in some human diseases (1-3). However, it is difficult to maintain the pluripotency of HuAECs for long periods in culture. Following primary isolation of HuAECs, pluripotency factors, including octamer-binding protein 4 (Oct4), SRY-box 2 (Sox2) and Nanog, are highly expressed; however, by the fifth passage, the expression levels of these factors are decreased to ~10% of the original levels, and after subsequent passages, HuAECs hardly express these factors. Furthermore, at this point, HuAECs can no longer be induced to differentiate into other adult cells, retaining only the phenotype and partial characteristics of epithelial cells (1). These issues have seriously hindered the use of HuAECs.

Ubiquitination is a post-translational modification process by which ubiquitin becomes covalently bound to a substrate protein by enzymatic action (4-8). The function of ubiquitination is to mark target proteins so that they are recognized and degraded by proteasomes (4-8). Comprising 76 amino acids, ubiquitin is a small polypeptide with a conserved sequence, which is present universally and ubiquitously in eukaryotic cells (4-8). The ubiquitination process begins with an E1 ubiquitin-activating enzyme, which, in an ATP-dependent reaction, activates a single free ubiquitin molecule by forming
a thioester bond between the Cys residue of the activity center of E1 and the C-terminus of ubiquitin. The activated ubiquitin is channeled to an E2 ubiquitin-conjugating enzyme, which, together with the recruited specific substrate protein and an E3 ubiquitin ligase, mediates the transfer of ubiquitin from E2 to the target protein. The recognition and degradation of ubiquitinated proteins by proteasomes is a non-specific process; therefore, the E3 ligase is a crucial factor in the process of protein degradation, determining the specificity of the protein degradation reaction. In most cases, the ubiquitination reaction involves one E1 ubiquitin-activating enzyme, and numerous E2 ubiquitin-conjugating enzymes and E3 ubiquitin ligases. Quantitatively, one E1 interacts with several E2 enzymes, whereas one E2 enzyme interacts with many more E3 ubiquitin ligases. The inverted ‘pyramid’ structure formed by E3, E2 and E1 is a large and complex signaling cascade network, which extensively and accurately regulates cell activities. The highly specific protein degradation mechanism in the cell is predominantly determined by E3 ubiquitin ligase, and all E3 ubiquitin ligases are capable of linking a target protein to a particular E2 enzyme. There are three major categories of E3 ubiquitin ligase: Those containing a homologous to the E6-AP carboxyl terminus (HECT) domain; those containing a RING domain; and those containing a U-box protein domain. The HECT domain-containing E3 ubiquitin ligases act mainly by forming a thioester bond with ubiquitin, which is essential to catalysis, whereas the RING domain-containing E3 ubiquitin ligases provide the binding sites for E2 enzymes and the substrate protein, so that the E2 enzyme can catalyze direct transfer of ubiquitin to the substrate (4-8). WW domain containing E3 ubiquitin protein ligase 2 (WWP2) (also known as atrophin-1-interacting protein 2) is a novel member of the E3 ubiquitin ligase family, which specifically induces ubiquitination of Oct4, ultimately leading to its degradation (9-14). Following the loss of ligase activity of WWP2 by a point mutation in the active site, the ubiquitination level of Oct4 is significantly reduced (9-14). Conversely, once a ubiquitin molecule is fused to the N-terminus of Oct4, a negative correlation is observed between Oct4 level and cellular ubiquitination level (9-14). In addition, when WWP2 is overexpressed in ESCs, it may inhibit the activity of endogenous Oct4 (9-14).

Based on these findings, the present study aimed to investigate whether microRNA (miR)-32 weakens the ubiquitination of Oct4 by specifically inhibiting WWP2 expression, thus maintaining the pluripotency of HuAECs and enhancing their capacity for differentiation into β islet-like cells by directed induction.

Materials and methods

HuAECs isolation and culture. The present study was approved by the ethics committee of Shanghai Geriatric Institute of Chinese Medicine, Longhua Hospital, Shanghai University of Traditional Chinese Medicine (Shanghai, China). From May 2015 to December 2015, human placentas were obtained from pregnant women (age, 28±3), who were negative for human immunodeficiency virus-I, hepatitis B and hepatitis C. The institutional ethics committee approved the use of human amnions. Amnion membranes were peeled mechanically from the chorions of placentas obtained from women that underwent an uncomplicated Cesarean section. The epithelial layers, with the basement membrane attached, were obtained and used to harvest HuAECs, as previously described, with some modification (15-17). Briefly, the membrane was placed in a 250-ml flask containing Dulbecco's modified Eagle's medium (DMEM) and was cut into 0.5-1.0 cm² segments using a razor. The segments were digested with 0.25% trypsin-EDTA at 37°C for 45 min. The resulting cell suspension was seeded in a 6-well plate in DMEM supplemented with 10% fetal calf serum (GE Healthcare, Chicago, IL, USA); penicillin (100 U/ml) and gentamicin (0.3 mg/ml) at 37°C in a humidified tissue culture incubator containing 5% CO₂. Successful infection of mutated miR (miR-Mut) and miR-32 was confirmed by RT-qPCR (Fig. 1).

Induction of HuAECs differentiation into β islet-like cells and C-peptide assay. According to a previous study (18), to induce their differentiation into insulin-producing β islet-like cells, HuAECs (1x10⁵) were seeded into a 10-cm dish and cultured in serum-free DMEM containing 25 mM D-glucose, N2 supplement (Gibco; Thermo Fisher Scientific, Inc., Waltham, MA, USA), 1% antibiotic-antimycotic and 10 mM nicotinamide (Sigma-Aldrich; Merck KGaA, Darmstadt, Germany) for 2 weeks; medium was changed twice a week. Cell culture supernatants were collected at appropriate time points, centrifuged to remove cell debris, and the C-peptide levels were assessed using a human C-peptide radioimmunoassay kit (China Diagnostics Medical Corporation, Beijing, China).

RNA extraction and reverse transcription-quantitative polymerase chain reaction (RT-qPCR) analysis. Total RNA was isolated from each cell sample using TRIzol® reagent (Invitrogen; Thermo Fisher Scientific, Inc.) according to the manufacturer’s protocol. RNA samples were digested with DNase I (Sigma-Aldrich; Merck KGaA), quantified and reverse transcribed into cDNA using a ReverTra Ace-α First Strand cDNA Synthesis kit (Toyobo Life Science, Osaka, Japan). A RealPlex4 real-time PCR detection system (Eppendorf, Hamburg, Germany) was used to conduct RT-qPCR, with SYBR-Green Real-time PCR Master mix (Toyobo Life Science) used as the detection dye. The RT-qPCR amplification conditions were as follows: 1 cycle of initial denaturation at 95°C for 2 min; and 40 cycles of denaturation at 95°C for 15 sec and annealing at 58°C for 45 sec and extension at 72°C for 42 sec; Then, 1 cycle of final extension at 72°C for 10 min. Target cDNA was quantified using the relative quantification method. The quantification cycle (Cq) method was used to determine gene expression relative to a control (calibrator), and steady-state mRNA levels were reported as the fold difference relative to the calibrator. For each sample, the marker gene Cq values were normalized using the formula \[ \Delta Cq = Cq_{\text{genes}} - Cq_{\text{18s ribosomal RNA (rRNA)}} \]. The relative expression levels were determined according to the following formula: \[ \Delta \Delta Cq = \Delta Cq_{\text{sample group}} - \Delta Cq_{\text{control group}} \]. The 2^(-\Delta\Delta Cq) method was used to calculate the relative expression levels of marker genes (19). The calibrator used was the 18s rRNA gene. The cDNA of each gene was amplified using primers shown in Table I.

Luciferase reporter assay. The luciferase reporter assay was performed according to a previously described method (15,20). NIH-3T3 cells were seeded at 3x10³/well in 48-well plates and were
cotransfected with 400 ng pRNAT-CMV-miR-32, pRNAT-CMV or pRNAT-CMV-miR-32-Mut, and 20 ng pGL-WWP2-3’ untranslated region (UTR)-wild-type (WT) or pGL-WWP2 3’UTR-Mut, and pRL-TK (Promega Corporation, Madison, WI, USA) using Lipofectamine® 2000 reagent (Invitrogen; Thermo Fisher Scientific, Inc.) according to the manufacturer’s protocol. After 48 h of transfection, luciferase activity was measured using a dual-luciferase reporter assay system (Promega Corporation).

**Immunofluorescence staining.** Cultured cells were washed three times with fetal calf serum and fixed with 4% paraformaldehyde (Sigma-Aldrich; Merck KGaA) for 30 min. After blocking, the cells were incubated with primary antibodies (Table II) overnight at 4°C, and then with goat anti-rabbit IgG H&L (Alexa Fluor® 555) pre-adsorbed (1:200; ab150086; Abcam, Cambridge, UK) and 5 µg/ml 4',6-diamidino-2-phenylindole (Sigma-Aldrich; Merck KGaA) for 30°C at 30 min. The cells were then washed thoroughly with Tris-buffered saline-Tween (TBST) [25 mM Tris/HCl (pH 8.0), 125 mM NaCl and 0.05% Tween-20] and viewed under a fluorescence microscope (DMI3000; Leica Microsystems, Inc., Buffalo Grove, IL, USA).

**Western blot analysis.** HuAECs were lysed in 2X loading lysis buffer [50 mM Tris-HCl (pH 6.8), 2% sodium dodecyl sulfate, 10% β-mercaptoethanol, 10% glycerol and 0.002% bromophenol blue]. The concentration of total protein was then determined by Enhanced BCA Protein Assay kit (Beyotime Institute of Biotechnology, Shanghai, China). The total proteins (15 µl, 25 µg/µl) from cultured cells were then separated by 12% SDS-PAGE and transferred onto Hybrid-polyvinylidene fluoride membranes (EMD Millipore, Bedford, MA, USA). The membranes were blocked with 5% (w/v) non-fat dried milk in TBST and were washed four times with TBST at room temperature (15 min/wash) prior to incubation with primary antibodies (Table II) in 5% (w/v) non-fat dried milk in TBST. After further washing, the membranes were incubated with horseradish peroxidase-conjugated goat anti-rabbit IgG secondary antibody (1:1,000; sc-2004; Santa Cruz Biotechnology, Dallas, TX, USA) at 37°C for 1 h. After washing four times with TBST at room temperature (15 min/wash), the immunoreactive protein bands were visualized.
Northern blot analysis. Northern blotting was performed according to a previously described method (15). For all groups, 20 µg good quality total RNA was analyzed on a 7.5 M urea, 12% polyacrylamide denaturing gel and was transferred to Hybond N⁺ nylon membrane (GE Healthcare). Ultraviolet light, for 30 sec at 1,200 mJoule/cm², was used to crosslink the membranes. The membranes were hybridized with the miR-32 antisense starfire probe, 5'-ATAACGCTGTAATGATTCAACGT-3' (Integrated DNA Technologies, Coralville, IA, USA), to detect the 22-nt miR-32 fragments, according to the manufacturer's protocol. After washing by Northern washing buffer (25 mM NaH2PO4·2H2O, 25 mM Na2HPO4·12H2O, 1 mM EDTA, 1% SDS, 5% formamide deionized), the membranes were exposed to Kodak XAR-5 films for 20-40 h (Sigma-Aldrich; Merck KGaA). All membranes were hybridized with a human U6 snRNA probe as a positive control, 5'-GCAGGGGCCATGCTATCTTC TCTGTATCG-3'. The exposure times for the U6 control probe varied between 15 and 30 min.

Co-immunoprecipitation (co-IP) assay. Cells were seeded at 3x10⁴/well in 6-well plates and were cultured until they reached 85% confluence. The cells were then lysed in 500 µl modified cell lysis buffer [20 mM Tris (pH 7.5), 150 mM NaCl, 1% Triton X-100, 1 mM EDTA, sodium pyrophosphate, β-glycerophosphate, Na3VO4 and leupeptin] (Beyotime Institute of Biotechnology) for western blotting and co-IP. After lysis, each sample was clarified by centrifugation to remove insoluble debris and the supernatant was preincubated with 20 µg protein A agarose beads (Beyotime Institute of Biotechnology) for 30 min at 4°C with agitation. Subsequently, the samples were centrifuged and transferred to a fresh 1.5 ml tube. The samples were incubated with antibodies (Table II) for 90 min prior to the addition of 20 µg protein A agarose beads to capture immune complexes. The pelleted beads were washed three times with 500 µl cell lysis buffer, dissolved in 4X SDS-PAGE sample loading buffer and heated for 10 min at 95°C.

KCl and tolbutamide stimulation experiments. According to the protocol of C-peptide RIA kit (China Diagnostics Medical Corporation), briefly, the 1x10⁴ cells/well were washed twice in DMEM (Gibco; Thermo Fisher Scientific, Inc.) containing 2 mM glucose (Sigma-Aldrich; Merck KGaA) and cultured in this medium for 1 h. The medium was then replaced by 2 ml DMEM containing glucose (0, 1, 5, 25 mM, respectively, for each well) or KCl (0, 10, 20, 40 mM, respectively, for each well; Sigma-Aldrich) or tolbutamide (0, 50, 100 mM, respectively, for each well; Sigma-Aldrich) for 60 min. The culture supernatants were collected after 12 h of incubation, centrifuged to remove debris, and concentration of C-peptide was tested by human C-peptide RIA kit.

Statistical analysis. Each experiment was performed at least three times. Data are presented as the means ± standard error. Student's t-test was used to evaluate differences between two groups. One-way analysis of variance was used to compare multiple groups. P<0.05 was considered to indicate a statistically significant difference.

Results

With increasing passage number, endogenous miR-32 expression and WWP2 expression are negatively associated in HuAECs. The results of the RT-qPCR analysis suggested that the mRNA expression levels of WWP2 were gradually increased in the HuAECs with continued passage (Fig. 2). Northern blot analysis confirmed that with continued passage of HuAECs, the hybridization signal of endogenous miR-32 became increasingly weaker (Fig. 2). These results indicated that endogenous miR-32 expression and WWP2 expression in HuAECs were negatively associated.

Targeted binding of miR-32 to the 3'UTR of WWP2 mRNA silences its expression. Sequence alignment, using bioinformatics analysis software (microRNA.org - Targets and Expression tools, http://34.236.212.39/microrna/home.do), identified a complementary binding site for mature miR-32 in the 3'UTR (between 1777 and 1783 bp) of WWP2 mRNA (Fig. 2). Luciferase reporter assays indicated that following overexpression of miR-32 in the NIH-3T3 cell line, the luciferase activity was significantly decreased when cells were cotransfected with WWP2 containing WT 3'UTR (Fig. 2). These results suggested that miR-32 could silence the mRNA expression of WWP2.

Exogenous overexpression of miR-32 effectively inhibits WWP2 expression in HuAECs. miR-32 was overexpressed in one group of HuAEC cells following infection with a lentivirus, whereas the negative control group of HuAECs were infected with miR-Mut. The two groups were cultured under the same conditions and passaged five consecutive times. Subsequently, RT-qPCR analysis indicated that, in HuAECs infected with miR-32, the mRNA expression levels of WWP2 were reduced at different passage numbers (passages 1, 2, 3, 4 and 5) compared with in the HuAECs infected with miR-Mut (Fig. 3). Furthermore, western blotting confirmed that at the 5th passage, miR-32-infected HuAECs exhibited a reduced abundance of WWP2 protein expression compared with in the miR-Mut-infected HuAECs (Fig. 3).

Exogenous overexpression of miR-32 inhibits Oct4 ubiquitination by WWP2 in HuAECs. RT-qPCR demonstrated that at the 5th passage, miR-32-infected HuAECs exhibited significantly increased expression levels of mRNAs encoding embryonic stem cell markers (Oct4, Sox2 and Nanog) compared with in the miR-Mut-infected HuAECs (Fig. 4). Furthermore, western blotting confirmed that at the 5th passage, miR-32-infected HuAECs exhibited significantly increased protein expression of embryonic stem cell markers (Oct4, Sox2 and Nanog) compared with in the miR-Mut-infected HuAECs (Fig. 4). In addition, western blotting of the co-IP samples indicated that, at the 5th passage, in miR-32-infected HuAECs Oct4 protein ubiquitination was reduced compared with in miR-Mut-infected HuAECs (Fig. 4). Immunofluorescence analysis revealed that the expression levels of Oct4, Sox2, Nanog and WWP2 were consistent with the western blotting
results (Fig. 4). These findings suggested that exogenous overexpression of miR-32 effectively inhibited Oct4 ubiquitination by WWP2 in HuAECs, and maintained the expression of transcription factors associated with ESC pluripotency.
Exogenous overexpression of miR-32 effectively maintains the capability of HuAECs to differentiate into β islet-like cells and express β islet-like cell markers. Under specific cell culture conditions, at the 5th passage, the two HuAEC cultures (infected with miR-32 or miR-Mut) were simultaneously induced to differentiate into β islet-like cells in vitro. After 21 days of induction, the results of qPCR suggested that, in the induced cell cultures, the mRNA expression levels of β islet-like cell biomarkers [insulin, pancreatic duodenal homeobox-1 (Pdx1), ISL LIM homeobox 1, paired box 6, NK2 homeobox 2, glucose transporter 2 and glucokinase] were significantly increased in miR-32-infected HuAECs compared with in miR-Mut-infected HuAECs (P<0.01 vs. miR-Mut HuAECs; P<0.05 vs. miR-Mut HuAECs; Student's t-test, n=3). Western blotting confirmed that, after induction, the protein expression levels of insulin and PDX1 were significantly increased in miR-32-infected HuAECs compared with in miR-Mut-infected HuAECs (Fig. 5). These results suggested that exogenous overexpression of miR-32 may effectively maintain the ability of HuAECs to differentiate into β islet-like cells and to express β islet-like cell markers.

Exogenous overexpression of miR-32 effectively enhances C-peptide secretion from β islet-like cells derived from HuAECs. C-peptide is secreted into the cell culture medium in amounts equal to those of insulin secreted by the pancreas; therefore, the C-peptide content of the supernatant of cell culture medium was determined, in order to investigate the synthetic insulin-secreting capability of β islet-like cells derived from the two HuAEC groups. Firstly, after directed induction of differentiation of HuAECs into β islet-like cells, the levels of C-peptide secreted by the two cell groups were analyzed at different time-points (1, 2 and 3 weeks). The
results demonstrated that, with increasing time after induction, the levels of C-peptide secreted by miR-32-infected HuAECs were much greater compared with those secreted by miR-Mut-infected HuAECs (Fig. 6 and Table III). Secondly, the levels of C-peptide secreted following stimulation of β islet-like cells with various concentrations of glucose were compared. The results indicated that with increasing glucose concentration, the amount of C-peptide secreted by β islet-like cells derived from miR-32-infected HuAECs was significantly increased compared with that secreted by cells derived from miR-Mut-infected HuAECs (Fig. 6 and Table III). In addition, following supplementation of the culture medium with various concentrations of KCl, the results indicated that, with increasing KCl concentration, the amount of C-peptide secreted by β islet-like cells derived from miR-32-infected HuAECs was much greater than that secreted by cells derived from miR-Mut-infected HuAECs (Fig. 6 and Table III). Furthermore, since blockage of the functionally sensitive intracellular $K_{ATP}$ ion channels may induce an increase in C-peptide secretion, tolbutamide, an ATP-sensitive $K_{ATP}$ ion channel inhibitor, was added to the culture medium of each cell group. The results indicated that with increasing tolbutamide concentration, C-peptide secretion of β islet-like cells derived from miR-32-infected HuAECs was much higher than that of cells derived from miR-Mut-infected HuAECs (Fig. 6 and Table III). These findings confirmed that exogenous overexpression of miR-32 could effectively enhance the C-peptide secretion of β islet-like cells.

### Table III. C-peptide release.

| Variable                   | miR-32 HuAECs (pM) | miR-Mut HuAECs (pM) |
|----------------------------|--------------------|--------------------|
| Induced time (weeks)       |                    |                    |
| 0                          | 0.445±0.215        | 0.270±0.270        |
| 1                          | 18.285±4.995       | 1.335±1.225        |
| 2                          | 48.505±4.065       | 1.950±0.930        |
| 3                          | 89.475±9.285       | 1.370±0.490        |
| Glucose concentration (mM) |                    |                    |
| 0                          | 8.400±2.150        | 8.385±2.905        |
| 1                          | 16.650±4.110       | 9.225±1.065        |
| 5                          | 47.015±6.825       | 10.375±1.705       |
| 25                         | 78.865±7.375       | 12.065±1.485       |
| KCl concentration (mM)     |                    |                    |
| 0                          | 1.515±0.495        | 1.590±0.400        |
| 10                         | 2.455±0.405        | 1.315±0.565        |
| 20                         | 5.245±1.235        | 1.640±0.390        |
| 40                         | 9.920±0.670        | 2.035±0.155        |
| Tolbutamide concentration (mM) |                  |                    |
| 0                          | 2.595±0.515        | 3.005±0.065        |
| 50                         | 4.860±0.690        | 3.580±0.590        |
| 100                        | 8.990±0.970        | 2.590±0.520        |

HuAECs, human amniotic epithelial stem cells; miR-32, microRNA-32. *P<0.01 vs. miR-Mut HuAECs; †P<0.05 vs. miR-Mut HuAECs.

Figure 5. miR-32 maintains the ability of HuAECs to differentiate into β islet-like cells. (A) Reverse transcription-quantitative polymerase chain reaction analysis suggested that, after induction, the mRNA expression levels of β islet-like cell biomarkers were increased in miR-32-infected HuAECs compared with in miR-Mut-infected HuAECs (*P<0.01 vs. miR-Mut HuAECs; †P>0.05 vs. miR-Mut HuAECs; Student’s t-test, n=3). (B) Western blot analysis demonstrated that, after induction, the protein expression levels of insulin and PDX1 were increased in miR-32-infected HuAECs compared with in miR-Mut-infected HuAECs (*P<0.01 vs. miR-Mut HuAECs; †P>0.05 vs. miR-Mut HuAECs; Student’s t-test, n=3). GCK, glucokinase; Glut2, glucose transporter 2; HuAECs, human amniotic epithelial stem cells; Isl1, ISL LIM homeobox 1; miR-32, microRNA-32; Mut, mutated; Nkx2.2, NK2 homeobox 2; Pax1, paired box 6; Pdx1, pancreatic duodenal homeobox-1.

**Discussion**

HuAECs are derived from trophoblasts, but retain the characteristics of ESCs (1-3). Although HuAECs are difficult to grow in vitro (they can be grown for a maximum of five passages), they have pluripotent capabilities, which are similar to stem cells (1-3). HuAECs are able to differentiate into numerous human tissues and cells that belong to the three human germ layers, under various induction conditions (1-3). In addition, they possess specific physiological and biochemical functions of adult cells; therefore, they are considered promising candidates for cell therapy (1‑3). However, it is difficult to maintain the pluripotency of HuAECs in vitro (1-3). The present study demonstrated that the expression levels of Oct4, WWP2 and Nanog, which are transcription factors associated with stem cell pluripotency, were markedly decreased with increasing passage number, leading directly to the loss of pluripotency of HuAECs and an inability to induce differentiation into certain adult cells. Therefore, investigating the mechanism underlying the maintenance of stem cell pluripotency may help to improve the culture efficiency of HuAECs and maintain their ‘stemness’. Previous
studies have indicated that the transcription factors associated with pluripotent stem cells serve important regulatory roles in in vitro and in vivo proliferation, the maintenance of pluripotency, and the directed differentiation of stem cells. The present study aimed to determine why the expression level of WWP2 was slowly increased in HuAECs alongside passage number. It has previously been reported that Oct4 activity is regulated by numerous factors (9,10,13,14). At the gene expression level, there are two regulatory pathways: Transcriptional modification and post-transcriptional modification. Generally, in adult cells, the Oct4 gene is inactivated, and epigenetic analyses indicated that the CpG islands in the gene promoter are highly methylated (9,10,13). In addition, binding sites in the promoter and in histones, including H3K27 and H3K9, are modified by methylation and deacetylation, which cause direct downregulation of gene transcription, ultimately affecting gene expression (9,10,13). These modifications are at the transcriptional level (9,10,13). At the post-transcriptional level, some preliminary studies have suggested that endogenous Oct4 protein is degraded in ESCs following prolonged culture in vitro via the main degradation pathway of protein ubiquitination (9,10,13,14). With continued passage of ESCs, WWP2 may be activated and bind to the Oct4 protein, thereby triggering subsequent ubiquitination and degradation, thus leading to loss of Oct4 protein expression and reduced pluripotency of ESCs (9,10,13,14). These results suggested that, in order to maintain Oct4 expression, blocking the expression and activity of WWP2 is crucial (9,10,13,14).

Based on these findings, the present study focused on the regulatory mechanism of WWP2 ubiquitin ligase in HuAECs, in order to provide a novel hypothesis regarding maintenance of the pluripotency of stem cells in vitro. The results confirmed that WWP2 is regulated by endogenous miR-32, particularly in the primary culture stage when miR-32 expression is relatively high. However, with consecutive passages of HuAECs, miR-32 expression was significantly reduced, whereas endogenous WWP2 expression was increased. HuAECs were then induced to overexpress exogenous miR-32 and were compared with HuAECs infected with miR-Mut. The results demonstrated that WWP2 expression in miR-32-infected cells was significantly decreased, whereas the expression of transcription factors associated with pluripotency (Oct4, Sox2 and Nanog) were significantly increased, thus suggesting that miR-32 may significantly inhibit WWP2 expression and promote the expression of pluripotency-associated transcription factors. Furthermore, the results of co-IP demonstrated that, crosslinking between WWP2 and Oct4 proteins was significantly increased in miR-Mut expressed HuAECs but not in miR-32 expressed HuAECs. These findings suggested that ubiquitination of Oct4 was enhanced, whereas miR-32 overexpression in HuAECs significantly inhibited endogenous WWP2 protein expression, and crosslinking between Oct4 and WWP2 was also decreased.

In the present study, HuAECs infected with either miR-32 or miR-Mut were induced to differentiate into β islet-like cells.
Molecular analyses indicated that following directed induction, the expression levels of β islet-like cell markers were significantly increased in miR-32-infected HuAECs compared with in miR-Mut-infected HuAECs. Furthermore, with increasing induction time, C-peptide secretion was also increased from miR-32-infected HuAECs compared with in miR-Mut-infected HuAECs. In addition, C-peptide secretion of the two HuAEC groups were investigated following stimulation with various concentrations of glucose. With increasing glucose concentration in the medium, C-peptide secretion from miR-32-infected HuAECs was significantly increased compared with that secreted by cells derived from miR-Mut-infected HuAECs. Furthermore, with increasing concentrations of KCl in the culture medium, C-peptide secretion from HuAECs. In addition, C-peptide secretion from β islet-like cells derived from miR-32-infected HuAECs was significantly increased compared with that from cells derived from miR-Mut-infected HuAECs. Furthermore, increasing concentrations of tolbutamide (a hypoglycemic drug) were added to the HuAEC groups, which increased C-peptide secretion from β islet-like cells derived from miR-32-infected HuAECs compared with that secreted by β islet-like cells derived from miR-Mut-infected HuAECs. These results confirmed that after WWP2 was silenced by miR-32 overexpression in miR-32-infected HuAECs, the responsiveness to sugar concentrations in the culture medium and C-peptide secretion from β islet-like cells derived from them were significantly increased compared with in cells derived from miR-Mut-infected HuAECs.

In conclusion, miR-32 may prevent the ubiquitination and degradation of Oct4 by inhibiting the expression of WWP2, thus promoting the maintenance of stem cell pluripotency during consecutive passages of HuAECs in vitro.

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Competing interests

The authors declare that they have no competing interests.

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