Characterization and identification of human immortalized granulosa cells derived from ovarian follicular fluid

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Abstract. Follicular fluid serves a crucial role in follicular development and oocyte maturation. Increasing evidence indicates that follicular fluid is rich in proteins and functional cells. In addition to oocyte cells, follicular fluid contains granulosa, thecal and ovarian surface epithelial cells. Granulosa cells (GCs) represent the predominant somatic cell type of the ovarian follicle and are involved in steroidogenesis and folliculogenesis. However, the long-term culture of GCs in vitro remains challenging. The present study aimed to extend the culture of GCs in vitro. Human GCs were collected from the follicular fluid of patients included in an in vitro fertilization program and cultured in the presence of conditioned medium obtained from mouse embryonic fibroblasts. GCs were cultured for over a year and 130 passages, and the population doubling time was ~22 h. Cells presented epithelial-like morphology and a cobblestone-like appearance when they reached confluence. Flow cytometric analysis demonstrated that cells expressed CD29, CD166 and CD49f but not CD31, CD34, CD45, CD90, CD105 or CD13. Immunofluorescence staining revealed that cells expressed follicle stimulating hormone receptor, luteinizing hormone receptor and cytochrome P450 aromatase, which was confirmed by reverse transcription-quantitative polymerase chain reaction. In the presence of androstenedione, cells secreted estradiol. In addition, estradiol level was further stimulated by dibutylryl cAMP treatment. In addition, intracellular cAMP and progesterone expression levels were upregulated by follicle stimulating hormone and/or human chorionic gonadotropin. Furthermore, cells survived in severe combined immunodeficiency mice following intra-ovarian injection. Histological analysis revealed that certain cells formed follicle-like structures. The results from the present study suggested that immortalized GCs may be a useful tool for further research on GC and improve the clinical application of drugs such as follicle-stimulating hormone or human chorionic gonadotropin.

Introduction

The development of ovarian follicles begins with the proliferation of granulosa cells (GCs), which change shape from flat to cubical and from single- to multi-layered cells (1,2). At the late stage of preantral follicle development, follicle-stimulating hormone receptor (FSHR) appears in GC membranes. The activity of GCs is regulated by follicle-stimulating hormone (FSH) and luteinizing hormone (LH) (3). The preovulatory follicle comprises follicular fluid, an oocyte, hundreds to thousands of GCs and two layers of surrounding theca cells. GCs of the preovulatory ovarian follicle represent a predominant somatic cell type of the ovarian follicle. GCs are responsible for the communication between the oocyte and the theca cells, which are regulated by the gonadotropins FSH and LH, produce estrogen and nurse the oocyte. Following ovulation, GCs become luteinized GCs, which produce large amounts of progesterone to support the corpus luteum and subsequent pregnancy (4).

Luteinized GCs have long been considered as terminally differentiated cells with a limited life span (5). Previous studies demonstrated that GCs present certain characteristics of stem cells (6-8). Lavranos et al (6) reported that bovine luteinized GCs are able to grow in colonies and retain the ultrastructural features of follicular GCs in an anchorage-independent culture system. Van Deerlin et al (7) revealed that luteinized GCs from a follicle are derived from a small number
of stem cells. Furthermore, Lavrans et al (8) demonstrated that the telomerase activity in the ovary originated mainly from preovulatory GCs and not the ovarian follicle oocyte, which supported the hypothesis that GCs could originate from a stem cell population. Additional studies demonstrated that GCs possess a multipotent differentiation capacity. Bukovsky et al (9) reported that porcine preovulatory GCs can convert into neural stem cells and differentiate into neurons. In addition, Kossowska-Tomaszcuk et al (3) demonstrated that luteinized GCs cultured in vitro can differentiate into neurons, chondrocytes and osteoblasts. These studies confirmed the presence of granulosa stem cells in the ovary (10-13).

Culture and expanding granulosa stem cells in vitro may be useful for basic research and clinical applications; however, cultivating human GCs in vitro remains a major challenge. Several attempts have been made to prolong the lifespan of GCs in culture, although these methods had limited success. These attempts included creating a three-dimensional culture system (14), supplementing follicular fluid into culture media (15), or adding growth factors into culture media (3). Mouse embryonic fibroblasts (MEFs) usually serve as a feeder cell layer for mouse and human embryonic stem cell cultures and can provide a suitable environment containing growth factors, cytokines and allowing cell-cell interactions, which maintain cells in an undifferentiated state (16-18). MEF-conditioned culture medium may therefore be helpful to support the growth of GCs in vitro. To test this hypothesis, follicular fluid from infertile women who underwent oocyte retrieval during in vitro fertilization (IVF) was collected. Cells from the follicular fluid were cultured in the presence of MEF-conditioned medium. Immortalized ovarian GCs were purified following >1 year culture. In addition, the phenotypic and functional features of the GCs were characterized.

Materials and methods

Collection of follicular fluid cells. Follicular fluid cells were collected from 109 infertile women who underwent IVF at the Department of Assisted Reproduction of the Ninth People's Hospital of Shanghai Jiaotong University School of Medicine between March 2013 and December 2016. The study included women aged between 22 and 32 years with tubal, unexplained or male factors of infertility, and excluded women who had ovarian cyst or tumor. Patients were first treated with controlled ovarian stimulation drugs such as 150-225 IU human menotropin (Anhui Fengyuan Pharmaceutical Co., Ltd.), 10 mg medroxyprogesterone acetate (Zhejiang Xianju Pharmaceutical Co., Ltd.) or 0.1 mg triptorelin (Decapeptyl Ltd.), 10 mg medroxyprogesterone acetate (Zhejiang Xianju Pharmaceutical Co., Ltd.) or 0.1 mg triptorelin (Decapeptyl®; Ferring Pharmaceuticals) per day for 7 to 10 days. Next, ovulation was triggered with 0.1 mg triptorelin and/or 2,000-5,000 IU human chorionic gonadotropin (Lizhu Pharmaceutical Trading Co., Ltd.). After 34-38 h of trigger administration, all oocytes in follicles with diameters >10 mm were retrieved by transvaginal ultrasound-guided aspiration (19).

Following the removal of the oocyte-corona-cumulus complexes, the fresh follicular fluid was centrifuged for 5 min at 524 x g and 4°C. Next, the cell pellet was collected and washed three times with PBS. Following this step, cells were isolated by density gradient centrifugation with 2 ml Percoll (Santa Cruz Biotechnology, Inc.) for 30 min at 2,095 x g and 4°C. Cells in the interphase layer were collected and washed three times with Dulbecco's-modified Eagle's medium (DMEM; Invitrogen; Thermo Fisher Scientific, Inc.). Cells were placed on 100-mm² tissue culture dishes in the presence of MEF-conditioned medium as described below. To obtain enough cells, follicular fluid from 5 to 10 patients was pooled into one primary cell culture.

The ovarian tissue sample was obtained from one female patient who underwent laparoscopic excision of dermoid cysts at the Department of Gynecology, Shanghai Ninth People's Hospital. The patient was 22 years old and the recruitment/collection date was February 2016.

The BMSCs were obtained from one male patient who suffered from femoral head necrosis and underwent BMSCs implantation procedures at the Department of Orthopedics, Shanghai Ninth People's Hospital. He was 23 years old and the recruitment/collection date was March 2016.

Human chondrocytes were obtained from a patient who underwent rhinoplasty procedures at the Department of Plastic and Reconstructive Surgery, Shanghai Ninth People's Hospital (20). She was a 27-year old female and the recruitment/collection date was April 2016.

Fresh follicle cells (FCs) were obtained from female patients who underwent IVF procedures at the Department of Assisted Reproduction, Shanghai Ninth People's Hospital. Their age distribution was between 24 and 32 years old (the median age was 29 years old). The date of recruitment/sample collection was between December 2015 and April 2016. The exclusion criteria were women with ovarian cysts or tumors.

The study was approved by the local Ethics Committee of the Ninth People's Hospital of Shanghai. All participants, including patients undergoing IVF, and patients from whom ovarian tissue samples, bone marrow mesenchymal stem cells (BMSCs), chondrocytes and fresh FCs were obtained, provided written informed consent.

Cell culture. Follicular fluid cells were cultured in a mixed culture medium that consisted of 70% basic culture medium and 30% MEF-conditioned medium. The basic culture medium comprised DMEM containing 4.5 g/l glucose (Gibco; Thermo Fisher Scientific, Inc.) and supplemented with 10% fetal calf serum (FCS; Gibco; Thermo Fisher Scientific, Inc.), 50 g/ml penicillin/streptomycin (Gibco; Thermo Fisher Scientific, Inc.), 3 mmol/l L-glutamine (Sigma-Aldrich; Merck KGaA) and 10 mM β-mercaptoethanol (Sigma-Aldrich; Merck KGaA). Cells were seeded at a density of 1.5x10⁶ cells/100 mm² bacterial-culture dish and incubated at 37°C and 5% CO₂. Medium was exchanged after 24 h and cells were passaged every 3-4 days. Images of primary passages 1, 2, 4, 50 and 99 were captured under an inverted light microscope (magnification, x100; Vert.A1; Zeiss GmbH).

The BMSCs were suspended in α-MEM (alpha-Minimum Essential Medium) containing 10% fetal bovine serum (Hyclone; GE Healthcare Life Sciences), 50 mg/ml sodium ascorbate (Sigma-Aldrich; Merck KGaA), antibiotic/antimycotic and 10⁻⁵ M dexamethasone (Sigma-Aldrich; Merck KGaA) and then seeded at a density of 1.5x10⁶ cells/100 mm² bacterial-culture dish and incubated at 37°C and 5% CO₂. Medium was exchanged every 2-3 days. BMSCs were passaged every 5-7 days.

Fresh FCs were cultured in DMEM containing 4.5 g/l glucose (Gibco; Thermo Fisher Scientific, Inc.) and
supplemented with 10% fetal calf serum (FCS; Gibco; Thermo Fisher Scientific, Inc.), 50 g/ml penicillin/streptomycin (Gibco; Thermo Fisher Scientific, Inc.). Medium was exchanged after 24 h and changed every 2 days.

MEFs were prepared in our laboratory as previously described (21). Briefly, MEFs were seeded at a density of 1x10^6 cells/100 mm² bacterial-culture dish and then cultured with DMEM medium supplemented with 10% FBS and 50 g/ml penicillin/streptomycin at 37°C and 5% CO₂. The MEFs were passaged every 3-4 days. From passages 2-5, the supernatant of MEFs was collected and centrifuged at 524 x g at 37°C for 5 min every 48 h to produce conditioned medium. The conditioned medium was then sterilized with 0.22-µm filters and frozen at -20°C until further use. Cells, including follicular fluid cells, BMSCs and MEFs, were passaged using 0.1% trypsin/EDTA (Invitrogen; Thermo Fisher Scientific, Inc.) once they reached 80% confluence.

**Growth curve of human GCs.** Cell counting was performed from passage 4 up to passage 34 with a hemocytometer. The population doubling time (DT) was determined according to the following formula: DT= t x [lg2/(lgNt-lgN0)] (15) where ‘t’ is the time between cell seeding and harvesting, ‘N0’ is the number of cells seeded and ‘Nt’ is the number of cells following culture. This determination was performed in three dishes for each experiment.

**Colonies formation assay.** Colony formation assays were performed as previously described (22). Briefly, cells at passage 8 were seeded at the density of 100 single cells per plate in triplicate. Following 20 days culture, cells were stained with crystal violet and the number of clones was counted by naked eye. The images of a large colony derived from a single cell was observed on days 1, 2, 3, 6, 12 and 15 under an inverted light microscope (magnification, x40; Vert.A1; Zeiss GmbH). The colony formation efficiency was calculated from three repeated experiments. The efficiency was calculated as mean ± SD.

**Flow cytometric analysis.** Cells at passages 10 and 30 were harvested for flow cytometric analysis. Prior to staining with the fluorescence conjugated antibodies, cells were incubated with 2.5 µg/test of Human BD Fc BlockTM at 4°C for 10 min (cat. no. 564219). Cells were incubated with phycoerythrin-conjugated CD13 (cat. no. 560998; 1:200 dilution), CD29 (cat. no. 556049; 1:200 dilution), CD31 (cat. no. 554061; 1:200 dilution), CD34 (cat. no. 550761; 1:200 dilution), CD45 (cat. no. 555755; 1:200 dilution), CD49f (cat. no. 555736; 1:200 dilution), CD90 (cat. no. 555596; 1:200 dilution), CD105 (cat. no. 560839; 1:200 dilution), CD166 (cat. no. 560903; 1:200 dilution) and human leukocyte antigen-ABC (HLA-ABC) antibodies (cat. no. 560168; 1:200 dilution) for 30 min at 4°C (antibodies were all from BD Biosciences). Cells were washed three times with PBS and analyzed with a flow cytometer (Epics Altra; Beckman Coulter, Inc.). Flow cytometry data were analyzed with CXP software (EXPO32 v1.2; Beckman Coulter, Inc.).

**Immunofluorescence staining.** FCs and BMSC at passages 10 and 60 were fixed with 4% cold paraformaldehyde (Sigma-Aldrich; Merck KGaA) in PBS for 15 min at 4°C and permeabilized with 0.25% Triton X-100 (Sigma-Aldrich; Merck KGaA) in PBS for 10 min. Fixed cells were blocked for 30 min at 37°C with PBS containing 1% bovine serum albumin (BSA; Sigma-Aldrich; Merck KGaA) and 10% goat serum (Sigma-Aldrich; Merck KGaA), and incubated with goat anti-human FSHR (cat. no. sc7798; 1:100), rabbit anti-human luteinizing hormone receptor (LHR; cat. no. sc25828; 1:100) or mouse anti-human cytomegalo virus P450 aromatase (CYP19A; cat. no. sc374176; 1:100; all from Santa Cruz Biotechnology, Inc.) antibodies at 4°C overnight. Cells were then incubated with secondary antibodies labelled with Alexa Fluor 555 for 30 min at 37°C. Specifically, the donkey anti-goat IgG (1:1,000; cat no: A21432) was used for binding the primary antibody goat anti-human FSHR, goat anti-rabbit IgG (1:1,000; cat no: A21428) was used for binding the primary antibody rabbit anti-human luteinizing hormone receptor and goat anti-mouse IgG (1:1,000; cat no: A21422) were used for binding the primary antibody mouse anti-human cytochrome P450 aromatase CYP19A. All secondary antibodies were purchased form Invitrogen (Thermo Fisher Scientific, Inc.). Nuclei were counterstained with DAPI for 5 min and 25°C (Invitrogen; Thermo Fisher Scientific, Inc.). Human BMSCs, which were isolated and expanded as previously described (23), served as a negative control.

**Reverse transcription (RT)-semi-quantitative PCR.** Human ovarian tissue and FCs served as positive controls while chondrocytes served as negative control. Total RNA was extracted from ovarian tissue, FCs, GCs at passage 10 or passage 60 and chondrocytes using TRIzol® (Invitrogen; Thermo Fisher Scientific, Inc.). Ovarian tissue was dissected into 5x5x5 mm samples and quickly placed into the sterile culture tube containing TRIzol® (1.5 ml) for 15 min then RNA was extracted from the homogenized solution. RT-PCR was performed using an RT-PCR kit (TakaRa Taq™ and Reverse Transcriptase XL (AMV); Takara Bio, Inc.) according to the manufacturer’s instructions. The temperature protocol for RT was as follows: 30°C for 10 min, 42°C for 60 min, 99°C for 5 min and 5°C for 5 min. The thermocycling conditions for PCR was as follow: 95°C for 3 min, 95°C for 30 sec, 60-62°C for 30 sec, 72°C for 30 sec and 29-34 cycles, 72°C for 10 min. The sequences of the analyzed genes were searched in the GenBank from the National Centre for Biotechnology Information (www.ncbi.nlm.nih.gov/genbank). Primers were synthesized by Sangon Biotech Co., Ltd. The sequences of the primers are listed in Table I. DNA polymerase used was TakaRa Taq™ (Takara Bio, Inc.) and reverse transcriptase used was Reverse Transcriptase XL (AMV) (Takara Bio, Inc.). In the procedure of semi-quantitative PCR, the thermocycling conditions were as following: FSHR at 60°C for 34 cycles; LHR at 60°C for 34 cycles; CYP19A at 62°C for 34 cycles; estrogen receptor-α (ER-α), 60°C, 29 cycles; estrogen receptor-β (ER-β), 60°C, 29 cycles; progesterone receptor (PR) at 60°C for 29 cycles and androgen receptor (AR) at 60°C for 29 cycles. The amplified products were separated on 1.2% agarose gels and visualized with ethidium bromide (Sigma-Aldrich; Merck KGaA).

**Measurement of intracellular cyclic adenosine-monophosphate (cAMP).** Intracellular cAMP following stimulation with recombinant human FSH (rhFSH; Gonal-f®;
cells were lysed and released their intracellular components. For the measurement of intracellular cAMP, cells were further cultured for 3 h in the presence or absence of 1.0% (w/v) BSA (Sigma-Aldrich; Merck KGaA), 0.5 mM 3-isobutyl-1-methylxanthine (Sigma-Aldrich; Merck KGaA) and 10⁻⁵ M forskolin (Sigma-Aldrich; Merck KGaA). The cells were treated with 10 μM androstenedione (4-androstene-3,17-dione; A₅; Jinchun Biochem Inc., Shanghai, China) and stimulated with 5 IU/ml rhFSH, 100 IU/ml HCG, 10⁻⁴ M dibutyryl cAMP [Bu₂-cAMP; Enzo Life Sciences] or 10⁻³ M forskolin for 48 h. The estradiol and progesterone levels in the culture medium were determined with the chemiluminescent immunoassay i2000SR (ARCHITECT®, Abbott Pharmaceutical Co. Ltd.). All experiments were conducted in triplicate.

**Measurement of the secreted steroid content in cell medium.** To measure the concentrations of estradiol and progesterone in the medium, cells were cultured in DMEM containing 5% charcoal-stripped FCS (GeneTex, Inc.) for 24 h at 37°C. Subsequently, cells were treated with 10 μM androstenedione (4-androstene-3,17-dione; A₅; Jinchun Biochem Inc., Shanghai, China) and stimulated with 5 IU/ml rhFSH, 100 IU/ml HCG, 10⁻⁴ M dibutyryl cAMP [Bu₂-cAMP; Enzo Life Sciences] or 10⁻³ M forskolin for 48 h. The estradiol and progesterone levels in the culture medium were determined with the chemiluminescent immunoassay i2000SR (ARCHITECT®, Abbott Pharmaceutical Co. Ltd.). All experiments were conducted in triplicate.

**Cell labeling.** Cells at passage 8 were labelled by transfection with an enhanced green fluorescent protein (eGFP) sequence using a lentivirus vector (pNL-EGFP/CMV/WPRE/ΔU3; SunBio, Inc.; http://www.sbo-bio.com.cn) according to the manufacturer's protocol. The cells at passages 8 were seeded into a 24-well plate at a density of 3.6x10⁴ cells/ml. After incubation for 24 h, the culture medium was removed, and the virus-containing medium was added into the cells at a multiplicity of infection (MOI) of 10. A total of 1.8 μl of 2X 10⁵ TU/μl lentiviral particles for transfecting 3.6x10⁴ cells, so the MOI was calculated as 10. Following 48-h of transfection, puromycin selection was performed. The viral solution was replaced with 1 mL DMEM medium with 1 μl Puromycin (2 mg/ml, SunBio, Inc.). Following incubation for 24 h, puromycin-containing medium was replaced with DMEM medium. The labeling efficiency was directly observed under a fluorescence inverted microscope (Vert.A1; Zeiss GmbH).

**Intra-ovarian transplantation.** eGFP-labelled cells were suspended in DMEM at the density of 1x10⁵ cells/ml. A total of 1x10⁵ cells were injected into the right ovary of 22 7-week-old female severe combined immunodeficiency (SCID) mice (Shanghai Sipuer-Bik Laboratory Animal Co., Ltd.) using a microinjector (Microsyringe). Mice were about 15-20 g and were housed at 21±2°C in a humidified atmosphere (40-70%) with 12-h light/dark cycles with free access to food and water. Meanwhile, 1x10⁵ cells were subcutaneously injected into two sides of the dorsal region of 4 SCID mice and 1x10⁵ cells were peritoneal cavity of 4 SCID mice. Animals were sacrificed by cervical dislocation. Ovarian tissues were harvested at 8 weeks post-transplantation. Ovarian tissues were fixed in 4% paraformaldehyde for 24 h at room temperature, then placed in freezing microtome, embedded at -25°C with an OCT embedding agent, the frozen section thickness was 6 μm. Fresh-frozen sections were used. The sections were observed under the fluorescence inverted microscope (Vert.A1; Zeiss GmbH). For H&E staining, fresh-frozen sections were first fixed with 10% neutral buffered formalin for 1 min at room temperature and then dehydrated in an alcohol series, which was as follow: 100% alcohol for 2 min, 75% alcohol for 2 min. Next, the sections were stained with Harris hematoxylin solution for 5 min, cAMP in the supernatant was measured by ELISA according to the manufacturer's protocol. All experiments were conducted in triplicate.

### Table I. Primer sequences used for reverse-transcription-semi-quantitative PCR.

| Name   | Primers (5'-3') | Size (bp) |
|--------|----------------|-----------|
| FSHR   | Forward GCGGAACCCCAACATCGGTGT | 248       |
|        | Reverse TGAAGAAATCTGCGAAGT     |           |
| LHR    | Forward TCAATGTTGTGGCCCTTTGCTA | 256       |
|        | Reverse TGGCGACAAGAATGTGATGG   |           |
| CYP19A | Forward CCATAAAGACCCGATTCCACCA | 396       |
|        | Reverse GCTGAGCCATAAATCGACAGAC |           |
| ER-α   | Forward GAGAGGTATCGTTGTTAGAGA | 207       |
|        | Reverse CCGATCACTACAGTAATAGT   |           |
| ER-β   | Forward TTCTCTTCTCTCTACACT     | 257       |
|        | Reverse ATGTGATAACCTGCGATGG    |           |
| PR     | Forward GGGATGAAAGCAGCTAGCTGT  | 248       |
|        | Reverse AGCATCCAGTGCTTCAACAA   |           |
| AR     | Forward GTGCTGGACAGCAACACAC    | 246       |
|        | Reverse GATCAGGGGCGGACTAGAGC   |           |
| β-actin| Forward GGACAGCATGGAGGAAT      | 185       |
|        | Reverse GATAAGCACAGCTGGATA     |           |

AR, androgen receptor; bp, base pair; CYP19A, cytochrome P450 aromatase; ER-α, estrogen receptor-α; ER-β, estrogen receptor-β; FSHR, follicle-stimulating hormone receptor; LHR, luteinizing hormone receptor; PRL, progesterone receptor. Following centrifugation at 931 x g and 4°C for 5 min, cAMP in the supernatant was measured by ELISA according to the manufacturer's protocol. All experiments were conducted in triplicate.
8 min and washed under running tap water for 1 min. Following that, the sections were counter-stained with DAPI for 5 min at room temperature. Finally, sections were observed under a light microscope (magnification, x200; Nikon, ECLIPSE Ni-E).

**Statistical analysis.** SPSS software (ver. 23.0, IBM Corp.) was used to analyze data. Data are presented as the mean ± standard deviation. One-way ANOVA was used to compare data among different groups, and Dunnett’s and Tukey’s post hoc tests were used to perform multiple comparisons between groups. P<0.05 was considered to indicate a statistically significant difference.

**Results**

**Human GCs cultured in vitro.** The morphological changes of the GCs in culture are presented in Fig. 1A. Cells with different morphologies were observed during early passages. From passage 4, a small group of epithelial-like cells surrounded by spindle-shaped fibroblastic cells was observed. Epithelial-like cells were observed with no obvious morphological changes at passage 99. (B) Growth curve of follicular fluid-derived human GCs. Data are presented as the mean ± standard deviation. (C) Colony formation of follicular fluid-derived human GCs. Scale bars, 100 µm. GCs, granulosa cells.

Figure 1. Cell morphology and proliferation. (A) Shape of follicular fluid-derived human GCs expanded with mouse embryonic fibroblasts-conditioned medium. From passage 4, a small group of epithelial-like cells surrounded by spindle-shaped fibroblastic cells were observed. Epithelial-like cells were observed with no obvious morphological changes at passage 99. (B) Growth curve of follicular fluid-derived human GCs. Data are presented as the mean ± standard deviation. (C) Colony formation of follicular fluid-derived human GCs. Scale bars, 100 µm. GCs, granulosa cells.
reached confluence (data not shown). No notable morphological changes were observed during the following passages. Images of primary, passages 1, 2, 4, 50 and 99 were captured under an inverted light microscope (magnification, x100). Out of 20 experiments, cells died eventually in 19 experiments. Only within one culture, a group of cells were able to proliferate and continue to grow for up to 130 passages. Images of primary, passages 1, 2, 4, 50 and 99 were captured under an inverted light microscope. The GCs growth curve between passages 4 and 34 is presented in Fig. 1B. Cell proliferation was stable and the average doubling time was ~22 h. To further test the cell proliferation capacity, colony formation assays were performed at passage 8. The formation of a large colony derived from a single cell was observed during the first 15 days of culture (Fig. 1C). Following seeding of 100 cells, 57, 53 and 59 clones were observed after 20 days of culturing in three repeated experiments (data not shown). The colony formation efficiency was calculated as 56.3±1.8%.

Surface marker expression profile. Expression profile of cell surface markers was determined at passages 10 and 30 by flow cytometry (Fig. 2). Mouse IgG1 was used as an isotype. Cells expressed mesenchymal cell markers, including CD29, CD166, CD49f and HLA-ABC but did not express hematopoietic cell markers, including CD13, CD31, CD34, CD45, CD90 and CD105. The expression profile was stable and no significant difference was observed between cells at passages 10 and 30.

Expression of FSHR, LHR, CYP19A, estrogen receptor (ER)-α, ER-β, progesterone receptor (PR) and androgen receptor (AR). To further confirm cell identity, the expression of FSHR, LHR and CYP19A were analyzed at passages 10 and 60 by immunofluorescence staining. As presented in Fig. 3A, GCs in P10, 60 and FCs were positive for FSHR, LHR, and CYP19A whilst BMSCs were negative for FSHR, LHR, and CYP19A. Furthermore, the expression of FSHR, LHR, CYP19A, ER-α, ER-β, PR and AR was confirmed by RT-PCR analysis (Fig. 3B). FSHR, LHR and CYP19A were expressed in GCs P10, 60, FCs and ovary tissue but not in mock and chondrocytes. ER-α, ER-β, PR and AR were expressed in GCs P10, 60, FCs, ovary tissue and chondrocytes but not in mock.

Intracellular cAMP accumulation following rhFSH and HCG stimulation. Intracellular cAMP accumulation was measured following 3-h stimulation with 5 IU/ml rhFSH, 100 IU/ml HCG or 5 IU/ml rhFSH+ 100 IU/ml HCG. Intracellular cAMP levels were significantly increased following rhFSH and/or HCG stimulation compared with untreated cells (Fig. 4A).
Steroidogenic activities of GCs in vitro. The steroidogenic activities of GCs were evaluated by measuring the estradiol and progesterone concentrations in the culture medium. The results revealed detectable levels of basal secretion of estradiol and progesterone in the medium. Following 48 h of treatment with A2, estradiol level in the medium was significantly increased, no additional changes were observed when cells were co-stimulated with rhFSH/HCG/forskolin. However, estradiol level was significantly increased following cell co-stimulation with A2 and Bu2cAMP (Fig. 4B). Furthermore, after 48 h of culturing, the levels of progesterone in medium significantly increased following the treatment with rhFSH and/or HCG in the presence of A2. But there was no difference when cells were treated with forskolin, Bu2cAMP combined with A2 or sole A2 administration (Fig. 4C).

Cell survival in SCID mice. Prior to cell transplantation, cells were transfected with an eGFP sequence using a lentivirus vector. Almost all cells expressed eGFP following eGFP transfection and puromycin selection (Fig. 5A). Cells were
Figure 4. Production of cAMP, estradiol and progesterone. (A) Production of cAMP. cAMP contents secreted from FSH-stimulated and/or HCG-stimulated cells were significantly higher than those from non-stimulated cells. Concentrations of (B) estradiol and (C) progesterone in the culture medium. The experiments were repeated three times. In Fig. 4B and C, the standard deviation was 0 in some groups. *P<0.001 vs. control; †P<0.001 vs. control + A2. Bu2cAMP, dibutyryl-cAMP; A2, 4-Androstene-3,17-dione; F, forskolin; rhFSH, recombinant human FSH; GCs, granulosa cells; HCG, human chorionic gonadotropin.

Figure 5. Ovaries of SCID mice 8 weeks following transplantation with eGFP-labelled cells. (A) Follicular fluid-derived human GCs were transfected with eGFP. Left image demonstrates GCs transfected with eGFP under white light, the right image was GCs transfected with eGFP under fluorescence. Almost all cells expressed eGFP following transfection with eGFP. Scale bars, 200 µm. (B) Murine ovaries 8 weeks following cell transplantation, the untransplanted ovaries were used as the control group. (C) Transplanted ovaries under fluorescence inverted microscope. The left black/white images were captured under white light. Red arrow, follicle-like structure in the transplanted ovary. Scale bars, 200 µm. (D) Follicle-like structures, granulosa-like cells and adipocyte-like cells in the transplanted ovaries observed following hematoxylin and eosin staining. The control group was ovaries which were not transplanted with eGFP-labeled GCs, so different stages of follicles can be observed in the ovaries. Scale bars, 100 µm. eGFP, enhanced GFP; GCs, granulosa cells; SCID, severe combined immunodeficiency.
subsequently injected into the right ovaries of the SCID mice. Ovarian tissues were harvested at 8 weeks post-transplantation, 12 tumors were formed in the injected ovaries of 22 SCID mice with no tumors formed in the un.injected ovaries of 22 SCID mice (three tumors and three control ovaries are presented in Fig. 5B). In addition, fluorescence microscopy of frozen tissue sections demonstrated that almost all tumors comprised eGFP-positive cells and that a few of follicle-like structures were observed (Fig. 5C). The presence of follicle-like structures was confirmed by histological analysis following hematoxylin and eosin staining (Fig. 5D). In addition, numerous adipocytes-like cells and some granulosa-like cells were identified in the tumors (Fig. 5D). No mass or tumor was detected in the SCID mice which were injected with cells into subcutaneous back or abdominal cavity (data not shown).

Discussion

The present study aimed to develop an improved culture medium to support GC proliferation in vitro. MEF-conditioned medium, which had commonly been used in mouse and human embryonic stem cell cultures (24,25), was tested. Cells from follicular fluid could be maintained up to 14 weeks in culture (data not shown), which was similar to findings previously reported (12,15). Although we tried to establish an immortal GCs cell line 20 times, only one clone was able to grow up to 130 passages without morphological changes. Because the cultured cells usually stopped proliferating eventually in the present study, a group of epithelial-like cells at passage 4 was discovered and was found to rapidly proliferate. At passage 8, a uniform cell population was obtained, which kept growing up to 130 passages without morphological changes. Cells proliferated in a stable manner, the average DT was ~22 h and the colony formation ability was ~56.3%. These results indicated the presence of stable GCs cultured in vitro for over one year.

Numerous GCs, theca cells and blood cells are present in the follicular fluid of patients included in IVF programs (26). Furthermore, contamination with vaginal and ovarian surface epithelial cells commonly occurs due to transvaginal follicular aspiration (27). To confirm the identity of the cell culture established in the current study, flow cytometric analysis, immunofluorescence staining and RT-qPCR analysis were performed. The results demonstrated that GCs expressed HLA-ABC, the most important human histocompatibility antigen (28), and also mesenchymal lineage markers including CD29, CD166 and CD49f (25) but were negative for hematopoietic cell markers including CD13, CD31 and CD45 (12), which indicated that cells were of mesodermal origin (29). In addition, cells were positive for FSHR, LHR and CYP19A. For the immunofluorescence staining experiment, BMSCs were used as the negative control. It has been commonly accepted that pituitary-derived hormones, including FSH and LH, act exclusively on the gonads and are only expressed on ovary and testis gonad cells (30); however, in the last decade, whether FSHR and LHR are expressed in nongonadal tissues/cells remains a controversy. Previous studies reported extremely low levels of FSH, LH receptors in nongonadal tissues/cells (31,32), including human umbilical cord and osteoclasts. Kumar (33) therefore highlighted the need for novel genetic models to investigate the extragonadal actions of FSH. Previous studies revealed no expression of FSH and LH receptors in umbilical cord and bone cells (34-36). In the present study, FSHR and LHR were not detected in BMSCs. In addition, experiments were performed where mock cells were only incubated with secondary antibodies and fresh GCs were used as a positive control. GCs are the only cell type in the female body possessing the FSHR (37). The expression of FSHR and LHR were neither observed on BMSCs nor on mock cells, which suggested that cells may be GCs. Kossowska-Tomaszczyk et al (3) revealed that long-term cultivated human GCs expressed human mesenchymal stem cell markers, including CD29, CD44, CD90, CD105, CD117 and CD166 but not CD73. Bruckova et al (15) reported that when follicular fluid was added to the culture medium, long-term cultured human GCs highly expressed human mesenchymal cell markers, including CD29, CD44, CD73, CD90 and CD166, and FSHR; however, they did not express CD31,CD34,CD49d, CD49e, CD106, CD184, CD197 or HLA-II. Furthermore, the immortalized GC line COV434 highly expresses HLA-1, slightly expresses FSHR, LHR and CD166, and does not express CD29, CD31, CD34, CD45, CD49d, CD49e, CD73, CD90, CD105, CD106 and HLA-II (15). The phenotype of the cells established in the present study was therefore similar to the GC line COV434 and the GC lines that have been previously reported.

In humans, immortalized GC lines can be established using ovarian tumor samples (COV434, KGN and HSOGT cell lines) or via gene transfection (SVOG-4o, HGL5, HO-23, GC1a, HGP53 and HGrC1 cell lines) (38). The functions of immortalized GCs, including their role in hormone secretion and their response to hormones, vary from cell to cell. For example, KGN, HGL5 and HGrC1 have been reported to secrete estrogen and progesterone (39-41); however, COV434 and HSOGT only produce estrogen (42-44), and HO-23, HGP53 and SVOG-4o only synthesize progesterone (45-47). Furthermore, COV434, KGN, HGP53 and HGrC1 cells respond to FSH stimulation (41-43,47,48) whereas SVOG-4o cells are sensitive to LH/HCG stimulation (45). The cell lines HGL5 (39), HO-23 (46) and GC1a (49) do not respond to either FSH or LH. Functional analyses of the immortalized GCs established in the present study demonstrated that intracellular cAMP accumulation could be upregulated by FSH/HCG stimulation. In addition, A₂ treatment induced estradiol secretion, combined A₂ and Bu-cAMP treatments significantly improved estradiol secretion, whereas A₂ treatment combined with FSH/HCG did not stimulate estradiol secretion. In addition, the present study demonstrated that progesterone secretion was induced by FSH and/or HCG treatment, but not by the PKA pathway activator forskolin (50) or by Bu-cAMP. These results, and the underlying mechanisms, should be further investigated in the future. FSH/HCG binding to their receptors activates the protein kinase A (PKA) pathway that stimulates cAMP accumulation (51,52). Cells established in the present study may act directly on FSHR and LH/HCG receptors which may activate the PKA pathway in order for FSH/HCG to exert their roles. However, certain other crucial factors are involved in the steroid synthesis process such as certain hormone glycosylation variants, potentiating antibodies and small molecule ligands (52), which could be further investigated in future studies. The long-term cultivated cells from the present study
may provide a model to study the physiological mechanisms of steroidogenesis and folliculogenesis.

To determine whether the cells established in the present study could survive in SCID mice, cells were injected into SCID mice ovaries. Tumors were observed at 8 weeks post-injection. However, no tumor was observed when the same number of cells was injected subcutaneously or intra-peritoneally in SCID mice (data not shown). This could be due to the blood supply in the ovaries being better compared with the other two locations or due to ovaries providing a unique environment for cell survival. In particular, this study reported that cells formed follicle-like structures. However, whether the surviving cells could respond to FSH/HCG stimulation in vivo was not investigated. In addition, the hormone levels of the tumor-bearing animals should be determined in the future.

Previous studies reported that GCs possess a multipotent differentiation capacity (9,11,13,53). In the current study, cells spontaneously differentiated into adipocyte-like structures following intra-ovarian injection. The multipotent differentiation capacity of this cell line is currently being investigated.

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Availability of data and materials

The datasets used and/or analyzed during the present study available from the corresponding author on reasonable request.

Authors’ contributions

AA and WZ designed the study, interpreted the results, wrote the manuscript and approved the final form of the manuscript. AA, ZT and YL conducted the experiments. SY performed the cell analysis. YC participated in study design, data interpretation, critical revision and approved the final form of the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate

The study was approved by the Ethics Committee of the Shanghai Ninth People’s Hospital affiliated with Shanghai Jiao Tong University School of Medicine. Written informed consent was provided by all participants prior to the study. Animal studies were approved by the Animal Experimentation Ethics Committee of the Shanghai Ninth People’s Hospital affiliated with Shanghai Jiao Tong University School of Medicine.

Patient consent for publication

All patients provided informed consent for the publication of their clinical data.

Competing interest

The authors declare that no conflicts of interest to declare.

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