Fertile offspring derived from mouse spermatogonial stem cells cryopreserved for more than 14 years

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BACKGROUND: Approximately 80% of childhood cancers can now be cured but a side effect of treatment results in about one-third of the surviving boys being infertile or severely subfertile when they reach reproductive age. Currently, more than 1 in 5000 men of reproductive age who are childhood cancer survivors suffer from this serious quality of life problem. It is possible to obtain a testicular biopsy before treatment to preserve the spermatogonial stem cells (SSCs) of the male by cryopreservation, but the results of long-term storage of SSCs on their subsequent functional ability to generate normal offspring has not been examined in any mammalian species. Moreover, it will be necessary to increase the number of these cryopreserved SSCs to remove any contaminating malignant cells and assure regeneration of spermatogenesis.

METHODS AND RESULTS: In this report, we demonstrate that long-term cryopreservation (>14 years) of testis cells from mouse, rat, rabbit and baboon safeguards SSC viability, and that these cells can colonize the seminiferous tubules of recipient testes. Moreover, mouse and rat SSCs can be cultured and re-establish complete spermatogenesis, and fertile mouse progeny without apparent genetic or epigenetic errors were generated by the sperm produced.

CONCLUSIONS: These findings provide a platform for fertility preservation in prepubertal boys undergoing gonadotoxic treatments.

Key words: cryopreservation / germ cells / spermatogenesis / spermatogonial stem cells / male infertility

Introduction

Chemotherapy or radiation treatment can cause extensive gonadal damage, resulting in infertility, emphasizing the need for men and boys to safeguard their fertility prior to cancer treatment (Jahnukainen et al., 2006; Brinster, 2007; Mitchell et al., 2009; Sauvat et al., 2009; Schlatt et al., 2010; Wynn et al., 2010, 2011; Jahnukainen et al., 2011a,b). Adult males can produce sperm for cryopreservation, whereas prepubertal boys cannot. It is estimated that at least 1 in 5000 males of reproductive age currently suffer from infertility or severely compromised fertility as a result of successful treatment of childhood cancer (Hewitt et al., 2003; Gincberg et al., 2010). Cryopreservation of testicular tissue from boys before treatment may allow for future transplantation of spermatogonial stem cells (SSCs) and re-establishment of fertility (Radford, 2000; Brinster, 2007; Geens et al., 2008; Ginsberg et al., 2010). However, it is unknown whether SSCs will survive cryopreservation for more than a few months, and the success of this approach is further complicated by the small tissue biopsy size; both of which may severely restrict the number of viable SSCs capable of re-establishing spermatogenesis (Geens et al., 2008; Wu et al., 2009). If SSCs indeed survive long-term cryopreservation, they would likely require in vitro expansion prior to transplantation, yet the effect of long-term storage and culture raise safety concerns regarding the unwanted genetic and epigenetic changes that can be transferred to the offspring (Radford, 2000; Alukal and Lipshultz, 2008; Wu et al., 2009).

Fifteen years ago, we reported that mouse SSCs could be cryopreserved for up to 5 months and re-establish spermatogenesis in the testes of infertile mice (Avarbock et al., 1996). Using a similar approach, progeny can be generated following 2–7 weeks of...
cryopreservation (Kanatsu-Shinohara et al., 2003). However, in those studies the storage period was short, and the availability of donor tissue was not restricted. We hypothesized that SSCs of most species can be cryopreserved for long periods and describe here the impact of long-term cryopreservation (~12–14 years) on the viability and function of SSCs from mouse, rat, rabbit and baboon.

Materials and Methods

Germ cell cryopreservation

Donor testis cells were isolated and cryopreserved between 1995 and 1998 as previously described (Avarbock et al., 1996). In brief, testes from ZFLacZ mouse (n = 53 mice), rat (n = 2), rabbit (n = 1) and baboon (n = 1) were enzymatically digested, and the donor testis cells resuspended in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal bovine serum (FBS), 2 mM glutamine, 6 mM lactate, 0.5 mM pyruvate, 30 units/ml penicillin and 50 μg/ml streptomycin (termed DMEM-c) at a concentration of 16–40 × 10^6 cells/ml. In drop-wise amounts, freezing medium comprising FBS, DMEM-c and dimethylsulfoxide (DMSO) (in a 1:3:1 ratio) was added to equal volumes of ZFLacZ mouse, rabbit and baboon testis cells (Avarbock et al., 1996). In contrast, ROSA mouse testes (n = 4 mice) were cut in half, placed in vials in 1× phosphate-buffered saline (PBS), and 10% DMSO added prior to freezing. Vials of cells for cryopreservation were first frozen at −70 °C and then placed in liquid N2 (−196 °C).

To thaw cryopreserved testis cells or pieces of testes, samples were removed from liquid nitrogen and immediately placed in a 34°C water bath and processed as previously described (Avarbock et al., 1996; Oatley and Brinster, 2006; Kubota and Brinster, 2008). In brief, thawed cells were washed three times in 1× Hank’s balanced salt solution (HBSS) (Invitrogen, Carlsbad, CA, USA), with a brief 7 min centrifugation (600g, 4°C) between each wash step in order to remove residual DMSO. Cryopreserved testis pieces derived from adult (~3 months old) ROSA mice were thawed, the tunica was removed and then digested for 10 min at 37°C in 0.25% trypsin EDTA (Invitrogen) containing 7 mg/ml DNase I (Sigma, St. Louis, MO, USA). Digestion was inhibited by the addition of 10% FBS (Invitrogen). Thawed testis cells for mouse, rat, rabbit and baboon were collected by centrifugation (7 min, 600g, 4°C) and resuspended in HBBS and then centrifuged (7 min, 600g, 4°C) twice through a sedimentation gradient comprising 30% Percoll (Sigma). Cells were again washed three times in HBSS to remove residual Percoll. Dye exclusion using trypan blue was used to determine the number of viable cells present in the cell suspension following thawing and after centrifugations in Percoll. This process resulted in a testis cell population with viability greater than 93% (Table I).

For mouse studies, testis cell suspensions were derived from either prepubertal (age 6–14 days) ZFLacZ pups or adult (~3 months old) ROSA mice. All testis cells were resuspended in mouse serum-free medium (mSFM) containing human gial cell-derived neurotrophic factor (GDNF) (20 ng/ml; R&D Systems, Minneapolis, MN, USA), rat GDNF family receptor α1 (GFRα1)-Fc fusion protein (150 ng/ml; R&D Systems) and human basic fibroblast growth factor (bFGF) (1 ng/ml; BD Biosciences, San Diego, CA, USA) (Oatley and Brinster, 2006; Kubota and Brinster, 2008). Both mouse strains contain the transgene β-galactosidase (LacZ); however, ROSA mice β-gal expression is constitutive (Nagano et al., 1999), whereas in ZFLacZ mice, β-gal expression is under the control of the zinc-finger gene on the Y chromosome (ZPY) promoter and expressed first in round spermatids (Zambrowicz et al., 1994). Cryopreserved rat testis cells were derived from prepubertal (~1 month) MTLacZ rats carrying the LacZ transgene under the control of the mouse metallothionein I (MT) promoter. The LacZ transgene is first expressed in spermatocytes in MTLacZ donor rats (Clouthier et al., 1996).

Testis cell transplantation

Recipient mice for transplantation were prepared as previously described (Brinster and Avarbock, 1994; Brinster and Zimmermann, 1994). In brief, adult mice were treated with the alkylating agent busulfan (50–60 mg/kg; Sigma), at least 6 weeks prior to transplantation of donor cells. For donor testis cells derived from ROSA or ZFLacZ mice, immunocompatible 129/SvCP × C57BL/6 or SJL × C57BL/6 F1 hybrid mice, respectively, were used as recipients (Brinster and Avarbock, 1994; Nagano et al., 1999). Immunodeficient Ncr Swiss nude mice (Taconic, Hudson, NY, USA) were used as recipients for transplantation of rabbit, baboon or MTLacZ rat testis cells. For transplantation into recipients, donor mouse testis cells were resuspended in mSFM at a concentration of 1 × 10^7 cells/ml (ZFLacZ) and 1 × 10^6 cells/ml (ROSA), and 10 μl of the donor testis cell suspension was microinjected into each recipient testis via the efferent duct, as previously described (Oatley and Brinster, 2006; Kubota and Brinster, 2008). Donor testis cells from rabbit, baboon or MTLacZ rats were resuspended in rat serum-free medium (rSFM) (Oatley and Brinster, 2006; Kubota and Brinster, 2008) at a concentration ranging from 3 to 5 × 10^7 cells/ml and 10 μl of the donor testis cell suspension was microinjected into each testis of a busulfan-treated (50 mg/kg) immunodeficient Ncr Swiss nude recipient mouse (Taconic) (Dobrinski et al., 1999; Nagano et al., 2001). Testis cells derived from either rabbit or baboon were each transplanted into five Ncr Swiss nude recipient mice. At the time of transplantation testis cell viability was ≥93% (Table I).

To compare differences in spermatogenesis between cryopreserved SSCs and freshly isolated SSCs, busulfan-treated recipient mouse testes were transplanted with freshly isolated or cryopreserved (~12 years) donor testis cells derived from ROSA mice (Table I). Two months after transplantation, the transplanted testes were removed and stained with 5-bromo-4-chloro-3-indolyl b-D-galactoside (X-gal). Colony number and colony length were determined using a dissecting microscope (Leica Microsystems, Bannockburn, IL, USA) and a Scale Master Classic (Scale Master Tech., NV, USA) measuring instrument (Fig. 1D). To further evaluate spermatogenesis, recipient mice were also transplanted with cryopreserved (~14 years) ZFLacZ testis cells (Table I). Three to 6 months after transplantation, the testes were stained with X-gal and the seminiferous tubules examined for positive X-gal staining, representing colonies of spermatogenesis. Histological sections of representative colonies in testes, stained with X-gal, were also prepared.

The impact of long-term cryopreservation on rabbit and baboon SSCs was also assessed using functional transplantation assays. Prior to transplantation into mice, rabbit testis cells were labeled with the fluorescent dye PKH26 (Sigma) according to the manufacturer’s directions. At 1 week and 2 months after transplantation, the recipient testes were removed and examined for clusters of red fluorescent cells. Specifically, tubules from these recipient testes were teased apart and clusters of red fluorescent cells were observed using a Nikon Eclipse TS100 microscope fitted with a Nikon Coolpix 4500 digital camera. Cryopreserved baboon testes cells transplanted into recipient testes were also assessed 1 week and 2 months after transplantation. The testes of recipient mice transplanted with baboon testis cells were removed, the tubules teased apart and immunostained for the expression of UCHL1 (PGP9.5; AbD Serotec, Raleigh, NC, clone #7863-0504), a protein previously shown to be expressed in spermatogonia of multiple species (Luo et al., 2009). The detection of primary antibody was carried out using a commercial horse-radish peroxidase detection kit (Histostain-SP; Invitrogen). Baboon germ cells contained within recipient tubules were visualized under bright field using the same microscope described above. All animal
Long-term freezing of spermatogonial stem cells

experimentation protocols were approved by the Institutional Animal Care and Use Committee of the University of Pennsylvania.

**Intracytoplasmic sperm injection**

The testes of busulfan-treated SJL × C57BL/6 recipient mice transplanted with total testis cells derived from ZFlacZ mice were isolated 3–6 months after transplantation (four testes from two recipient mice were used for each microinsemination experiment; mouse #A6090 and A6091). The testes were harvested, the tunica removed and the tissue washed twice with 1 × HBSS. Testes were incubated with X-gal solution, without any fixation, for 30–60 min, until positive, light-blue staining revealed colonies of spermatogenesis generated from ZFlacZ stem cells. The testes were rapidly washed three times in 1 × HBSS and the stained seminiferous tubules were carefully dissected and pooled. The tubules were digested in 0.25% trypsin EDTA (Sigma) containing 1 mg/ml collagenase IV (Sigma) for 15 min with occasional pipetting to disperse cells. Digestion was inhibited by 10% FBS and the cell suspension was filtered through a 40 μm cell strainer; cells were then washed twice by centrifugation (7 min, 600 g, 4°C) using 1 × HBSS. The cell pellet was suspended in nuclear isolation medium (NIM) containing 6% polyvinyl alcohol (PVA) and sonicated (power-setting = 2; Heat Systems-Ultrasonic, model #W-220F) for 10 s to remove the sperm tails (Kimura and Yanagimachi, 1995). Cells were placed in drops of NIM on a dish containing mineral oil, and individual sperm heads were collected by micromanipulation under the microscope. The donor-derived sperm heads were microinjected into oocytes derived from dilute brown non-agouti (DBA) × C57BL/6 F1 mice using a piezoelectric actuator (PrimeTech, Ibaraki, Japan). Injected oocytes were cultured overnight in Brinner’s medium at 37°C in a humidified atmosphere of 5% O2, 5% CO2 and 90% N2, before being transferred to the oviducts of Day 1 pseudo-pregnant ICR Swiss female mice, as previously reported (Brinner, 1965; Kimura and Yanagimachi, 1995).

**Cell culture**

*In vitro* culture of mouse and rat SSCs was carried out, as previously described (Oatley and Brinster, 2006; Kubota and Brinster, 2008). Following thawing and enrichment of viable cells through a Percoll sedimentation gradient as described above, the SSCs contained within cryopreserved testis cells were enriched using Thy-1 Antibody-Conjugated Magnetic Microbeads (MACS; Cat#130-042-201, Miltenyi Biotec, Auburn, CA, USA). All cultures were maintained at 37°C, 5% CO2 in air (Oatley and Brinster, 2006; Kubota and Brinster, 2008). The donor-derived sperm heads were microinjected into oocytes derived from dilute brown non-agouti (DBA) × C57BL/6 F1 mice using a piezoelectric actuator (PrimeTech, Ibaraki, Japan). Injected oocytes were cultured overnight in Brinner’s medium at 37°C in a humidified atmosphere of 5% O2, 5% CO2 and 90% N2, before being transferred to the oviducts of Day 1 pseudo-pregnant ICR Swiss female mice, as previously reported (Brinner, 1965; Kimura and Yanagimachi, 1995).

**DNA isolation**

A small piece of mouse liver was used to isolate genomic DNA. Tissue was incubated with proteinase K overnight and then DNA was isolated using the Qiagen DNeasy Blood & Tissue Kit (Qiagen, Valencia, CA, USA).

**Array-based comparative genomic hybridization**

To evaluate the possible deleterious effects prolonged cryopreservation and assisted reproduction technologies (ARTs) may have on the DNA of F0 progeny, we used array-based comparative genomic hybridization (CGH) for analysis of genomic DNA. ICSI-derived progeny were generated from *ZFlacZ* sperm (DBA × C57BL/6 oocyte), where the ZFlacZ transgenic mice were originally generated from a SJL × C57BL/6 background. A total of five ICSI-derived mice (three males and two females) that expressed the β-galactosidase transgene were used in the analysis. To ensure that the reference DNA encompassed all the mouse genetic backgrounds, three groups of DNA were collected for CGH as ‘pooled reference DNA’. Specifically, DNA, in equal amounts, from adult mice of C57BL/6, SJL and DBA strains were pooled as the ‘pooled reference DNA’. DNA from adult, wild-type ZFlacZ progeny was collected as ‘control wild-type DNA’; DNA from ZFlacZ progeny derived from cryopreserved testis cells via ICSI was collected as ‘cryopreserved ICSI-derived DNA’. An appropriate amount of DNA was amplified.

<table>
<thead>
<tr>
<th>Table I Testis cells stored at −196°C for ~11–14 years.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Species</strong></td>
</tr>
<tr>
<td>-----------------</td>
</tr>
<tr>
<td>Mouse (Mus musculus)</td>
</tr>
<tr>
<td>Mouse (Mus musculus)</td>
</tr>
<tr>
<td>Rat (Rattus norvegicus)</td>
</tr>
<tr>
<td>Rabbit (Oryctolagus cuniculus)</td>
</tr>
<tr>
<td>Baboon (Papio anubis)</td>
</tr>
</tbody>
</table>

DMEM, Dulbecco’s Modified Eagle’s Medium; FBS, fetal bovine serum; DMSO, dimethylsulfoxide; PBS, phosphate-buffered saline.

Details in ‘Materials and Methods’ section.

Half pieces of whole testes were cryopreserved in freezing medium. Following thawing, tissue pieces were digested to obtain a cell suspension as described in Materials and Methods.

Percentage of frozen testis cells recovered following several wash steps, including centrifugation through two separate Percoll density gradients (see ‘Materials and Methods’ section): ZFlacZ transgenic = 9.8%; ROSA lacZ transgenic = 1.1%; Rat = 4.25%; Rabbit = 14.8% and Baboon = 12.4%.
according to the Agilent array CGH protocols prior to final labeling, microarray processing and feature extraction. 4 × 180 K SurePrint G3 CGH Microarray Kits (Agilent Technologies, Wilmington, DE, USA, Cat# G4125A), were applied. Agilent feature extraction files were imported into Partek Genomics Suite v6.5. Signals were processed from extracted green (Cy3) and red (Cy5) for each position on each array. ‘Loess’ normalization per array-position was applied, which resulted in normalized intensities for each experimental sample versus the ‘pooled reference DNA’ (Cy3 labeled). Inter-sample variation was visualized using principal component analysis ≥ observed gender difference as the only defined attribute that separated the samples. Grossly, ‘cryopreserved ICSI-derived progeny’ versus ‘control wild-type progeny’ did not separate samples, and no gross differences were observed for all chromosomes from plotted averages of probe intensities for ‘cryopreserved ICSI-derived progeny’ versus ‘cryopreserved ICSI-derived progeny’. For detection of significant regions differentiating ‘cryopreserved’ versus ‘control’, no regions were found to have detectable difference at default parameters (P ≤ 0.01, fragment length 300, min 10 probes, 10% high and low excluded).

DNA methylation analyses

A total of 1 μg of DNA was bisulfite-treated using the EpiTect Bisulfite Kit (Qiagen) following the manufacturer’s protocol. The Luminometric Methylation Assay was performed and analyzed as previously described using 500 ng of genomic DNA and performed in quadruplicate (Pilsner et al., 2010). Pyrosequencing for methylation analysis of the differentially methylated regions (DMRs) of H19 and Snrpn was performed using 50 ng of bisulfite-treated DNA for PCR reactions. The H19 primer sequences were as follows: forward primer 5′-GGTTAGGGATATGTATTTTTAGGTT-3′, biotinylated reverse primer 5′-CTCATAAACCATACTAAACAT-3′. The Snrpn primer sequences were as follows: forward primer 5′-GGTAGTTTGTGGTAGGATAT-3′, biotinylated reverse primer 5′-ACTAAATCCACAACAACCCACACTAA CCT-3′. The PyroMark PCR Kit (Qiagen) was used following the
manufacturer’s protocol in a 25 μl reaction. PCR conditions were: 95°C for 15 min followed by 45 cycles of 95°C for 15 s, 55°C for 30 s and 72°C for 15 s. Each sequencing assay was carried using 10 μl of the biotinylated PCR product with either the H19 sequencing primer 5’-TGTTAAAGATTGTTGTG-3’ or Snp1 sequencing primer 5’-AAAAATGTGAGTATGTTAGTTA-3’. Pyrosequencing was performed using PyroMark Q96MD (Qiagen) system following the manufacturer’s protocol and the PyroMark Gold 96 Reagents Kit (Qiagen). Methylation was analyzed using Qiagen’s Pyro Q-CpG software. All assays analyzing DNA methylation at imprinted genes were performed in duplicate.

Cryopreservation studies
We used testes cells from 6- to 14-day-old ROSA mouse pups and examined cryoprotectants previously reported to aid post-thaw viability. We initially tested the effect that the cryoprotectants DMSO and glycerol have on cell viability. To mSFM medium, which contained 0.2% bovine serum albumin (BSA; MP Biomedical, Solon, OH, Cat# 194774, Lot# R14551), GDNF, GFRα1 and bFGF, either DMSO (10% final concentration) or glycerol (1 M final concentration) was added. In addition, FBS (20% final concentration) was added to one-half of the tubes containing DMSO or glycerol. Cells were frozen at an uncontrolled or at two different controlled rates (i.e. Protocols 1 and 2), as previously reported (Kvist et al., 2006; Keros et al., 2007). The controlled rate of freezing was obtained using the BioCool II controlled rate freezer (FTS systems Inc., Warminister, PA, USA). In Protocol 1 (Keros et al., 2007), the rate of cooling was 1°C min⁻¹ with a 5 min hold at 0°C, followed by cooling at a rate of 0.5°C min⁻¹ to −8°C and held for 10 min, before being cooled to −40°C at a rate of 0.5°C min⁻¹. The viols were held for 10 min at −40°C following which cell freezing continued to −70°C at a rate of 7°C min⁻¹ (Keros et al., 2007). Protocol 2 was adapted from Kvist et al. (2006), with some modifications. In brief, freezing of testes cells was initiated at a rate of 2°C min⁻¹ to −9°C, followed by a 5 min hold time. Cells were frozen to −40°C at a rate of 0.3°C min⁻¹, following which cells were further frozen to −70°C overnight. The uncontrolled rate of freezing was obtained by placing testes cells in a Nalgene Cryo Freezing container (Thermo Fisher Scientific Inc., Pittsburgh, PA, USA) in a −70°C freezer overnight, which results in a freeze rate of ~1°C min⁻¹, as described by manufacturer. In all cases, cells were subsequently stored for 48 h in liquid N₂ (~196°C). Following thawing in a 34°C water bath, testes cells were washed several times with 1 × HBSS, and viability was determined by trypan blue dye exclusion. Sucrose (0.07 M final concentration) and Y-27632, a selective inhibitor of the Rho-associated protein kinase p160 ROK (10 μM final conc., Calbiochem, San Diego CA, USA), were also evaluated for their added ability to safeguard SSCs through the freezing process (Gauthaman et al., 2010). In addition, the cryoprotectant activity of polyvinylpyrrolidone (1% PVP; Sigma, cat# P-0930) was assessed as a replacement for FBS (Kim et al., 2008). To investigate the effect that different freezing conditions have on SSC viability, thawed testis cells were transplanted into recipient testes as described above.

Statistical analyses
All data were processed using the Statistical Package for the Social Sciences, v18.0 (SPSS Inc., Chicago, IL, USA), and are present as mean ± SEM, except for the CGH array experiments. To compare the difference between groups, univariate ANOVA was conducted using a Tukey’s honest significant difference post hoc analysis. A difference was considered significant when the P-value was ≤0.05.

Results
Testis cells from mouse, rat, rabbit and baboon were stored at ~196°C between 1995 and 1998. Upon thawing, the viability of testis cells was low, with viability ranging from 1.5 to 17% (Table I). We attributed this low testis cell viability to a lack of robust freezing methods that had not yet been developed at the time of freezing. Nonetheless, viable testis cells could be enriched to greater than 93% from these samples by centrifugation through Percoll (Table I). In these studies, we used two mouse models, ROSA and ZFlacZ, both expressing the lacZ transgene (see ‘Materials and Methods’ section). Transplantation of donor-derived SSCs into recipient testes is the only method for unequivocally demonstrating SSC presence (Brinster and Avarbeck, 1994), and to provide an assessment of the effect of cryopreservation on donor testis cell populations that would be comparable with cells derived from prepubertal human males, testis cells from ZFlacZ mouse pups (age 6–14 days), frozen for ~14 years, were thawed and transplanted into recipient testes of mice in which endogenous spermatogenesis had been destroyed (Table I). Three months after transplantation, LacZ expression was determined by X-gal staining of tubules, and we observed that donor-derived SSCs generated colonies of spermatogenesis (Fig. 1A). To evaluate the dynamics of donor SSC colonization in recipient testes, previously frozen (~12 years) testis cells from adult (~3 months old) ROSA mice were transplanted into the testes of busulfan-treated recipient mice. Two weeks after transplantation, X-gal staining of recipient testes showed that cryopreserved (~12 years) ROSA mouse testis cells formed cell–cell associations in transplanted recipients comparable with recipients transplanted with freshly isolated ROSA mouse testis cells (Fig. 1B). Two months after transplantation, the

### Table II Fertility of ZFLacZ F₀ mice progeny generated by ICSI or by natural mating from donor cells cryopreserved for ~14 years.

<table>
<thead>
<tr>
<th>F₀ mouse #</th>
<th>ICSI or natural mating</th>
<th>F₀ Sex</th>
<th>Offspring*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ICSI</td>
<td>♂♂</td>
<td>9 pups in one litter (5♂, 4♂♂)</td>
</tr>
<tr>
<td>2</td>
<td>ICSI</td>
<td>♂♂</td>
<td>11 pups in one litter (7♂, 4♂♂)</td>
</tr>
<tr>
<td>3</td>
<td>ICSI</td>
<td>♂♂</td>
<td>9 pups in one litter (6♂, 2♂♂)</td>
</tr>
<tr>
<td>4</td>
<td>ICSI</td>
<td>♀♀</td>
<td>23 pups in two litters (11♂, 12♂♂)</td>
</tr>
<tr>
<td>5</td>
<td>ICSI</td>
<td>♀♀</td>
<td>18 pups in two litters (7♂, 11♂♂)</td>
</tr>
<tr>
<td>6</td>
<td>Natural mating</td>
<td>♀♀</td>
<td>7 pups in one litter (2♂, 5♂♂)</td>
</tr>
<tr>
<td>7</td>
<td>Natural mating</td>
<td>♀♀</td>
<td>26 pups in two litters (12♂, 14♂♂)</td>
</tr>
</tbody>
</table>

*The testes of recipient mice (#A6090 and A6091) were transplanted with cryopreserved (~14 year) ZFLacZ donor testis cells. Blue-stained, LacZ positive testis cells from both mice were collected and pooled for generation of ICSI-derived progeny (F₂ mouse ID# 1–5). ICSI-derived zygotes were transferred into two pseudo-pregnant ICR Swiss females to produce progeny; the three male F₂ progeny were derived from one female (F₁ mouse ID # 1–3), and the two female F₂ progeny were from the second female (F₁ mouse ID # 4, 5). A recipient male (#A6026) transplanted with cryopreserved ZFLacZ donor testis cells was housed with females and produced two F₂ progeny through natural mating (F₂ mouse ID # 6 and 7).

 Females often lose one or two pups in their first litter and in cases where total litter size does not correspond to the individual numbers of ♂ and ♀ progeny per litter; it is because sex was not determined until weaning (~3 weeks).
number of colonies of spermatogenesis produced in recipient testes transplanted with cryopreserved (∼12 years) or freshly isolated ROSA testis cells was 2.2 ± 0.58 colonies/10⁴ cells transplanted and 3.5 ± 0.67 colonies/10⁴ cells transplanted, respectively (Fig. 1C, mean ± SEM, P = 0.19). Moreover, the length of colonies generated from transplanted SSCs represents a critical assessment of SSC functional ability (Ryu et al., 2006), and no significant difference was observed in the length of spermatogenic colonies in recipient seminiferous tubules transplanted with either cryopreserved (2.83 ± 0.22 mm) or freshly isolated (2.48 ± 0.23 mm) ROSA mouse testis cells (Fig. 1D; P = 0.65). These findings indicate that long-term cryopreserved SSCs are able to re-establish normal spermatogenic colonies that are comparable with those from freshly isolated SSCs.

To demonstrate unequivocally that functional sperm are produced from cryopreserved (∼14 years) mouse testis cells, colonies of spermatogenesis derived from the transplantation of previously frozen ZFlacZ donor testis cells (Fig. 1E) were isolated, digested and sperm heads collected for ICSI into oocytes (Fig. 1F). After 24 h incubation, 2-cell zygotes developed, which were transferred into surrogate females. The efficiency of generating 2-cell zygotes using sperm derived from freshly isolated testes in two control experiments was 21/40 and 38/44 (2-cell zygotes/total eggs injected). In three

Figure 2 Analyses of F₀ progeny generated by ICSI revealed no significant genetic or epigenetic alterations. (A) To measure changes in DNA copy number between F₀ progeny generated by ICSI and control wild-type progeny obtained through natural mating we used array-based CGH array analysis. Using the statistical parameters defined in the Materials and Methods, no significant chromosomal deletions or duplications were observed between these two groups. (B) To further elucidate possible alterations in the genome of F₀ progeny generated by ICSI, we evaluated changes in whole genome DNA methylation using the luminometric methylation assay. The methylation percentages of ICSI-generated progeny (72.1 ± 0.44%) were not significantly different from control wild-type mice obtained through natural mating (70.7 ± 0.14%). Pyrosequencing of H19 (C) and Snrpn (D) differentially methylated regions, comparing DNA methylation percentages between ICSI-derived progeny (49.6 ± 1.86 and 38.5 ± 1.96%, respectively) and control wild-type mice obtained through natural mating (49 ± 1.90 and 42 ± 1.12%, respectively) also did not yield any significant difference between groups. Values are mean ± SEM.
experimental injection experiments, the development of 2-cell zygotes using sperm derived from recipient testes transplanted with cryopreserved (≏14 years) testis cells was 27/38, 24/71 and 27/76, respectively (2-cell zygotes/total eggs injected). The difference between the two experimental groups was not significant (P = 0.319). Importantly, sperm derived from the transplantation of previously frozen ZFlacZ donor testis cells produced a total of five pups (three males and two females) that carried the lacZ transgene, which was present in donor testis cells (Fig. 1G). All of the mice appeared normal (Fig. 1H). Additionally, two males were produced through natural mating of a recipient male transplanted with cryopreserved testis cells. Importantly, these F_0 males and females were fertile and produced fertile F_1 generations that appeared normal (Table II).

ART and germ cell cryopreservation raise concerns that deleterious genetic and epigenetic changes may be introduced into offspring (Alukal et al., 2008). Therefore, we screened the genomes of F_0 progeny (three males and two females) generated by ICSI that expressed the LacZ transgene using CGH array analysis and compared this with control wild-type mice (Fig. 2A). To assess possible epigenetic aberrations, we evaluated genome-wide DNA methylation in ICSI and control progeny and found no differences between the two groups (Fig. 2B). Additional pyrosequencing DMRs in the H19 and Snrpn imprinted loci, which are associated with the Beckwith–Wiedemann syndrome and Prader–Willi syndrome (Bartolomei, 2009; Mak et al., 2010), respectively, also showed no significant difference between ICSI and control progeny (Fig. 2C and D). These results indicate that ICSI-derived mouse offspring generated from SSCs cryopreserved for ≏14 years were not significantly different from natural offspring in these important genetic and epigenetic characteristics.

The efficient recovery of SSCs from prepubertal human testis biopsies for later transplantation and restoration of fertility is limited by the size of the initial tissue sample, as well as the small number of testis cells that possess stem cell potential (≏1 SSC is present in 3500 testis cells) capable of re-establishing spermatogenesis (Tegelenbosch and de Rooij, 1993; Wu et al., 2009). To overcome this obstacle, the SSC population contained within the testis biopsy will require in vitro expansion prior to transplantation in order to enable successful restoration of fertility. Since the in vitro conditions for rodent SSC culture are well defined (Kubota and Brinster, 2008), we evaluated the ability for...
mouse and rat testis cells cryopreserved for ~12–14 years to proliferate in culture. Cryopreserved ROSA (~12 years) and ZFLacZ (~14 years) mouse testis cells, as well as MT LacZ (~14 years) rat testis cells generated characteristic clumps of germ cell cultures containing SSCs. Cryopreserved ROSA (~12 years) and ZFLacZ (~14 years) mouse testis cells, and MT LacZ (~14 years) rat testis cells, following

Table III Comparison of DMSO and glycerol as a cryoprotectant for testis cells.

<table>
<thead>
<tr>
<th>Cryoprotectants</th>
<th>Freezing protocol</th>
<th>% cell viability Protocol 1</th>
<th>% cell viability Protocol 2</th>
<th>% cell viability uncontrolled rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>10% DMSO</td>
<td>1 week</td>
<td>53.06 ± 6.97</td>
<td>71.04 ± 1.45</td>
<td>71.81 ± 1.56</td>
</tr>
<tr>
<td></td>
<td>2 months</td>
<td>71.64 ± 1.63</td>
<td>73.03 ± 1.87</td>
<td>73.38 ± 3.17</td>
</tr>
<tr>
<td>1.4 M glycerol</td>
<td>1 week</td>
<td>10.89 ± 2.11</td>
<td>10.90 ± 1.15</td>
<td>13.62 ± 3.41</td>
</tr>
<tr>
<td></td>
<td>2 months</td>
<td>17.28 ± 1.63</td>
<td>15.49 ± 1.82</td>
<td>16.29 ± 1.79</td>
</tr>
</tbody>
</table>

*All cryoprotectants were in mouse serum-free medium (mSFM) with added growth factors glial cell-derived neurotrophic factor (GDNF), GDNF family receptor α1 (GFRα1) and basic fibroblast growth factor (bFGF). (mean ± SEM, n = 3).

Protocol 1 = Keros et al. (2007); Protocol 2 = Kvist et al. (2006). Values with different superscripts are significantly (P < 0.05) different (Tukey’s honest significant difference).
Y-27632 to the freezing medium although the mode of action requires additional experimentation.

The original freezing medium contained a small amount of BSA (0.1%) from the mSFM used for SSC maintenance and culture and a significant amount of FBS (20%), which would be an undesirable component of medium used for cells to be reintroduced into humans. Therefore, we tested the effect of PVP as a possible replacement for FBS in the freeze medium on testis cell viability and found that removal of FBS results in a small (13.7%) but significant reduction in testis cell viability. The removal of sucrose or Y-27632 had no additional effect in the absence of FBS. However, addition of 1% PVP resulted in a large (30.2%), significant reduction in testis cell viability in the absence of FBS.

The generation of donor-derived spermatogenesis following transplantation of cryopreserved testis cells provides a critically important parameter to measure SSC survival, which may be different from total testis cell viability, and is the critical end-point. We tested three of the original five freezing media in Fig. 3D to specifically examine the influence of FBS and PVP on SSCs. The absence of FBS had no effect on the number of SSCs present but the presence of PVP significantly decreased colony formation, a definitive measure of SSC number (Fig. 3E). Therefore, while FBS in the freezing medium significantly decreased the viability of mouse testis cells (Fig. 3D), the viability of mouse SSCs was not affected by the absence of FBS (Fig. 3E). A freeze medium based on mSFM with BSA (0.12%), 10% DMSO, sucrose and Y-27632 resulted in 60.3% viability of cryopreserved prepubertal mouse testis cells and generation of 32.27 ± 1.63 colonies per 10^5 cryopreserved testis cells transplanted (Fig. 3D and E). These parameters may serve as a useful framework for clinical applications involving freezing of prepubertal human testis cells.

**Table IV** Comparison of sucrose and Y-27632 as cryoprotectants for testis cells.

<table>
<thead>
<tr>
<th>Cryoprotectants*</th>
<th>Freezing protocol</th>
<th>% cell viability Protocol 2</th>
<th>% cell viability uncontrolled rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>10% DMSO + 20% FBS</td>
<td>63.38 ± 1.85a</td>
<td>68.80 ± 1.31a</td>
<td></td>
</tr>
<tr>
<td>10% DMSO + 20% FBS + 0.07M sucrose</td>
<td>69.25 ± 0.37a</td>
<td>67.59 ± 1.26a</td>
<td></td>
</tr>
<tr>
<td>10% DMSO + 20% FBS + 0.07M sucrose + 10 μM inhibitor Y-27632</td>
<td>76.89 ± 1.37a</td>
<td>79.06 ± 1.79a</td>
<td></td>
</tr>
</tbody>
</table>

*All cryoprotectants were in mSFM with added growth factors (GDNF, GFRα1 and bFGF) (mean ± SEM, n = 3).

Protocol 2 = Kivist et al. (2006). Different superscripts indicate significant (P < 0.05) differences among groups (Tukey’s honest significant difference).


**Discussion**

Gonadal damage is a relatively common consequence of treatments used to cure pediatric cancer, and male germ cells are particularly susceptible to injury by cytotoxic drugs and radiation therapy (Ginsberg et al., 2010). As treatment regimens for pediatric malignancies have improved, survivors are entering their reproductive years, and maintenance of fertility is extremely important with regard to long-term quality of life (Hewitt et al., 2003). Cryopreservation of testicular tissue, prior to the start of cancer therapy, provides a novel approach to preserve a patient’s future reproductive potential (Goossens and Tournaye, 2007; Jahnukainen et al., 2007, 2011a,b). Unfortunately, this approach is currently a limited option for many parents and patients because many practical and ethical concerns need to be addressed. Since 10 years or more may elapse before a cryopreserved testicular tissue biopsy is recovered to reestablish fertility, it is essential to understand the impact that long-term cryopreservation may have on the viability and function of SSCs. Our previous studies have shown that following cryopreservation of mouse testis cells for several months the SSCs within the cell population can restore spermatogenesis in the testes of recipient mice (Avarbock et al., 1996). However, longer periods, comparable to those that would be necessary in the human clinical setting, have not been evaluated for the ability of the frozen stem cells to reestablish spermatogenesis and produce viable, fertile progeny. Moreover, possible genetic and epigenetic abnormalities that may occur from cryopreservation have not been evaluated. Our current findings demonstrate that following long-term (~12–14 years) cryopreservation the SSC population of mouse, rat, rabbit and baboon remains viable. Cryopreserved mouse and rat testis cells reestablished spermatogenesis in recipient testes, and recipients of cryopreserved mouse testis cells could generate normal appearing, fertile progeny without apparent genetic or epigenetic errors. These findings provide experimental evidence for physicians to use in counseling parents and young patients regarding the usefulness of a testis biopsy to preserve the potential for future fertility.

Since long-term cryopreservation of testis cells and SSCs is a feasible clinical option, it is critical to improve the associated methods. Whereas previous studies have attempted to define optimal methods for freezing spermatids and spermatozoa (Ogonuki et al., 2006), little is known about conditions that maximally preserve undifferentiated germ cells containing the SSC population. In particular, the use of animal sera, such as FBS, for cryopreserving human testis cells may restrict clinical application because of the possible transmission of viral and prion-related vectors (Unger et al., 2008). In this study, we demonstrate that a simple medium containing 10% DMSO, sucrose and Y-27632A maintains SSC viability and function, and that withdrawal of FBS from the freezing medium does not significantly impact thaw viability and function of cryopreserved SSCs under the conditions developed.

These findings have important clinical implications as treatment regimens for pediatric cancers continue to improve, and the data presented here strongly support the feasibility of prolonged SSC cryopreservation and subsequent restoration of fertility for prepubertal boys undergoing cancer therapy.

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**Authors’ roles**

X.W., S.M.G., and R.L.B. designed experiments. X.W. and M.R.A. performed experiments. X.W., S.M.G. and R.L.B. analyzed data and wrote the paper; L.K.A. and M.S.B. performed and analyzed DNA methylation studies. J.W.T. analyzed CGH array profile.

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**Conflict of interest**

None declared.

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