Abstract. Circular RNAs (circ) have been reported to serve crucial roles in the regulation of cancer occurrence and development. The present study aimed to investigate the role of circ-ACACA in the progression of cervical cancer (CC). The expression levels of circ-ACACA in several CC cell lines were first determined using reverse transcription-quantitative PCR. circ-ACACA expression was subsequently knocked down to evaluate its effects on the viability, proliferation, apoptosis, invasion and migration of CC cells using MTT, colony formation, TUNEL, transwell and wound healing assays, respectively. 13C-labeling of intracellular metabolites and analysis of glucose consumption and lactate production were performed to determine the levels of glycolysis. In addition, the expression levels of endoplasmic reticulum oxidoreductase 1α (ERO1α; ERO1A) and glycolysis-related proteins were analyzed using western blotting. The binding interactions among circ-ACACA, microRNA (miR)-582-5p and ERO1A were validated using dual-luciferase reporter assays. Subsequently, rescue experiments were performed to determine the potential underlying mechanism by which circ-ACACA affected CC cell functions. The results revealed that circ-ACACA expression was significantly upregulated in CC cells and silencing of circ-ACACA significantly reduced the proliferation, invasion and migration, and promoted the apoptosis of CC cells. Knockdown of circ-ACACA markedly inhibited glycolysis in CC cells. However, the effects of silencing of circ-ACACA on CC cells were reversed following transfection with the miR-582-5p inhibitor or pcDNA3.1-ERO1A overexpression plasmid. In conclusion, to the best of our knowledge, the present study was the first to investigate the role of circ-ACACA in CC progression. The results suggested that circ-ACACA may promote CC tumorigenesis and glycolysis by targeting the miR-582-5p/ERO1A signaling axis. Therefore, circ-ACACA may be a promising biomarker for CC diagnosis and treatment.

Introduction

Cervical cancer (CC) is the second most common type of cancer in women aged 20-39 years worldwide, and represents a serious threat to public health (1). Each year, 265,700 individuals die from the disease globally, and thus, efforts have been made to improve the timely diagnosis and effective treatment of CC (2,3). Currently, hysterectomy has become increasingly popular and is widely used as a treatment for early-stage CC (4). Targeted therapies have also become an effective strategy for CC treatment (5).

Circular RNAs (circRNAs/circ), a novel class of non-coding RNAs (ncRNAs), are characterized by a covalently closed loop without a 5' cap and 3' polyadenylated tail, and are generated through abnormal transcription and splicing (6). Accumulating evidence has demonstrated that circRNAs exert effects on multiple biological and pathological processes, suggesting that they may be involved in the occurrence and development of numerous diseases, including cancer (7,8). For example, circ-acetyl-CoA carboxylase α (ACACA) has been previously reported to promote the proliferation, migration and glycolysis of non-small cell lung cancer (NSCLC) cells by modulating the PI3K/AKT signaling pathway (9). However, to the best of our knowledge, the role of circ-ACACA in CC remains to be determined.

miRNAs are small ncRNA molecules that can sponge target genes and regulate their expression (3). A previous study demonstrated that miR-582-5p inhibits the proliferation and invasion of NSCLC cells by downregulating Notch1 expression (10). In addition, bone metastasis of prostate cancer is inhibited by the combined effects of miR-582-3p and miR-582-5p via modulation of TGF-β signaling.
transduction (11). Long ncRNA urothelial cancer associated 1 targets miR-582-5p and promotes the progression and drug resistance of bladder cancer cells through the inhibition of autophagy related 7-mediated autophagy (12). The expression levels of miR-582-5p have also been reported to be down-regulated in endometrial and gastric cancer, and miR-582-5p overexpression inhibits cell proliferation and promotes apoptosis by targeting AKT3 (13). Furthermore, the inhibitory effect of miR-582-5p on the proliferation of colorectal cancer and hepatocellular carcinoma cells has been identified to be achieved by targeting certain genes (14). However, to the best of our knowledge, the role of miR-582-5p in CC has not yet been reported.

Endoplasmic reticulum oxidoreductase 1α (ERO1α; ERO1A) is an oxidase located in the endoplasmic reticulum, which promotes the formation of disulphide bonds in granulocyte-colony stimulating factor (15). Accumulating evidence has demonstrated the close association between upregulated expression levels of ERO1A and poor prognosis in multiple types of cancer, such as pancreatic cancer, cholangiocarcinoma and breast cancer (16-19). In a previous study, knockdown of ERO1A expression reduced the proliferation, migration and tumorigenesis of CC cells by downregulating H₂O₂-associated epithelial-to-mesenchymal transition (20). Additionally, ERO1A has been demonstrated to affect the functions of pancreatic cancer cells by activating the Wnt/β-catenin signaling pathway to enhance the progression of pancreatic cancer (17). A previous study also demonstrated that upregulated expression levels of ERO1A are associated with the poor prognosis of gastric cancer (21). Furthermore, endoplasmic reticulum stress-dependent ERO1A expression has been demonstrated to enhance aerobic glycolysis in pancreatic cancer (16). Hypoxia-inducible ERO1A also promotes the development of colorectal cancer by regulating integrin-β1 and integrin-β1-associated signaling in colorectal cancer cells (22).

Therefore, the present study aimed to investigate the potential role of circ-ACACA in CC and its regulatory effects on the miR-582-5p/ERO1A signaling axis.

Materials and methods

Cell lines and culture. Cervical squamous cell carcinoma cell lines (Ca Ski and SiHa), an endocervical adenocarcinoma cell line (HeLa) and the normal cervical epithelial cell line (End1/E6E7) were obtained from The Cell Bank of Type Culture Collection of The Chinese Academy of Sciences. All cells were cultured in DMEM (Gibco; Thermo Fisher Scientific, Inc.) and maintained in an atmosphere of 37˚C with 5% CO₂.

Cell transfection. pIRES vectors containing short hairpin RNA (shRNA)-circ-ACACA#1, shRNA-circ-ACACA#2, shRNA-negative control (NC), miR-582-5p mimic (50 nM; sense, 5'-UUACAGUUCACAGUUACU-3' and antisense, 5'-UUACUGUUGUGAAACUCUUAU-3'), mimic-NC (50 nM; sense, 5'-UUCUCCGACUGUCAGUTT-3' and antisense, 5'-ACGUGACAGCGUUCGGAGATT-3'), miR-582-5p inhibitor (50 nM; 5'-AGUACUGUGUACACAGUAAUU-3'), NC inhibitor (50 nM; 5'-CAGUACUUUGUAGUACAAA-3'), pcDNA3.1-ERO1A and pcDNA3.1-NC (empty plasmid) were purchased from Shanghai GenePharma Co., Ltd. HeLa cells (1x10⁴ cells per well) were plated and incubated in six-well plates for 24 h and then transfected for 48 h with the corresponding plasmids, oligonucleotides or NCs using Lipofectamine® 2000 reagent (Invitrogen; Thermo Fisher Scientific, Inc.) at 37˚C. The transfected cells were used for subsequent experiments at 48 h after transfection. Untreated cells were used as the control group.

Reverse transcription-quantitative PCR (RT-qPCR). Total RNA was extracted from CC cells using TRIzol® reagent (Invitrogen; Thermo Fisher Scientific, Inc.) according to the manufacturer's protocols. Total RNA was reverse transcribed into cDNA using the PrimeScript™ Strand cDNA synthesis kit (Takara Biotechnology Co., Ltd.). The temperature protocol for this step was as follows: 70˚C for 5 min, 37˚C for 5 min and 42˚C for 1 h. qPCR was subsequently performed using a Power SYBR® Green PCR Master mix (Invitrogen; Thermo Fisher Scientific, Inc.) on an ABI 7500 Real-Time PCR Detection system (Applied Biosystems; Thermo Fisher Scientific, Inc.). The following thermocycling conditions were used: Initial denaturation at 95˚C for 10 min; followed by 40 cycles of denaturation at 95˚C for 15 sec and annealing at 60˚C for 1 min; and a final extension of 10 min at 72˚C. Primers pairs used in this study were as follows: circ-ACACA forward, 5'-GTG GCT TTT G AAG GAG CTG TC-3' and reverse, 5'-CAGA CATGTGGACACCTTGA-3'; miR-582-5p forward, 5'-GGCGGTACAGTTCCACAC-3' and reverse, 5'-CTCACA AACTGTGGTCGGAGA-3'; ERO1A forward, 5'-ATGACATCAGCCATGTGGA-3' and reverse, 5'-CATG CTTGGTTCCACTTGAA-3'; GAPDH forward, 5'-ACA CTTTTGGATCTGGGAAAG-3' and reverse, 5'-GCCATC AGCCACACATTTC-3'; and U6 forward, 5'-GGAAACGGT CACGAGATTGAC-3' and reverse, 5'-TGGAACGC TACGAGATTGCCG-3'. The miRNA expression level was normalized to U6 and GAPDH was applied as the internal control of mRNAs, following quantification using the 2−ΔΔCq method (23).

RNase R treatment assay. Six units of RNase R (Geneseead Biotech, Inc.) were added into every 2 μg RNA and incubated at 37˚C for 20 min. After RNase R treatment, the mRNA expression of circ-ACACA and ACACA was detected by RT-qPCR.

Cell cytoplasm/nucleus fraction isolation. Cell cytoplasm/nucleus fraction isolation was performed using the Nuclear/Cytosol Fractionation Kit (Cell Biolabs, Inc.). Briefly, extracted RNAs from the cytoplasm or nucleus were determined by RT-qPCR which was performed in the previous RT-qPCR section. The relative expression levels of circ-ACACA, ACACA, nuclear control transcript (U6) and cytoplasmic control transcript (β-actin) were measured. β-actin and U6 were used as the internal control for cytosolic and nuclear fractions, respectively.
**MTT assay.** The viability of transfected HeLa cells was detected using MTT reagent (Sigma-Aldrich; Merck KGaA). Briefly, cells (6x10³ cells/well) were plated into 96-well plates and incubated for 24, 48 or 72 h. At each time point, 10 µl MTT was added to each well. Following incubation, 100 µl DMSO (Sigma-Aldrich; Merck KGaA) was added to each well to dissolve the purple formazan crystals. The absorbance of each well was measured at a wavelength of 490 nm using a microplate reader (Bio-Rad Laboratories, Inc.).

**Colony formation assay.** A total of 2x10³ HeLa cells/well were seeded into a six-well plate and maintained in a humidified incubator with 5% CO₂ at 37°C. Following 48 h of incubation, the plates were washed with PBS twice, and then colonies were fixed with 10% paraformaldehyde for 10 min at room temperature and stained with 0.1% crystal violet for 15 min at room temperature. After the plates were washed with phosphate buffer saline (PBS) at room temperature for 1 min and dried, the number of colonies was counted under an inverted light microscope.

**Cell invasion assay.** To detect the ability of cell invasion, Transwell plates with a 8-µm pore insert precoated with Matrigel (BD Biosciences) overnight at 37°C were obtained. A total of 2x10⁴ transfected HeLa cells were trypsinized and plated into the upper chamber of transwell plates with serum-free DMEM. The lower chambers were filled with 800 µl DMEM supplemented with 10% FBS. Following 48 h of incubation at 37°C, the invasive cells were fixed with 95% ethanol for 20 min at 37°C and stained with 0.1% crystal violet for 10 min at 37°C. The number of cells in five randomly selected fields of view was counted under a light microscope.

**Wound healing assay.** HeLa cells were plated into 96-well plates at a density of 4x10³ cells/well and cultured until they reached 80% confluence. The cells were washed with PBS three times and then cultured in medium supplemented with 1% FBS for 24 h. An artificial wound was created across the cell monolayer by scratching the center of all wells with a 10-µl pipette tip. Migratory cells were visualized using an inverted light microscope. ImageJ software (version 1.52r; National Institutes of Health) was used to determine the wound healing rate.

**TUNEL assay.** Following culture for 24 h, 1x10⁴ HeLa cells were seeded in a 6-well plate. Cells were fixed with 4% paraformaldehyde at 4°C for 20 min, permeabilized with 0.1% Triton X-100 in PBS and subsequently incubated with TUNEL reagents (EMD Millipore). Cell nuclei was stained with diamobenzene for 10 min at room temperature. The apoptosis of TUNEL-positive cells was observed under a fluorescence microscope (magnification, x200; Olympus Corporation) and cells were counted in five randomly selected microscopic fields.

**Glycolysis assay.** Glycolysis was determined using a Seahorse XF Glycolytic Rate assay kit (Agilent Technologies, Inc.) on a Seahorse XFe96 analyzer (Agilent Technologies, Inc.) according to the manufacturer’s protocol. Briefly, 2x10⁴ HeLa cells per well transfected with shRNA-circ-ACACA, miR-582-5p, shRNA-circ-ACACA + miR-582-5p inhibitor or respective NCs were plated into a 96-well plate. After the probes were calibrated, 10 mmol glucose, 10 µmol oligomycin and 50 mmol 2-deoxyglucose were serially injected to detect the extracellular acidification rate (ECAR). Data were analyzed using Seahorse XFe24 Wave 2.2 software (Agilent Technologies, Inc.).

**Glucose consumption and lactate production assays.** HeLa cells and cell culture medium were collected following transfection for 48 h. The concentration of glucose in the cell culture medium was determined using a Glucose assay kit (Sigma-Aldrich; Merck KGaA), whereas the lactate concentration was measured using a Lactic Acid assay kit (Seebio), according to the manufacturer’s protocols.

**Measurement of intracellular metabolite generation.** ¹³C-labeled intracellular metabolites were identified as previously described (24). Briefly, 1x10⁴ HeLa cells were incubated with 2 g/l ¹³C-labeled glucose for 2 h at 37°C. Metabolites were subsequently extracted and evaluated using a liquid chromatography system equipped with a TripleTOF® 5600 mass spectrometer (SCIEX; AB SCIEX). Electrospray positive ion mode was used along with the following parameters: ion voltage, 5,500 V; declustering potential, 80 V; source temperature, 60°C; curtain gas, 35 psi; and 100-1,000 m/z.

**Western blotting.** Total protein was extracted from cells using RIPA lysis buffer (Beyotime Institute of Biotechnology). Protein concentration was determined using a BCA kit (Beyotime Institute of Biotechnology) and protein samples (40 µg protein/lane) were separated via 12% SDS-PAGE. The separated proteins were subsequently transferred onto PVDF membranes (EMD Millipore) and the membranes were blocked with 5% skimmed milk for 1.5 h at room temperature. The membranes were then incubated with primary antibodies overnight at 4°C. Following the primary antibody incubation, the membranes were rinsed with TBS-0.2% Tween 20 and incubated with the goat anti-rabbit horseradish peroxidase-conjugated secondary antibody (dilution, 1:2,000; cat. no. ab205718; Abcam) at room temperature for 1 h. GAPDH was used as the internal loading control. Protein bands were visualized using an ECL detection system (Thermo Fisher Scientific, Inc.) and densitometric analysis was performed using ImageJ software (version 1.52r; National Institutes of Health). The following primary antibodies were used: anti-hypoxia-inducible factor 1α (HIF-1A; cat. no. 36169T; 1:1,000; Cell Signaling Technology, Inc.), anti-lactate dehydrogenase A (LDHA; cat. no. 3582T; 1:1,000; Cell Signaling Technology, Inc.), anti-glucose transporter type 1 (GLUT1; cat. no. 12939S; 1:1,000; Cell Signaling Technology, Inc.), anti-hydrogenase A (LDHA; cat. no. 3582T; 1:1,000; Cell Signaling Technology, Inc.), anti-ERO1A (cat. no. 3264T; 1:1,000; Cell Signaling Technology, Inc.), anti-glucose transporter type 1 (GLUT1; cat. no. 12939S; 1:1,000; Cell Signaling Technology, Inc.), anti-ERO1A (cat. no. 3264T; 1:1,000; Cell Signaling Technology, Inc.) and anti-GAPDH (cat. no. 5174T; Cell Signaling Technology, Inc.) and densitometric analysis was performed using ImageJ software (version 1.52r; National Institutes of Health). The following primary antibodies were used: anti-hypoxia-inducible factor 1α (HIF-1A; cat. no. 36169T; 1:1,000; Cell Signaling Technology, Inc.), anti-lactate dehydrogenase A (LDHA; cat. no. 3582T; 1:1,000; Cell Signaling Technology, Inc.), anti-glucose transporter type 1 (GLUT1; cat. no. 12939S; 1:1,000; Cell Signaling Technology, Inc.), anti-ERO1A (cat. no. 3264T; 1:1,000; Cell Signaling Technology, Inc.), anti-GAPDH (cat. no. 5174T; 1:1,000; Cell Signaling Technology, Inc.) and anti-GAPDH (cat. no. 5174T; 1:1,000; Cell Signaling Technology, Inc.) GAPDH was used as an internal control.

**Dual-luciferase reporter assay.** The binding site between circ-ACACA and miR-582-5p as well as miR-582-5p and ERO1A was predicted using the Encyclopedia of RNA Interactomes.
To validate the binding relationship between miR-582-5p and circ-ACACA or ERO1A, wild-type (WT) or mutant (MUT) 3’-untranslated region sequences of circ-ACACA or ERO1A were cloned into a pmiRGLO dual-luciferase miRNA target expression reporter (Promega Corporation). HeLa cells were subsequently transfected with miR-582-5p mimic or mimic-NC and WT or MUT reporters using Lipofectamine® 2000 reagent (Invitrogen; Thermo Fisher Scientific, Inc.). The relative luciferase activity was measured at 48 h post-transfection using a Dual Luciferase Reporter assay system (Promega Corporation). And firefly luciferase activity was normalized to Renilla luciferase activity.

Statistical analysis. All experiments were repeated independently in triplicate. Statistical analysis was performed using SPSS v20.0 software (IBM Corp.). Data are presented as the mean ± SD. Statistical differences between two groups were determined using an unpaired Student’s t-test, while one-way ANOVA followed by Tukey’s post hoc test was used for comparisons among ≥3 groups. P<0.05 was considered to indicate a statistically significant difference.

Results
circ-ACACA expression is upregulated in CC cells. To investigate the role of circ-ACACA in CC, its expression levels in CC cells were first determined using RT-qPCR. The results revealed that the mRNA expression levels of circ-ACACA were upregulated in CC cells compared with End1/E6E7 cells (Fig. 1A). The expression levels of circ-ACACA were the highest in HeLa cells among the cell lines examined. Therefore, HeLa cells were used as the CC cell model in subsequent experiments. As shown in Fig. 1B, circ-ACACA was not dissolved, while ACACA mRNA was dissolved by RNase R. Furthermore, circ-ACACA was expressed at a higher level in the cytoplasm compared with the nucleus (Fig. 1C).

These results suggested that circ-ACACA expression may be upregulated in CC cells.

Silencing of circ-ACACA inhibits the proliferation, invasion and migration, while promoting the apoptosis of CC cells. circ-ACACA expression was subsequently knocked down to determine its effects in CC cells. The expression levels of circ-ACACA were downregulated following transfection with shRNA-circ-ACACA#1 and shRNA-circ-ACACA#2 compared with shRNA-NC, especially in shRNA-circ-ACACA#2. Therefore, shRNA-circ-ACACA#2 was selected for use in subsequent experiments (Fig. 2A). The effects of transfection of shRNA-circ-ACACA on CC cells were determined using functional assays. MTT and colony formation assays demonstrated that cell viability and proliferation were decreased in HeLa cells transfected with shRNA-circ-ACACA as compared with the shRNA-NC group (Fig. 2B and C). Furthermore, transfection with shRNA-circ-ACACA increased the levels of apoptosis compared with the shRNA-NC group (Fig. 2D and E). circ-ACACA silencing also notably inhibited cell invasion and migration compared with the shRNA-NC group (Fig. 2F-I). These results suggested that silencing of circ-ACACA may inhibit the proliferation, invasion and migration, and promote the apoptosis of CC cells.

Silencing of circ-ACACA suppresses glycolysis in CC cells. It was subsequently investigated whether knockdown of circ-ACACA led to the attenuation of glycolysis. As displayed in Fig. 3A, ECAR was notably decreased after transfection with shRNA-circ-ACACA when compared with the shRNA-NC group. Lactate production and glucose consumption assays were performed to evaluate the effect of circ-ACACA on glycolysis. As shown in Fig. 3B and C, the transfection with shRNA-circ-ACACA markedly decreased the production of lactate and increased the levels of glucose compared with the shRNA-NC group. In addition, HeLa cells were incubated with 13C-labeled glucose and liquid chromatography-mass
spectrometry was performed to gain more in-depth insights into the metabolic flux of glucose. As shown in Fig. 3D, the ratio of 13C-marked glucose was notably decreased following circ-ACACA-knockdown relative to the shRNA-NC group. As expected, the expression levels of glycolysis-related proteins, including HIF-1A, LDHA, GLUT1 and HK2, were markedly downregulated following the knockdown of circ-ACACA compared with those in the shRNA-NC group (Fig. 3E). These
results indicated that silencing of circ-ACACA may inhibit glycolysis in CC cells.

**circ-ACACA negatively regulates the expression levels of miR-582-5p.** To determine the regulatory effect of circ-ACACA on the expression levels of miR-582-5p, the expression levels of miR-582-5p were first analyzed in various CC cell lines. The results revealed that miR-582-5p expression was downregulated in CC cells compared with End1/E6E7 cells. **Figure 3.** Silencing of circ-ACACA suppresses glycolysis in cervical cancer cells. (A) ECAR after HeLa cells were transfected with shRNA-circ-ACACA. ***P<0.001 vs. shRNA-NC. (B) Lactate levels and (C) glucose content after HeLa cells were transfected with shRNA-circ-ACACA. (D) Metabolite analysis and (E) glycolysis-related protein expression after HeLa cells were transfected with shRNA-circ-ACACA. *P<0.01, ***P<0.001. circ-ACACA, circular RNA acetyl-CoA-carboxylase α; ECAR, extracellular acidification rate; GLUT1, glucose transporter type 1; HIF1A, hypoxia-inducible factor 1α; HK2, hexokinase-2; LDHA, lactate dehydrogenase A; NC, negative control; shRNA, short hairpin RNA; OM, oligomycin; 2-DG, 2-deoxy-D-glucose.
miR-582-5p by transfection with the miR-582-5p mimic, a dual-luciferase reporter assay was used to demonstrate the binding relationship between circ-ACACA and miR-582-5p (Fig. 4C and D). As shown in Fig. 4E, knockdown of circ-ACACA markedly upregulated the expression levels of miR-582-5p compared with those in the shRNA-NC group. These data suggested that circ-ACACA may negatively regulate miR-582-5p expression in HeLa cells.

ERO1A is a target gene of miR-582-5p. It was hypothesized that circ-ACACA may exert its promoting effects on the progression of CC via the miR-582-5p/ERO1A signaling axis. To verify this hypothesis, the expression levels of ERO1A in various CC cell lines were determined. As illustrated in Fig. 5A and B, ERO1A expression was notably unregulated in CC cells compared with End1/E6E7 cells. Since ERO1A was predicted to target and bind with miR-582-5p using ENCORI database. A binding association between ERO1A and miR-582-5p was predicted using ENCORI (Fig. 5C) and verified using a dual-luciferase reporter assay (Fig. 5D). Subsequently, RT-qPCR and western blotting demonstrated that transfection with the miR-582-5p mimic markedly reduced the mRNA and protein expression levels of ERO1A compared with the shRNA-NC group (Fig. 5G and H). These findings suggested that ERO1A may be a target of miR-582-5p, and that circ-ACACA may upregulate the expression levels of ERO1A by sponging and negatively regulating miR-582-5p expression.
Knockdown of miR-582-5p or overexpression of ERO1A reverses the effects of circ-ACACA silencing on CC cell proliferation, viability, invasion, and migration. To investigate whether circ-ACACA could affect the functions of CC cells in vitro via the miR-582-5p/ERO1A signaling axis, ERO1A was overexpressed in CC cells. The expression levels of ERO1A were examined by western blotting and RT-qPCR after pcDNA3.1-ERO1A was transfected into HeLa cells. Expression levels of ERO1A were examined by western blotting and RT-qPCR after pcDNA3.1-ERO1A was transfected into HeLa cells. (A) Expression levels of ERO1A were examined by western blotting and RT-qPCR after pcDNA3.1-ERO1A was transfected into HeLa cells. (B) miR-582-5p expression was evaluated by RT-qPCR after transfection with miR-582-5p inhibitor. **P<0.01, ***P<0.001. (C) Cell viability was detected using an MTT assay. *P<0.05, ***P<0.001 vs. shRNA-NC; ###P<0.001 vs. shRNA circ-ACACA+NC inhibitor; ΔΔΔP<0.001 vs. shRNA circ-ACACA+pcDNA 3.1-NC. (D) Cell proliferation was determined using a colony formation assay. (E) Cell apoptosis was evaluated using TUNEL staining. Magnification, x200. (F) The apoptosis rate of TUNEL assay. (G) Cell invasion was evaluated using Transwell assay. (H) The number of invaded cells. Magnification, x100. (I) Cell migration was assessed by wound healing assay. (K) The cell migration rate. Magnification, x100. "P<0.001. circ-ACACA, circular RNA acetyl-CoA-carboxylase α; ERO1A, endoplasmic reticulum disulphide oxidase 1α; miR, microRNA; NC, negative control; OD, optical density; RT-qPCR, reverse transcription-quantitative PCR; shRNA, short hairpin RNA.
and these effects were subsequently reversed by transfection with the miR-582-5p inhibitor or pcDNA3.1-ERO1A (Fig. 7A). Furthermore, the results of the lactate production and glucose consumption assays revealed that the decreased lactate production and enhanced glucose levels induced by transfection with shRNA-circ-ACACA were partially recovered following the transfection with the miR-582-5p inhibitor or pcDNA3.1-ERO1A (Fig. 7B and C). Finally, the shRNA-circ-ACACA-induced decreases in the ratio of 13C-marked glucose and expression levels of glycolysis-related proteins were partially reversed following transfection with the miR-582-5p inhibitor or pcDNA3.1-ERO1A (Fig. 7D and E). Overall, these results suggested that knockdown of miR-582-5p or overexpression of ERO1A alleviates the effects of circ-ACACA silencing on CC cell proliferation, viability, invasion, migration and apoptosis.

Discussion
ncRNAs have been suggested to serve important roles in the progression of numerous types of tumor, such as prostate, non-small cell lung, colorectal and cervical cancer (25-29). circRNAs, which are single-stranded ncRNAs, have been revealed to be involved in the development of multiple cancer types (30). Due to the increased stability, evolutionary conservation and high abundance of circRNAs, they have been established as essential players in numerous physiological and pathophysiological processes (31). Previous research has demonstrated that circ-ACACA expression is upregulated in NSCLC tissues and cells, and silencing of circ-ACACA hinders the cellular functions of NSCLC cells, including proliferation, invasion and migration, via regulation of the miR-1183 and PI3K/AKT signaling pathway (9). Therefore,
it was hypothesized that circ-ACACA may also serve as an oncogene to promote the tumorigenesis of CC. The results of the present study revealed that circ-ACACA expression was significantly upregulated in CC cells, and circ-ACACA was expressed at a higher level in the cytoplasm compared with the nucleus. Silencing of circ-ACACA inhibited the viability, proliferation, invasion and migration, and enhanced the apoptosis of CC cells.

Alteration of energy metabolism, especially abnormal activation of the glycolysis pathway has been observed in diverse human cancer types, including CC (32-36). Cancer cells are dependent on the glycolytic pathway to meet their high energy demands; thus, increased glycolysis is a hallmark of cancer cells (37). Notably, knockdown of circ-ACACA could hinder proliferation and migration of NSCLC cells and also reduce the glycolysis rate (9). In the present study, the effect of circ-ACACA on glycolysis in CC cells was determined. Higher lactate levels have been previously identified to increase the risk of recurrence and metastasis in patients with CC, and are associated with poor overall survival (38). Furthermore, high lactate content in the tumor microenvironment has been revealed to induce highly acidic conditions, which hinders the function of several chemotherapeutic drugs, such as tamoxifen, cisplatin and adriamycin (39,40).

In the present study, silencing of circ-ACACA reduced the ECAR and lactate levels. Increased glucose uptake is another characteristic of cancer cells (37), and the present study revealed that glucose uptake was decreased in HeLa cells transfected with shRNA-circ-ACACA. HIF-1A, LDHA, GLUT1 and HK2 serve important roles in glycolytic metabolism (39). The present study revealed that transfection with shRNA-circ-ACACA downregulated the expression levels of these factors, indicating that shRNA-circ-ACACA may inhibit glycolysis. Therefore, these results indicated the suppressive effect of silencing of circ-ACACA on glycolysis and the progression of CC.

The unique cellular stability and ability of circRNAs to sponge miRNAs and proteins suggests their potential as novel molecular markers for targeted therapies for cancer (40). miRNAs are known to serve an important role in the pathogenesis of most types of cancer (41). Efforts have been made to determine the effect of miR-582 on cancer development (42). A previous study reported that miR-582-5p is differentially expressed during the development of multiple types of cancer, indicating that the expression levels of miR-582-5p depend on the specific type of cancer tissue or cells (42). The results of the present study demonstrated that the expression levels of miR-582-5p were downregulated in CC cells, and dual-luciferase reporter assays verified the binding between circ-ACACA and miR-582-5p. RT-qPCR analysis revealed that the silencing of circ-ACACA markedly upregulated the expression levels of miR-582-5p, demonstrating the negative association between circ-ACACA and miR-582-5p.

circRNAs have been reported to act as miRNA sponges in circRNA/miRNA/mRNA signaling axes (43). In the present study, bioinformatics analysis was used to predict that ERO1A could interact with miR-582-5p, which prompted further investigations to determine the role of ERO1A in CC cells. ERO1A, which is a protein involved in the oxidative protein folding of molecules involved in cancer progression, has been reported to be induced by hypoxia in HeLa cells (44). The expression levels of ERO1A have been reported to be upregulated in numerous types of cancer cells compared with normal cells, such as breast, pancreatic and colorectal cancer (19,17,22), which was consistent with the present findings that CC cells exhibited upregulated ERO1A expression compared with End1/E6E7 cells. Notably, transfection with the miR-582-5p mimic downregulated the expression levels of ERO1A in HeLa cells. Whether circ-ACACA affected the functions of HeLa cells via regulation of the miR-582-5p/ERO1A signalling axis was subsequently investigated. The results demonstrated that transfection with shRNA-circ-ACACA decreased the proliferation, invasion and migration, and promoted the apoptosis of HeLa cells, while these effects were reversed following transfection with the miR-582-5p inhibitor or pcDNA3.1-ERO1A.

In conclusion, to the best of our knowledge, the present study was the first to investigate the role of circ-ACACA in CC development and provided evidence to suggest that the knockdown of circ-ACACA may decrease proliferation, invasion, migration and glycolysis, and promote apoptosis in HeLa cells via targeting of the miR-582-5p/ERO1A signalling axis. These results highlight the potential of circ-ACACA as a novel biomarker for CC. However, the lack of experiments regarding the clinical value of circ-ACACA/miR-582-5p/ERO1A in CC tissue samples to determine its clinical applicability, as well as in vivo anti-tumor effects in an animal model are limitations of the present study. Additionally, based on a previous study in NSCLC (9), the circ-ACACA/miR-1183/PI3K/protein kinase B signaling pathway and whether there is a direct interaction between circ-ACACA and miR-1183 should also be explored in CC in future studies. Therefore, comprehensive analysis is required in the future.

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Availability of data and materials

All data generated or analyzed during this study are included in this published article.

Authors' contributions

DH and CL interpreted the data and performed experiments. DH collected the data, searched the literature, designed the study and wrote the manuscript. CL revised the manuscript. Both authors read and approval the final manuscript. DH and CL confirm the authenticity of all the raw data.

Ethics approval and consent to participate

Not applicable.
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