Abstract. Background/Aim: Cryopreservation of cell lines has been widely used in the laboratory; however, cryopreservation of organs is still considered to be difficult. The submandibular gland (SMG) of fetal mice is one of the best-characterized organs. We investigated the conditions for cryopreserving SMG rudiments. Materials and Methods: Embryonic day 13 SMG rudiments were cryopreserved with or without a cryoprotectant. They were thawed and incubated in DMEM/F12 medium. Moreover, the influence of EGF stimulation on the signaling cascade after frozen-thawing the rudiments was analyzed by Western blotting. Results: When SMG rudiments were cryopreserved without a cryoprotectant, all cells in the rudiments died. However, the SMG rudiments that had been preserved in a cryoprotectant showed branching morphogenesis. Additionally, the responsiveness of signaling cascades to EGF did not differ between frozen with a cryoprotectant and non-frozen rudiments. Conclusion: Cryopreservation might be a useful technology for preserving tissues from small organs, such as fetal SMG rudiments.

Cryopreservation of cell lines has been widely used in the laboratory for in vitro study of cell systems. Cryopreservation is also useful for germ cells in medical care, for organic long-term preservation of induced pluripotent stem cells (iPS cells), and in animal husbandry (1-4). When cells are frozen without using a cryoprotective agent (CPA), most of the constituent cells die. This is because the water molecules inside and outside of the cells crystallize, and the cytoarchitecture, such as cell membranes, is physically destroyed (5, 6). CPAs contain chemical compounds, such as dimethyl sulfoxide (DMSO), glycerol, sucrose and ethylene glycol, and the compositions of CPAs differ depending on the cells to be frozen (5). Despite all the available protocols for successful cryopreservation of different cell types, cryopreservation of organs and tissues is still considered difficult.

Mouse SMG rudiment development begins on embryonic day 12 (E12) when the epithelial tissue of the floor of the mouth sinks towards the direction of the neck. On E13, the SMG rudiment is oval shaped, approximately 500 μm, and it rapidly forms a glandular body by repeated cleft formation and stalk elongation, called branching morphogenesis (BrM). BrM is studied in SMG rudiments on filters floating in serum-free medium in ex vivo organ culture systems (7-10). The epithelial cells constituting SMG rudiments promote BrM when stimulated with cell growth factors. Kashimata, et al. (1997) have reported that epidermal growth factor (EGF) strongly promotes BrM of the SMG epithelium (11), while Hoffman et al. (2002) have reported that fibroblast growth factor (FGF) also promotes BrM of the SMG epithelium (12). EGF and FGF are secreted mainly from the mesenchyme, and growth factor receptors are expressed in the epithelial cells of SMG rudiments (12, 13). Following binding of EGF or FGF to their respective receptors, signaling pathways, such as extracellular signal-regulated kinase-1/2 (ERK1/2), classical mitogen-activated protein kinase (MAPK) and phosphoinositide 3-kinase (PI3K)-AKT cascades are activated (11-15). Therefore, EGF and FGF induce cell proliferation and differentiation in the developing mouse SMG through activation of these signaling cascades (11-15).

Recently, Ogawa et al. (2013) prepared SMGs from immature epithelial and mesenchymal cells obtained from fetal mouse SMGs and transplanted them into mice whose salivary glands had been removed (16). They, then, demonstrated secretion of saliva from the transplanted SMGs. Since steady progress is being made in regenerative therapy of organs, including the salivary gland, the need for...
further development of cryopreservation technologies for organs will almost certainly increase in the future.

In this study, we investigated whether CPAs protect cryopreserved SMG rudiments and induce the morphological changes of BrM in thawed and incubated rudiments, and whether EGF-responsive phosphorylation of ERK1/2 and AKT occurs in SMG rudiments cryopreserved with CPAs.

Materials and Methods

Cryopreservation of SMG rudiments. Pregnant mice (ICR strain) were purchased from Japan SLC (Hamamatsu, Japan). The day of discovery of a vaginal plug was taken as embryonic day 0 (E0). E13 SMG rudiments were removed from fetuses as described previously (13, 17). The SMG rudiments were placed in 4 different CPAs: i) DMEM/F12 medium (Gibco Invitrogen Cell Culture, Carlsbad, CA, USA) containing 100 units/ml penicillin (Gibco), 100 μg/ml streptomycin (Gibco), 150 μg/ml vitamin C (Sigma-Aldrich, St. Louis, MO, USA), and 50 μg/ml transferrin (Sigma-Aldrich), ii) DMEM/F12 containing 10% DMSO (Sigma-Aldrich), iii) TC Protector (TC) (DS pharma Biomedical Co., Osaka, Japan), and iv) CELLBANKER1 (CB) (ZENOAQ, Fukushima, Japan). One SMG rudiment was put into 200 μl of each CPA solution in a cryogenic tube (Thermo Fisher Scientific) and then stored in a deep freezer at -80°C for 24 h. Mr. Frosty is a container whose inner temperature slowly decreases at a rate of 1˚C/min in a –80˚C deep freezer. All animal handling procedures were conducted in accordance with the Guideline for Experimental Animals of the Asahi University (Protocol No. 18-007).

Culture of cryopreserved SMG rudiments. The frozen SMG rudiments were quickly thawed by adding 900 μl of 37°C DMEM/F12 medium, followed by pipetting. Then the SMG rudiments were washed two times with 1 ml of DMEM/F12 medium, placed on a membrane filter (Nuclepore membrane, 0.1 μm pore size; Whatman International, Brantford, UK) floating in 400 μl of DMEM/F12 medium, and cultured in 5% CO2, 95% air and 80% humidity at 37°C ex vivo. The cultured SMG rudiments were photographed at 0, 24, 48 and 72 h following incubation. Subsequently, the endpieces of the SMG rudiments were counted, and the area of the epithelium of the SMG rudiments was measured.

Western blotting for signaling proteins. E13 SMG rudiments were cryopreserved in CB and thawed 24 h later, as described above. The cryopreserved SMG rudiments and non-frozen control SMG rudiments were cultured for 0-72 h. Some of cryoprotective agents (CPA) were protected the cells of SMG rudiments although the protective effects were different from each other. The black arrow indicates a hypertrophy-like structure. Scale bar=200 μm, TC: TC Protector; CB: CELLBANKER1.
rudiments were cultured on membrane filter for 24 h, as described above. Subsequently, EGF 20 ng/ml (R & D System Inc., Minneapolis, MN, USA) was added to the medium, followed by incubation for 0, 10 or 30 min. The rudiments were washed with ice-cold PBS containing 1 mM Na₃VO₄ and phenylmethylsulfonyl fluoride (PMSF, Sigma-Aldrich), and homogenized with cell lysis buffer (Cell Signaling Technology Inc.). Centrifugation was performed at 15,000 rpm for 30 min at 4°C, and the resulting supernatants were collected. The protein concentrations were measured using the Pierce 660 nm Protein Assay Reagent (Thermo Fisher Scientific). Aliquot protein samples (2 μg) were subjected to SDS-PAGE and transferred to polyvinylidene difluoride (PVDF) membranes (BIO-RAD, Hercules, CF, USA). Specific proteins on the membrane were detected by probing with the specific primary monoclonal or polyclonal antibodies described above, followed by secondary antibodies conjugated to horseradish peroxidase. The proteins were detected using ECL Select Western Blotting Detection Reagents (Amersham Biosciences, Buckinghamshire, UK). The intensities of the Western blot bands were analyzed using a CS analyzer (ATTO Corp., Tokyo, Japan). All Western blots were repeated at least three times.

**Statistical analysis.** One-way ANOVA followed by Dunnett’s multiple comparison test was used for comparing experimental groups with the control. Values of $p<0.05$ and $p<0.01$ were considered to be statistically significant.

![Figure 2. Influence of cryoprotective agents on branching morphogenesis of cultured SMG rudiments. The number of endpieces (A) and the area of the epithelium (B) of E13 cultured SMG rudiments held under the different conditions were determined. All SMG rudiments were cultured for 72 h. Values represent the mean±SD of four independent experiments. N.D.: not detected; Quick: quick freezing (using a freezing tube box); Slow: slow freezing (using a Mr. Frosty). *p<0.05, **p<0.01.](image-url)
Results

Culture of cryopreserved SMG rudiments. Figure 1 shows the morphological changes in cultured rudiments of non-frozen control SMGs and thawed SMGs that had been cryopreserved at −80 °C for 24 h in each of the 4 CPAs. The area of non-frozen control E13 SMG rudiments increased (Figure 1A), whereas SMG rudiments that had been cryopreserved in DMEM/F12 medium at −80 °C for 24 h showed no changes at all under both quick- and slow-freezing conditions (Figure 1B). The number of endpieces and the area of the epithelium in cultured rudiments that had been cryopreserved in DMEM/F12 medium at −80 °C for 24 h could not be determined under both quick- and slow-freezing conditions, as the border between the epithelium and mesenchyme of SMG rudiments cryopreserved without any CPA disappeared (Figures 2A and B). However, the SMG rudiments cryopreserved in 10% DMSO under both quick- and slow-freezing conditions grew slowly during incubation (Figure 1C) compared to the non-frozen control SMG rudiments (Figure 1A). Therefore, some epithelial cells and mesenchymal cells in the rudiments cryopreserved in 10% DMSO had remained alive, and BrM occurred during incubation (Figure 1C), even though BrM appeared abnormal, with smaller and fewer epithelial endpieces in the rudiments, resulting in hypertrophy. The number of endpieces and the area of the epithelium of the rudiments cryopreserved in 10% DMSO increased under both quick- and slow-freezing conditions (Figures 2A and B), and those values differed significantly compared to the non-frozen control SMG rudiments (p < 0.01).

The protective effects of TC and CB as CPAs for SMG rudiments were stronger than that of 10% DMSO. As shown...
in Figure 1, the morphological changes of SMG rudiments cryopreserved in TC and CB under both quick- and slow-
freezing conditions were similar to the changes in the non-
frozen control SMG rudiments during 72-h culture (Figure 1A). The number of endpieces and the epithelial area of
cultured SMG rudiments after cryopreservation in TC and
CB (Figure 2A) under both quick- and slow-freezing
conditions were smaller compared to the non-frozen control
SMG rudiments. There were significant differences in the
number of endpieces between SMGs cryopreserved in TC
under quick-frozen and non-frozen control SMGs (\(p<0.01\)),
between SMGs cryopreserved in TC under slow-frozen and
non-frozen control SMGs (\(p<0.05\)), and between SMGs
cryopreserved in CB under quick-frozen and non-frozen
control SMGs (\(p<0.01\)). However, no significant differences
were seen between the number of endpieces in SMGs
cryopreserved in CB under slow-freezing and non-freezing
conditions and in the control SMGs (Figure 2A). Also, no
significant differences were seen between the epithelial area
of the SMGs cryopreserved in CB under slow-freezing
conditions and the non-frozen control SMGs (Figure 2B). On
the other hand, there were significant differences between
SMGs cryopreserved in TC under quick- and slow-freezing
conditions and the non-frozen control SMGs (\(p<0.05\)), and
between SMGs cryopreserved in CB under quick-freezing
conditions and the non-frozen control SMGs (\(p<0.01\)). This
means that BrM of SMG rudiments cryopreserved at -80°C
for 24 h in CB under slow-freezing conditions was normal,
similar to the non-frozen control SMGs. In particular,
following a 24-h culture cryopreservation in CB, the SMG
rudiments looked healthier compared to the other
cryopreserved rudiments (Figure 1E).

Cell signaling systems of cryopreserved SMG rudiments. It
is well known that EGF stimulates ERK1/2 and AKT
phosphorylation and induces BrM in cultured SMG
rudiments (15, 17, 18). We investigated whether functional
ERK1/2 and AKT signaling cascades remained activated in
cryopreserved SMG rudiments. As shown in Figure 3A,
ERK1/2 in cultured non-frozen control SMG rudiments was
phosphorylated at 10 and 30 min following the addition of
EGF. EGF increased the phosphorylation levels significantly,
by more than 3-fold (\(p<0.01\), Figure 3B). EGF also
stimulated ERK1/2 phosphorylation in the cultured SMG
rudiments after cryopreservation in CB (Figure 3A).
The levels of phosphorylation increased by 2.2-fold at 10 min
and by 1.6-fold at 30 min, respectively (\(p<0.05\), Figure 3B).

Figure 4 shows our findings regarding the AKT cascade
in the SMG rudiments. EGF stimulated AKT
phosphorylation in the cultured non-frozen control SMG
rudiments. The levels of phosphorylation were increased 1.9-
fold at 10 min and 1.7-fold at 30 min after EGF stimulation
(\(p<0.05\), Figure 4B). EGF stimulation also increased AKT
phosphorylation in CB-cryopreserved SMG rudiments, by
1.7-fold at 10 min and 1.4-fold at 30 min (\(p<0.05\), Figure 4B). These results show that functional, EGF-responsive ERK1/2 and AKT signaling cascades remain active in SMG
rudiments cryopreserved in CB.

Discussion

Cryopreservation methods have already been developed for
mammalian cell lines and are widely used in the laboratory
for decades. Cryopreservation is also used for oocytes,
embryonic stem (ES) cells and iPS cells (1-3). CPAs, such as
DMSO, glycerol, sucrose and polyethylene glycol are
available for cryopreservation of many types of cells (5).
However, there is no general method of cryopreservation for
tissues, except for testicular and ovarian tissues (19, 20).
Therefore, cryopreservation of organs and/or organ rudiments
remains difficult.

The fetal mouse SMG is a useful model for studying
organogenesis, epithelial-mesenchymal interactions and
BrM, and it is able to undergo BrM even in ex vivo systems
(21-23). Since BrM, which is driven by epithelial-
mesenchymal interactions, can be observed in the culture
systems of SMG rudiments, there is no need for addition of
cell growth factors or serum (18, 24, 25). We tried to
determine the damage caused by cryopreservation of organs
and the effects of various CPAs on epithelial BrM using
incubated SMG rudiments.

Most of the cells in SMG rudiments that were frozen
without any CPA died. The border between the epithelium
and mesenchyme of SMG rudiments cryopreserved without
any CPA was found to have disappeared when they were
cultured ex vivo. However, when cryopreserved in 10% DMSO,
some cells in the SMG rudiments survived, and a few epithelial endpieces of cryopreserved SMG
rudiments were observed after 24 h in culture; and
even increased during longer culture periods. This means
that not only the cells of the epithelium and mesenchyme
of the SMG rudiments, but also the 3D structure of the
extracellular matrix, was partly protected by 10% DMSO.
DMSO has been previously reported to be effective for
the cryopreservation of cell lines (5, 26). Our present
results show that DMSO is also effective for the
cryopreservation of SMG rudiments. However, the
morphological changes were quite different compared to
the non-frozen control SMG rudiments. The rate of BrM
was very slow in culture until 24 h, and a hypertrophy-
like structure was observed in endpieces of the SMG
epithelium, probably due to inhibition of cleft formation
and decreased numbers of endpieces.

The protective effects of TC and CB as CPAs were
greater than that of the 10% DMSO. The morphological
changes in cultured SMG rudiments that had been
cryopreserved in TC and CB were very similar to those of the non-frozen control rudiments. However, the morphological changes after 72 h of culture were totally different compared to those of the 10% DMSO group, although the SMG rudiments cryopreserved in TC showed a slower rate of BrM. The results for the area of the epithelium and the number of endpieces of the cultured SMG rudiments did not differ significantly between the non-frozen control SMG rudiments and the SMG rudiments cryopreserved in CB by slow freezing in a Mr. Frosty. Although TC and CB are sold worldwide as CPAs for the cryopreservation of cell lines, their ingredients are not disclosed. The reagent descriptions state that CB contains serum but TC does not. The difference in protective effect between CB and TC may, thus, be due to the presence or absence of serum.

BrM of SMG rudiments is regulated by many factors, including cell growth factors, adhesion molecules and extracellular matrix (23, 27). These factors are known to stimulate intracellular signaling cascades, including mitogen-activated protein kinase (MAPK) and AKT. We investigated whether these cell signaling cascades were damaged by the cryopreservation of SMG rudiments. Western blotting analysis clearly showed that EGF stimulated the phosphorylation of ERK1/2 in SMG rudiments cryopreserved in CB by a Mr. Frosty. A similar result was obtained for AKT phosphorylation. Although the phosphorylation levels of ERK1/2 and AKT by EGF in cryopreserved rudiments were somewhat lower compared to the levels of the non-frozen control SMG rudiments, the EGF-responsive activation systems for both signaling cascades were at least still present, even after cryopreservation. There have been few research reports on cryopreservation of organs to date. In this study, slow freezing in CB significantly protected the biological functions of SMG rudiments from fetal mice.

Our present findings open the possibility that larger or mature organs may be effectively cryopreserved subsequent to further development of suitable cryopreservation reagents and freezing conditions.

Conflicts of Interest

The Authors have no conflicts of interest to declare regarding this study.

Authors’ Contributions

KA and MK prepared cryopreservation of SMG rudiments. KA, YM and MK performed the experiments of culture of frozen SMG rudiments. YO and KS performed the western blotting and analyzed the data. AS performed morphological analyses of cultured SMG rudiments. All authors (KA, YO, KS, AS, YM and MK) organized the research and wrote the manuscript.

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