INTRODUCTION

Ovarian germline stem cells (OGSCs) differentiate into oocytes before birth and thus, their existence in the postnatal ovaries is controversial in most mammalian species (Telfer et al., 2005). In contrast, the existence of OGSCs in adult fish ovaries has been demonstrated (Nakamura et al., 2010) and the production of offspring through transplantation of adult female-derived OGSCs has been achieved in zebrafish (Wong et al., 2011; Wong et al., 2013) and trout (Yoshizaki et al., 2010; Lee et al., 2016). Thus, fish OGSCs can be a useful resource for the study of...
of developmental biology, genetics and biotechnology like primordial germ cells (PGCs) and spermatogonial stem cells (SSCs). However, their utilization in the fields has been restricted due to a lack of knowledge regarding their in vitro culture implying that the effort should be made to establish a stable fish OGSC culture system.

Previously, Wong et al. demonstrated that rainbow trout splenic feeder cells expressing zebrafish leukemia inhibitory factor (LIF) and zebrafish ovarian somatic feeder cells expressing zebrafish fibroblast growth factor 2 (FGF2), glial cell derived neurotrophic factor (GDNF), gonadal soma derived factor (GSDF) or LIF could not only maintain germline competency of zebrafish OGSCs but also support their growth in vitro (Wong et al., 2013). For Japanese medaka (Oryzias latipes), we, recently, used the method that coats solid surface with polymers to enhance the culture outcome of OGSCs. As a result, coating the dishes with polydopamine, a polymer generated from dopamine through oxidation, influenced positively on the adhesion, survival and potency maintenance of enriched OGSC populations under feeder-free culture condition (Jeong et al., 2019). However, this method was effective only for the maintenance of medaka OGSCs in culture, and it did not induce significant growth of them unlike the feeder-dependent culture system that was used for zebrafish OGSC culture. This suggests that it is necessary to evaluate the effects of feeder-dependent culture system on medaka OGSC culture to know the more optimal basis for establishing long-term culture of OGSCs in medaka model.

For this reason, this study was performed to investigate the effects of feeder cells on the primary culture of O. latipes ovarian cell population, which were enriched by percoll density gradient centrifugation and differential plating, to find the clues to develop the feeder-dependent culture system for medaka OGSCs. In order to do that, we first examined the optimal tissue source to be able to supply feeder cell lines stably and then the enriched ovarian cell populations were cultured on the established feeder cell lines to evaluate its effectiveness on cell proliferation. Additionally, a feeder cell line artificially expressing medaka FGF2 was prepared and its effectiveness to O. latipes ovarian cell culture was also evaluated.

**MATERIALS AND METHODS**

**Animals**

Adult Japanese medaka (O. latipes) was purchased from a local aquarium and reared in 20 L tanks at 26°C. Fish were fed two or three times a day with a commercial diet for flounder larvae (EWHA, Busan, Korea) and the photoperiod was maintained at light for 14 h and darkness for 10 h. For the experiments, adult females were separated from the others and starved for at least 24 h. All experiments using fish were conducted in compliance of ethical guidelines from the Institutional Animal Care and Use Committee (IACUC) of Pukyong National University, which approved our research proposal (approval number: 2016-07).

**Primary culture of the cells derived from three tissues**

In order to establish feeder cell lines for in vitro culture of O. latipes ovarian cell populations, primary cultures were carried out with the cell populations from embryos, fins and ovaries. Embryos at stage 32 to 36 were disinfected with 70% (v/v) ethanol (SK Chemicals, Sungnam, Korea) for 10 sec and washed twice with Dulbecco’s phosphate buffered saline (DPBS; Gibco, Grand Island, NY, USA). After the removal of chorion and egg yolk with syringe needles, an embryo was transferred into each well of 0.1% (w/v) gelatin (Sigma-Aldrich, St. Louis, MO, USA)-coated 96-well culture plates (Thermo Scientific, Vernon Hills, IL, USA) filled with culture medium. Each embryo was dissociated by pipetting 15 times using a syringe equipped with 26 gauge needle and the dissociated cells were cultured. To obtain fins and ovaries, fish were anesthetized with 0.1% (v/v) 2-phenoxyethanol (Sigma-Aldrich) and disinfected with 70% (v/v) ethanol. For fin cell culture, caudal fins were removed from fish and cut into approximately 2 mm² using a surgical blade and tissue fragments were placed on 0.1% (w/v) gelatin-coated 35 mm culture dishes (SPL Life Sciences, Pocheon, Korea). After pressing the tissue fragments with coverslips, they were cultured in culture media. To isolate ovarian cells, ovaries were washed twice with DPBS containing 1% (v/v) penicillin and streptomycin (P/S; Gibco) and disassociated mechanically with sterile scissors. Subsequently, enzymatic dissociation was conducted in Leibovitz’s L-15 medium (L15; Gibco) supplemented with 500 U/mL collagenase type I (Worthington Biochemical Corporation,
Lakewood Township, NJ, USA) for 1 h at 28°C. After dissociation, an equal volume of L15 containing 10% (v/v) fetal bovine serum (FBS: Gibco) was added and the cell suspension was filtered on a 40 μm cell strainer (Falcon, Corning, NY, USA) to remove debris. Finally, the cells were washed twice with DPBS containing 1% (v/v) P/S and then used for experiments. For culture, the ovarian cells isolated were resuspended in culture medium and seeded on 0.1% (w/v) gelatin-coated culture dishes. Culture media were L15 containing 20% (v/v) FBS and 1% (v/v) P/S regardless of the type of cell populations. All cell populations were cultured at 28°C with an air atmosphere and culture medium was changed every two or three days. When the cells reached more than 80% confluency, they were subcultured at a ratio of 1:2 to 1:3.

**Preparation of *O. latipes* FGF2 expression vectors**

To obtain *O. latipes* FGF2 cDNA, total RNA was extracted from adult testes using RNeasy Plus Mini Kit (Qiagen, Valencia, CA, USA) and cDNA was synthesized from 1 μg total RNA using GoScript reverse transcription system (Promega, Madison, WI, USA) according to the manufacturer’s instructions. After PCR amplification of the CDS region, a target band (468 bp) was extracted using gel extraction kit (Genotech, Daejeon, Korea) and ligated into a vector of All in One™ PCR Cloning kit (Biocart, Daejeon, Korea) according to the manufacturer’s instructions. Then, the insert-ligated vector was cloned into *E. coli* DH5α and the clones were analyzed by a commercial sequencing service (CosmoGenetech, Seoul, Korea) after plasmid extraction using Plasmid Extraction Mini Kit (Favorgen, Pingtung, Taiwan). FGF2 cDNA from the clones that have same sequences with that in the NCBI GenBank database was subcloned into pcDNA5-Flag-Med25 (Addgene, Cambridge, MA, USA) using the restriction sites of Nhe I and Not I. The primer sequences and PCR conditions used in this study are listed in Table 1.

**Establishment of a stable cell line that expresses foreign FGF2**

An established embryonic cell line was transfected with 1 μg FGF2 expression vectors using 1 μL lipofectamine 2000 (Invitrogen, Carlsbad, CA, USA) and the transfected cells were selected by treating 600 μg/mL hygromycin (Invitrogen) for 4 weeks. The colonies formed were harvested using cloning cylinders (Corning Life Sciences, Corning, NY, USA) and cultured individually. After establishment of stable cell lines, FGF2 mRNA expression was confirmed by RT-PCR analysis. To detect the protein expression, a cell line, of which FGF2 mRNA expression was confirmed, was used. Cells harvested from a 100 mm culture dish (SPL Life Sciences) were washed twice with DPBS. Then cell lysates were obtained by a sonication in 1% (w/v) triton X-100 (Sigma-Aldrich) in DPBS and protein concentration of the cell lysates was measured using Pierce BCA protein assay kit (Thermo Scientific) according to the manufacturer’s instructions. The denatured protein was loaded on 15% acrylamide SDS gels, electrophoresed for 1.5 h at 100

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**Table 1. Primer sequences used in this study**

| Genes                  | Primer sequences (5’ > 3’)             | Product size (bp) | PCR condition | Accession number |
|------------------------|----------------------------------------|-------------------|---------------|-----------------|
| FGF2 for cloning       | Forward, ATGGCTACGGGAGAAAATCACC        | 468               | 35 cycles of 94°C for 30 s, 51°C | XM_004086561.2   |
|                        | Reverse, TTAGTACTTGGCAGACATAAGGC       |                   | for 30 s, and 72°C for 45 s     |                 |
| FLAG-tagged FGF2 for cloning | Forward, GC ACTGCTACGGGAGAAAATCACC | 520               | 28 cycles of 94°C for 30 s, 65°C |                |
|                        | Reverse, CGGCGGCTACCAATCGTCTCATCTTGTAAATC |                   | for 30 s, and 72°C for 45 s     |                 |
| Endogenous FGF2 for RT-PCR | Forward, TGTCCTGGTATCATTTGTTGTC | 186               | 35 cycles of 94°C for 30 s, 55°C | XM_004086561.2   |
|                        | Reverse, CTCCTCGTATACACAGCA            |                   | for 30 s, and 72°C for 45 s     |                 |
| FLAG-tagged FGF2 for RT-PCR | Forward, ATGGCTACGGGAGAAAATCACC | 489               | 35 cycles of 94°C for 30 s, 48°C |                |
|                        | Reverse, GTGTCATCTTTGAATC              |                   | for 30 s, and 72°C for 45 s     |                 |
| Nanos2 for RT-PCR      | Forward, GGTGCAACAACATGGTGAGT          | 262               | 35 cycles of 94°C for 30 s, 56°C | NM_001104447.1   |
|                        | Reverse, CTCGCAAGAGTGGTATGTTAAT        |                   | for 30 s, and 72°C for 45 s     |                 |
| Vasa for RT-PCR        | Forward, GAGAAGTTTCCGACCCACAG          | 177               | 35 cycles of 94°C for 30 s, 58°C | NM_001104676.1   |
|                        | Reverse, AATGTTTGGGGAGGTTCA            |                   | for 30 s, and 72°C for 45 s     |                 |
| β-actin for RT-PCR     | Forward, CCACCATGTACCGGGGAAATC         | 153               | 35 cycles of 94°C for 30 s, 58°C | NM_001104808.1   |
|                        | Reverse, GCTGGAAGGTTGGACAGAG           |                   | for 30 s, and 72°C for 45 s     |                 |
Enrichment of germ cell populations

To obtain ovarian cell populations, 10 ovaries were subjected to mechanical and enzymatic dissociation as mentioned in the part of ovarian cell isolation. Then, to enrich germ cell populations by removing red blood cells and anchorage-dependent cells, the retrieved cells were washed twice with DPBS containing 1% (v/v) P/S and loaded onto the top of layered Percoll (Sigma-Aldrich) solution consisting of 20, 25, 30, 35, 40, 50 and 60% in a 15 mL conical tube (Falcon). After centrifugation at 800×g for 30 min, density fractions of 20-40% were harvested. After being washed twice with DPBS, the cells were resuspended in L15 supplemented with 10% (v/v) FBS and 1% (v/v) P/S. Then, 2-5 × 10⁶ cells were seeded on a 35 mm petri dish (SPL Life Sciences) coated with 0.1% (w/v) gelatin and incubated for 15 h at 28°C. The cells that were floating or loosely bound were harvested and used for further experiments. These cells were designated as “enriched ovarian cell population” after this.

In vitro culture of enriched ovarian cell populations

To investigate the effect of feeder cells on the growth of ovarian cells, 1.5 × 10⁴ live cells from the enriched ovarian cell populations were seeded on feeder cells or feeder-free condition. For the culture on feeder cells, 5 × 10⁴ feeder cells were seeded on a well of 24-well plates (SPL Life Sciences) and mitotically inactivated by treating 10 μg/mL mitomycin-C (Sigma-Aldrich) for 3 h before use. Prior to initiation of culture, all culture plates with or without feeder cells were incubated with ovarian germline stem cell medium (OGSM) for 24 h at 28°C in an air atmosphere. Enriched ovarian cell populations were labeled with 6 μM PKH26 (Sigma-Aldrich) for 3 min and cultured in OGSM consisting of L15 supplemented with 25 mM HEPES (Sigma-Aldrich), 6 mg/mL D-(+)-glucose (Sigma-Aldrich), 1% (v/v) glutamax (Gibco), 1% (v/v) non-essential amino acids (Gibco), 0.5% (w/v) BSA (Sigma-Aldrich), 5% (v/v) FBS, 0.25% (v/v) trout serum (Caisson Laboratories, Smithfield, UT, USA), 1 μg/mL medaka embryo extract, 1% (v/v) P/S, 25 μg/mL bovine insulin (Sigma-Aldrich), 100 μg/mL recombinant human apo-transferrin (Sigma-Aldrich), 10 ng/mL recombinant human basic fibroblast growth factor (bFGF: Gibco), 10 ng/mL recombinant human glial cell-derived neurotrophic factor (Peprotech, Rocky Hill, NJ, USA), 5% (v/v) Knockout serum replacement (Gibco), 50 μM ascorbic acid (Sigma-Aldrich), 50 μM β-mercaptoethanol (Gibco), and 2 mM sodium selenite (Sigma-Aldrich) at 28°C in an air atmosphere. Medaka embryo extract was extracted as described previously (Choi and Gong, 2018). Half of the culture medium was replaced with fresh one every three days. For measurement of the cell number after culture for 10 days, the cells were harvested by treatment with 0.05% trypsin-EDTA (Gibco) and the number of fluorescent cells was counted using a hemocytometer (Paul Marienfeld GmbH & Co. KG, Lauda-Königshofen, Germany) under a TS-100F microscope equipped with a fluorescent unit (Nikon, Tokyo, Japan).

Statistical analysis

The statistical analyses were performed using SPSS version 18 (IBM-SPSS, Chicago, IL, USA). The data were analyzed by One-way-ANOVA or t-test followed by Duncan’s method. Significant differences among groups were determined when p < 0.05.

RESULTS

Establishment of feeder cell lines

To find optimal tissue source to be able to supply feeder cell lines, culture of the cells from three tissues including embryonic tissue, fin, and ovary was performed. All cell populations regardless of tissue sources showed primary adherence to substrates and of those, more than 80% were subcultured at least once (Table 2). However, cell death was observed after the removal of fin fragments and after 2 weeks of culture in fin- and ovary-derived cells, respec-
Ryu and Gong. Medaka Ovarian Cell Culture on Feeder Cell Layer

The cell populations derived from fins and ovaries could not reach to passage 5 in culture and were just maintained for $18.2 \pm 6.0$ and $20.8 \pm 13.4$ days, respectively. On the contrary, $18 (66.7\%)$ out of $27$ cell populations from embryonic tissues was cultured to more than passage $5$ and $9 (33.3\%)$ out of $27$ was cultured to more than passage $20$. Twenty two ($81.5\%)$ out of $27$ cell populations was maintained in culture for more than $50$ days. These indicated that the embryonic tissues were an optimal tissue source to derive the cell lines that can be used as feeder cells among three tissues tested. Three embryonic cell lines (named as EC1, EC2, and EC3) were used as feeder cells in this study. These ECs contained two cell types that were morphologically different: fibroblast-like cells and epithelial-like cells. In EC1 and EC3, fibroblast-like cells were predominant whereas epithelial-like cells were predominant in EC2 (Fig. 1A). These three ECs did not express germline stem cell-specific Nanos2 and germ cell-specific Vasa genes indicating that germline cells were not included in these ECs (Fig. 1B).

**Effects of feeder cell lines on in vitro culture of enriched ovarian cell populations**

To examine the effects of feeder cells on the culture of ovarian cells, enriched ovarian cell populations that were labeled with fluorescent PKH26 were cultured on feeder cells derived from each of three ECs (EC1, EC2, and EC3) and the change of cell number was investigated after culture for $10$ days. The enriched ovarian cell populations cultured on feeder cells formed cell aggregates unlike those cultured under feeder-free condition and those cultured on EC1 and EC3 showed a significant increase in cell number after culture for $10$ days compared to those cultured on feeder-free condition ($p < 0.05$; Fig. 2).

**Effects of FGF2-expressing feeder cell line on in vitro culture of enriched ovarian cell populations**

The FGF2 expression vector was transfected into EC3 and stable cell line expressing FGF2 was subsequently established by antibiotic selection. RT-PCR analysis confirmed that the established cell line expressed the mRNA of foreign FGF2 whereas it did not express endogenous FGF2 (Fig. 3A). Moreover, the expression of FLAG-tagged

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**Table 2. Culture outcomes of cell populations derived from three different tissues of *Oryzias latipes***

| Tissues | Culture methods     | Trials | No. (%) $^+$ of cell populations initially attached | No. (%) $^+$ of cell populations subcultured to Passage 1 | Passage 5 | Passage 20 | No. (%) $^+$ of cell populations maintained $\geq 50$ days |
|---------|---------------------|--------|--------------------------------------------------|--------------------------------------------------------|---------|---------|--------------------------------------------------|
| Embryo  | Dissociated cell culture | 27     | 27 (100)                                         | 22 (82)                                                | 18 (67) | 9 (33)  | 22 (81)                                          |
| Fin     | Explant culture      | 5      | 5 (100)                                          | 4 (60)                                                 | 0 (0)   | 0 (0)   | 0 (0)                                            |
| Ovary   | Dissociated cell culture | 8      | 10 (100)                                         | 8 (80)                                                 | 0 (0)   | 0 (0)   | 0 (0)                                            |

$^+$Percentage of trials.
FGF2 (18.6 kDa) was detected in the stable cell line by western blot analysis (Fig. 3B). To test the effects of FGF2-expressing feeder cell line (named as EC3F) on ovarian cell culture, enriched ovarian cell populations were cultured on EC3 or EC3F feeder cells. Furthermore, the effects of recombinant human bFGF as a medium supplement were also tested by removing it from OGS. As the results, the cultured ovarian cells formed cell aggregates during in vitro culture regardless of treatment groups (Fig. 4A). After culture for 10 days, the number of cells increased from 1.08 to 1.37-fold relative to the initial cell number seeded according to the experimental groups (Fig. 4B). No significant difference was detected among experimental groups indicating that both human recombinant bFGF and medaka FGF2 produced by EC3F were not effective to induce ovarian cell proliferation.
DISCUSSION

To develop a suitable culture condition for a specific cell type, several points including physical aspects, nutrients, signaling molecules such as hormones and cytokines, substrate condition and culture method should be considered (Freshney, 2010). Feeder cells have been often utilized to develop or improve the culture condition for certain cells, as they can supply several soluble signaling molecules and extracellular matrices to target cells during in vitro culture (Hongisto et al., 2012; Villa-Diaz et al., 2013). In mammals, the cells derived from embryos such as mouse embryonic fibroblasts have been frequently used for the culture of GSCs (Kanatsu-Shinohara et al., 2003; Kubota et al., 2004). In the present study, embryonic cell lines were established and two of them significantly promoted the growth of enriched ovarian cell populations when used as feeder cells. This would have been due to the extracellular matrices and cytokines secreted from the feeder cells. For establishment of embryonic cell lines, the embryos at stages 32 to 36 were used and they are known to express various kinds of hormones and growth factors such as GSDF and insulin (Assouline et al., 2002; Shibata et al., 2010). Based on this, the specific molecules that positively influenced to ovarian cells in culture needs to be found for the optimization of OGSC culture system.

To achieve more active proliferation of ovarian cells, a feeder cell line artificially-expressing FGF2 was developed. Because of the amino acid dissimilarities between mammals and fish, mammalian growth factors supplemented in culture media may have low or no activity on fish GSCs (Kawasaki et al., 2012). Thus, testing fish growth factors instead of mammalian ones is a major step for the establishment of a more suitable culture condition for fish OGSCs. FGF2, a member of the FGF family, is an important factor for the self-renewal of mammalian SSCs (Ishii et al., 2012; Takashima et al., 2015). The protein identity of FGF2 between human and medaka is 73% and it was reported that recombinant human FGF2 supports SSCs from fish including medaka, trout and zebrafish at 1-100 ng/mL concentrations (Hong et al., 2004; Shikina et al., 2008; Kawasaki et al., 2012). In addition, it was reported that zebrafish FGF2 supports the colony formation and proliferation of zebrafish SSCs and OGSCs during in vitro culture (Wong and Collodi, 2013; Wong et al., 2013). In our current study, culture of enriched ovarian cell populations on FGF2-expressing feeder cells did not show any significant promoting effects on cellular growth. This could be caused by the difference between species. Otherwise, the quantity of FGF2 secreted from the feeder cells might not sufficient to promote the growth of ovarian cells. Further experiment using the purified FGF2, the concentration of which can be adjusted, will be helpful. In addition, it may be necessary to use the feeder cells expressing multiple growth factors based on a previous report that showed that the feeder cells expressing two growth factors effectively promoted zebrafish OGSC proliferation compared to the feeder cells expressing a single growth factor (Wong et al., 2013).

In the present study, the effects of feeder cells on the culture of medaka ovarian cell populations were examined. The feeder cells derived from medaka embryonic tissues were able to support ovarian cell proliferation during short culture period. The results from this study will provide fundamental information for development of the optimal conditions for OGSC culture and will contribute to the utilization of OGSCs in various areas.

CONFLICTS OF INTEREST

No potential conflict of interest relevant to this article was reported.

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AUTHOR CONTRIBUTIONS

JH Ryu performed the experiments and wrote the paper; SP Gong conceived and designed the experiments and wrote the paper.

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