Resurrection of a Bull by Cloning from Organs Frozen without Cryoprotectant in a −80°C Freezer for a Decade

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Abstract

Frozen animal tissues without cryoprotectant have been thought to be inappropriate for use as a nuclear donor for somatic cell nuclear transfer (SCNT). We report the cloning of a bull using cells retrieved from testicles that had been taken from a dead animal and frozen without cryoprotectant in a −80°C freezer for 10 years. We obtained live cells from defrosted pieces of the spermatic cords of frozen testicles. The cells proliferated actively in culture and were apparently normal. We transferred 16 SCNT embryos from these cells into 16 synchronized recipient animals. We obtained five pregnancies and four cloned calves developed to term. Our results indicate that complete genome sets are maintained in mammalian organs even after long-term frozen-storage without cryoprotectant, and that live clones can be produced from the recovered cells.

Introduction

In several mammalian species, cloned animals have been successfully produced by somatic cell nuclear transfer (SCNT) [1–5]. The integrity of the genome in the donor nuclei is essential for the full-term development of such cloned animals. Therefore, somatic cells of endangered or extremely valuable animals have been cryopreserved for maintenance of genetic resources [6–8]. To maintain the viability of mammalian cells for long-term storage, the cells are usually frozen-stored with appropriate cryoprotectants [9]. However, in cases in which donor individuals are already dead or their species extinct, cryopreserved cells with intact nuclei are not always available. There have been attempts to rescue animal genetic resources from dead cells or non-cryoprotected frozen specimens. Freeze-dried spermatozoa [10] and nuclei of spermatids in frozen mouse testes without cryoprotectant [11], when injected into oocytes, could produce normal pups, indicating that genomic integrity was maintained in some dead male gametes after freezing. Moreover, production of viable SCNT offspring from dead [12] and heat-denatured [13] somatic cells suggested that live donor cells were not required for SCNT. Recently, Li and Mombaerts [14] established mouse nuclear transfer embryonic stem (ntES) cells from embryos cloned from nuclei of dead cells after freezing without cryoprotectant. Moreover, blastocyst development has been reported after nuclear transfer with freeze-dried somatic sheep cells [15]. However, such cells are usually severely damaged by ice crystals and osmotic stress [9]. Very recently, Wakayama et al. reported the generation of cloned mice from nuclei recovered from a body that had been frozen without cryoprotectant [16]. They showed that SCNT embryos from nuclei of frozen brain and tail blood could develop to term, although the developmental potential was very low for other frozen tissues [16]. Therefore it is unclear whether the nuclei in the other frozen tissues remained intact.

A bull named Yasufuku was one of the most important and famous sires in the history of breeding Wagyu cattle due to his contributions to the improvement of the quality of marbling, which is a major characteristic of Wagyu beef. Yasufuku died of senility at an age of 13.5 years in September 1993. His testicles were collected 12 hours after his death, then wrapped in aluminum foil and placed in a −80°C freezer without cryoprotectant for 10 years. The testicles were then transferred to liquid nitrogen without cryoprotectant for another 3 years. We examined whether intact and cultivable somatic cells could be retrieved from the testicles and whether the nuclei from such cells could contribute to the development of viable offspring after SCNT with enucleated oocytes. In this study, we succeeded in obtaining four live cloned calves from these frozen organ cells. Three calves are healthy though one died two days after birth. To our knowledge, this is the first report of the resurrection of a dead elite livestock specimen from a non-cryoprotected frozen organ by cloning.

Results

Retrieving cells from bull testicles frozen without cryoprotectant for one to four months

Before experimenting with tissues from Yasufuku, we conducted some preliminary experiments with fresh frozen testicles. We collected testicles from three 12- to 15-month-old bulls and froze
them at $-80^\circ$C without any special treatment in a freezer for one to four months. We then dissected the frozen testicles into different parts, caput epididymis, cauda epididymis, spermatic cords and testes. The parts were thawed, minced and digested with collagenase and dispase and then cultured. In our preliminary experiments, we used Dulbecco’s modified Eagle’s medium (DMEM) or $\alpha$-minimum essential medium ($\alpha$-MEM) to obtain primary cultures from the frozen testicles. However, no cells grew from the thawed tissue. It was possible that cells in the thawed tissue might be with quite low proliferating activity even when they were alive. Therefore we selected MF-start$^{TM}$ that was developed to readily obtain initial outgrowth of cells with low proliferating activity at primary culture. We obtained live and culturable cells from both the caput epididymis and the spermatic cords but not from the cauda epididymis or the testes. Most of the culturable cells proliferated actively and populations expanded, suggesting that the cells were in normal condition. We used these cells to produce SCNT embryos by electrofusion with enucleated oocytes. Fibroblasts taken from bovine car tissue were used as controls. The SCNT embryos were then cultured for 168 hours [17]. The SCNT experiments were repeated 3 times. The SCNT embryos reconstructed from cells derived from the frozen testicles did not differ significantly from SCNT embryos generated from control cells with regard to blastocyst rates from cultured embryos (22.1% and 20.2%, p > 0.05, Student’s t-test), number of cells in blastocysts [130±43 and 121±43 (mean±SD), p > 0.05, Student’s t-test], or ratio of the number of cells in the inner cell mass relative to the total number of cells in the blastocysts (21.1% and 22.6%, p > 0.05, Student’s t-test).

Our results indicated that culturable somatic cells could be retrieved from bovine organs that had been frozen without cryoprotectant and that the cells could be used for nuclear transfer.

Retrieving cells from Yasufuku’s testicles

We isolated the caput epididymis and the spermatic cords from Yasufuku’s frozen testicles and cut them into several small pieces (Fig. 1). We thawed, minced and digested the pieces of tissue as described above, and cultured the precipitates. We obtained four cultures of viable cells that were derived from two pieces of frozen tissue. As shown in Figure 2, the cells appeared to be either fibroblast-like (cell lines A and C; Figs. 2A and 2C, respectively) or epithelial-like (cell lines B and D; Figs. 2B and 2D, respectively).

We used primary cultures of cell lines A and B for SCNT, and we cryopreserved cell lines C and D in a cell cryopreservation solution (Cellbanker; Mitsubishi Kagaku-Iatron, Tokyo, Japan), and then subcultured them with five passages. Cells of line C were enlarged and flattened, and their proliferative activity was reduced after freezing and thawing. However, cells of line D appeared to be normal and maintained proliferative activity even after freezing and thawing. A TUNEL assay (In situ cell death detection kit; Roche, Basel, Switzerland) revealed that most (71.4 to 75.0%) of the nuclei in cells of line C were apoptotic, while only a small percentage of the nuclei in cells of line D were apoptotic (1.3% to 5.4%), and the latter percentage did not differ significantly from that obtained with serum-starved control cells (4.1%, p < 0.05, chi-square test). Moreover, chromosomal numbers were normal (60) in most (55%, 38/69) of the cells of line D.

Cloning Yasufuku

We produced SCNT embryos by electrofusion of cells of three different cell lines (A, B and D) with enucleated bovine oocytes. SCNT blastocysts were produced from each cell line (Table 1). When we transferred six blastocysts that had been cloned from cell line A into recipient animals, three recipients became pregnant. Although one fetus was aborted due to mummification after six months, two developed to term and male calves were delivered. The first calf was born on 30 November 2007 and is still alive at the time of writing (Fig. 3A). The second calf from a vitrified SCNT embryo was born on 5 March 2008 (Fig. 3B), but he died after two days. When we transferred six blastocysts cloned from cell line D into six recipients, two recipients, which received vitrified embryos, delivered two healthy male calves on 22 and 31 July 2008 (Fig. 3C). DNA microsatellite analysis indicated that both the cloned calves and the aborted mummified fetus were derived from the cultured cells used as donors for SCNT, that the cells were derived from Yasufuku’s testicles, and that the cells were different from those of the recipient animals (Tables 2 and 3).

Discussion

In this study, we demonstrated that normal and actively proliferating cells can be retrieved from mammalian organs that have been frozen and stored without cryoprotectant for more than a decade and that normal healthy offspring can be produced from...
Parturition was induced by injection of prostaglandin F2α after 287 days of gestation and the recipient animal delivered this calf two days after induction. The calf’s birth weight was 18.5 kg and he remains healthy at the time of writing. (B) A male calf, derived from a vitrified SCNT embryo, that was delivered by Caesarean section on 5 March 2008, after 286 days of gestation. The calf’s birth weight was 47.5 kg; he died two days after birth. (C) Male calves derived from vitrified SCNT embryos. The calf with an ear tag “c66” was born on 31 July 2008, at 288 days of gestation. Its birth weight was 30 kg. Parturitions were induced as described above. Both remain healthy at the time of writing.

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These results indicated that the nuclei of mouse somatic cells frozen even without cryoprotectant can be reprogrammed by SCNT. The combination of SCNT and ntES cell techniques may improve the probability of obtaining cloned animals from frozen tissues or organs in which the cells have been extensively damaged. In contrast to these reports, we obtained live, actively proliferating and normal cells from an organ that had been collected from a dead animal and frozen in a −80°C freezer for 10 years, and we produced four viable cloned cattle from five pregnancies by a conventional single-step SCNT technique. Our results indicate that small numbers of organ cells had withstood and/or evaded cryoinjury from ice crystals and/or osmotic stress within the organ and that nuclei in the cells were intact after defrosting the non-cryoprotected frozen organs. Our findings are consistent with the fact that frozen morselized human bone has been found to contain live cells [18]. Cells in certain tissues may be able to withstand cryoinjury, even when injury is caused by freezing without cryoprotectant. Spermatic cords consist mainly of fat tissue, blood vessels, nerve tissue, muscle tissue and connective tissue. We do not know which of these tissues, in the present study, had contained live cells after freezing. Further studies are needed to identify the organs or tissues from which live cells can be efficiently collected.

Gametes, zygotes, embryos and/or cell cultures derived from elite livestock or endangered animals have often been cryopreserved in ‘Gene Banks’ [7,8]. Our results suggest that intact and
living mammalian cells can be readily preserved by storing organs or tissues in a −80°C freezer without any special treatment. Therefore, it might be possible to rescue viable cells from frozen tissues or organs of dead animals or extinct species. Recently, clones have been successfully produced by cross-species nuclear transfer in several endangered species [12,19,20]. These results, together with our results suggest the possibility of restoring extinct species, such as woolly mammoths, if live cells can be retrieved from an organ or animal that has been frozen in a freezer or in the Siberian permafrost [21].

Materials and Methods

Freezing testicles

Testicles after castration were immediately wrapped in aluminum foil, and then frozen in a −80°C freezer without cryoprotectant. The testicles were stored in the freezer for one to four months. Yasufuku’s testicles were taken from his scrotum 12 hours after his death, then wrapped in aluminum foil, frozen in a −80°C freezer without cryoprotectant, Ten years later, his testicles were plunged into liquid nitrogen and stored for 3 years.

Table 2. DNA microsatellite analysis in 13 polymorphic loci.

| BM1824 | BM2113 | ETH10 | ETH225 | ETH3 | INRA023 | SPS115 |
|--------|--------|-------|--------|------|---------|--------|
| Yasufuku’s frozen semen | 180,188 | 135,137 | 274,274 | 145,147 | 117,117 | 198,208 | 246,256 |
| Frozen testicles | 180,188 | 135,137 | 274,274 | 145,147 | 117,117 | 198,208 | 246,256 |
| Donor cultured cells | 180,188 | 135,137 | 274,274 | 145,147 | 117,117 | 198,208 | 246,256 |
| Mummified fetus | 180,188 | 135,137 | 274,274 | 145,147 | 117,117 | 198,208 | 246,256 |
| Recipient1 | 180,182 | 131,135 | 274,274 | 145,145 | 115,117 | 198,208 | 246,256 |
| Recipient2 | 180,180 | 123,135 | 274,274 | 145,145 | 117,117 | 198,208 | 246,256 |
| Recipient3 | 180,180 | 125,137 | 268,274 | 145,145 | 117,117 | 212,212 | 248,252 |
| Recipient4 | 182,188 | 133,135 | 272,274 | 143,147 | 115,117 | 198,212 | 252,256 |
| TGLA122 | TGLA126 | TGLA227 | TGLA53 | DIK023 | DIK010 |
| Yasufuku’s frozen semen | 148,150 | 111,117 | 77,93 | 157,167 | 90,100 | 185,195 |
| Frozen testicles | 148,150 | 111,117 | 77,93 | 157,167 | 90,100 | 185,195 |
| Donor cultured cells | 148,150 | 111,117 | 77,93 | 157,167 | 90,100 | 185,195 |
| Mummified fetus | 148,150 | 111,117 | 77,93 | 157,167 | 90,100 | 185,195 |
| Recipient1 | 148,150 | 111,117 | 77,77 | 165,177 | 90,100 | 185,195 |
| Recipient2 | 148,148 | 117,117 | 77,93 | 167,177 | 100,100 | 185,193 |
| Recipient3 | 148,150 | 111,117 | 77,93 | 157,167 | 90,100 | 185,195 |
| Recipient4 | 148,148 | 111,117 | 77,93 | 157,159 | 84,90 | 185,185 |
| Recipient5 | 148,150 | 111,117 | 77,93 | 157,167 | 90,100 | 185,195 |
| Recipient6 | 148,150 | 111,117 | 77,93 | 167,177 | 100,104 | 193,195 |
| Recipient7 | 148,150 | 111,117 | 77,93 | 157,167 | 90,100 | 185,195 |
| Recipient8 | 140,150 | 111,117 | 77,93 | 165,167 | 84,100 | 185,195 |

Table 2 continued...

Values different from those of donor cultured cells are indicated in boldface. a. Polymorphism in 13 microsatellites was examined by the Maebashi Institute of Animal Science, Livestock Improvement Association of Japan, Inc (LIAJ). b. The probability that the genotype of a non-cloned animal would completely match the donor cells at these 13 loci is less than 10^−12.

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Cell culture

Primary cultures of cells from thawed tissue were generated as described earlier [22]. Frozen tissue was thawed quickly by putting it into saline at 42°C. Thawed tissue was minced (~5-mm in diameter) and incubated at 39°C for 2 h in Dulbecco’s modified Eagle’s medium (DMEM) that contained 0.1% collagenase (Invitrogen, Carlsbad, CA, USA) and 0.2% dispase (Invitrogen, Carlsbad, CA, USA). After filtration of the resultant suspension through 250-μm nylon mesh, the filtrate was centrifuged at 250 g for 5 min. Then the precipitate was resuspended in MF-start™ medium (Toyobo, Osaka Japan) and incubated at 38.5°C in an atmosphere of 5% CO2 in air with high humidity. After incubation for five days, the medium was then replaced with AmnioMAX™II complete medium (Invitrogen, Carlsbad, CA, USA) to induce the rapid proliferation of cells. Ten days later, after outgrowths had formed, the medium was replaced with MF-medium® (Toyobo, Osaka Japan).

Nuclear transfer

SCNT by electrofusion of somatic cells with enucleated bovine oocytes was performed as described earlier [17]. The SCNT embryos were activated with 5 μM ionomycin for 5 min and then
treated with 10 μg/ml cycloheximide in modified synthetic oviduct fluid medium (mSOFM) without KH2PO4 until 6 h post fusion (hpf) at 39°C in 5% CO2, 5% O2 and 90% N2 with high humidity. Following activation, the SCNT embryos were cultured in mSOFM until 6 h post fertilization (hpf) at 39°C.

Table 3. DNA microsatellite analysis in 18 polymorphic loci.

| DNA marker | DIK069 | DIK024 | DIK102 | DIK097 | DIK106 | DIK068 | DIK039 | DIK096 | BM6026 |
|------------|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| Yasufuku’s frozen semen | 163,163 | 239,245 | 135,135 | 190,190 | 109,109 | 152,160 | 195,195 | 256,258 | 167,167 |
| Donor cultured cells | 163,163 | 239,245 | 135,135 | 190,190 | 109,109 | 152,160 | 195,195 | 256,258 | 167,167 |
| Mummified fetus | 163,163 | 239,245 | 135,135 | 190,190 | 109,109 | 152,160 | 195,195 | 256,258 | 167,167 |
| Cloned offspring1 | 163,163 | 239,245 | 135,135 | 190,190 | 109,109 | 152,160 | 195,195 | 256,258 | 167,167 |
| Cloned offspring2 | 163,163 | 239,245 | 135,135 | 190,190 | 109,109 | 152,160 | 195,195 | 256,258 | 167,167 |
| Cloned offspring3 | 163,163 | 239,245 | 135,135 | 190,190 | 109,109 | 152,160 | 195,195 | 256,258 | 167,167 |
| Cloned offspring4 | 163,163 | 239,245 | 135,135 | 190,190 | 109,109 | 152,160 | 195,195 | 256,258 | 167,167 |

DNA microsatellite analysis

Microsatellite analysis was performed, for parentage testing, for 13 microsatellites by the Maebashi Institute of Animal Science, Livestock Improvement Association of Japan, Inc. (LIAJ: Maebashi, Japan) as described elsewhere [2,24]. Independently, we investigated an additional 18 microsatellites [23] to confirm the results. Moreover, additional 11 microsatellites were examined by LIAJ to confirm the results (data not shown).

Supporting Information

Alternative Language Text S1 Japanese Translation of the Full Text by YH

Found at: doi:10.1371/journal.pone.0004142.s001 (0.07 MB DOC)

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Author Contributions

Conceived and designed the experiments: YH KS TO AI KS. Performed the experiments: YH NH ST NK KS. Wrote the paper: YH KS.
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