Effect of pressure at primary drying of freeze-drying mouse sperm reproduction ability and preservation potential

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Abstract

Freeze-dried spermatozoa are capable of participating in normal embryonic development after injection into oocytes and thus useful for the maintenance of genetic materials. We recently reported that long-term preservation of freeze-dried mouse spermatozoa by conventional methods requires temperatures lower than −80 °C. Successful permanent preservation of mouse spermatozoa at much higher temperatures requires thorough investigation of the freeze-drying procedure. Thus, we examined the relationship between the pressure at primary drying and the preservation potential of freeze-dried mouse spermatozoa. Three different primary drying pressures were applied to evaluate the effect of pressure on freeze-dried spermatozoa under varying storage conditions and the rate of development measured. The developmental rate of embryos to the blastocyst stage from intracytoplasmic sperm injection by freeze-dried spermatozoa without storage was 59% (337/576), 71% (132/187), and 33% (99/302) respectively. When stored at 4 °C for 6 months, the rate was 13% (48/367), 50% (73/145), and 36% (66/182) respectively. These results show that primary drying pressure is an influential factor in the long-term preservation of freeze-dried mouse spermatozoa.

Introduction

In the last decade, numerous reports have shown that freeze-dried mouse spermatozoa are capable of participating in normal embryonic development after injection into oocytes (Wakayama & Yanagimachi 1998, Kusakabe et al. 2001, Kaneko et al. 2003a,b, Ward et al. 2003, Kawase et al. 2005). Freeze-dried mouse spermatozoa are most efficiently stored for extended periods, several to tens of decades, at temperatures lower than −80 °C (Kawase et al. 2005). However, maintaining such low temperatures over a long period of time puts the samples at risk of loss from technical difficulties (e.g. power failure) and requires a relatively high initial investment. Although freeze-dried spermatozoa can be stored for 1.5 years (Ward et al. 2003), this length of time is insufficient for either the maintenance of genetically modified mouse strains or mutant mice in saturation mutagenesis projects. Long-term preservation at ambient temperatures would be ideal. The freeze-drying process is of importance, especially the primary drying process; however, no study of pressure at primary drying of spermatozoa has been reported. Here, we focused on the pressure at primary drying and found 0.37 mbar to be the optimum pressure for preservation of freeze-dried mouse spermatozoa at much higher temperatures.

Materials and Methods

Animals

F1 (B6C3F1) mice were purchased from Clea Japan (Tokyo, Japan). All the mice were housed in polycarbonate cages and maintained under a specific pathogen-free environment in light-controlled lights on from 0500 to 1900 h and air-conditioned rooms (temperature, 24 ± 1 °C; humidity, 50 ± 10%). The mice had free access to standard laboratory chow (CE-2, Clea Japan). The Institutional Animal Care and Use Committee of Chugai Pharmaceutical reviewed the protocols and confirmed that the animals used in this study were cared for and used under the Guiding Principles for the Care and Use of Research Animals promulgated by Chugai Pharmaceutical.
Freeze-drying and preservation of spermatozoa

The procedure for freeze-drying was essentially the same as that described by Kaneko et al. (2003a) and Kawase et al. (2005). The six epididymides from three B6C3F1 male mice were removed and a dense sperm mass was squeezed out of each cauda epididymis from a cut made with scissors. The total sperm mass was gently placed into 9 ml EGTA Tris–HCl-buffered solution (50 mM EGTA, 50 mM NaCl, and 10 mM Tris–HCl, pH 8.0; Kaneko et al. 2003a) in a tube (352059, Becton Dickinson Labware, NJ, USA) and kept at 37 °C for 10 min. The sperm suspension at a concentration of 15–38 × 10^6 cells/ml was divided into 18 aliquots. Each aliquot of 500 µl was put into an amber vacuum vial for freeze-drying (V-2B, Nichiden-rika Glass Co. Ltd., Kobe, Japan). The vials were plunged into liquid nitrogen (~−196 °C) for 5 min and then transferred to a programmable freeze-dryer (BETA2-16, Martin Christ Gefriertrocknungsanlagen GmbH, Osterode am Harz, Germany), which had been pre-cooled to −30 °C. The freeze-drying conditions consisted of primary drying at a pressure of 0.04 mbar for 8 h, 0.37 mbar for 13 h, or 1.03 mbar for 13 h and secondary drying at a pressure of 0.001 mbar for 6 h. More time is required for complete drying when lower pressures are provided at primary drying. The inside pressure of the vials at the time of sealing was 0.001 mbar and pressure reduction was within 5 min. The vials were stored at 4 or 30 °C until use. Immediately before ICSI, the vials of freeze-dried spermatozoa were unsealed, and the spermatozoa were hydrated by adding 500 µl sterile distilled water. To maintain a similar composition of sperm suspension before and after freeze-drying, distilled water was added rather than a medium such as Hepes-buffered culture medium.

Comet assay for DNA damage

DNA damage of the spermatozoa from freeze-drying and subsequent preservation was assessed by single-cell gel electrophoresis (comet assay; Hughes et al. 1997, Steel et al. 1999, Cho et al. 2003). Evaluation of the shape of the DNA ‘comet’ tail and migration pattern gives an assessment of DNA damage. The sperm suspension was suspended in Comet LMAgarose (1% low-temperature melting agarose, Trevigen, Gaithersburg, MD, USA) at a ratio 1:10 (v/v). With the addition of molten LMAgarose melting agarose, Trevigen, Gaithersburg, MD, USA) at a suspended in Comet LMAgarose (1% low-temperature medium such as Hepes-buffered culture medium. The sperm suspension was immediately placed on a CometSlide (Trevigen). The slides were placed flat in a refrigerator at 4 °C for 10 min and then submerged in 23 ml lysis solution (Trevigen) at 4 °C for 60 min. Next, 2.5 ml 10 mM dithiothreitol was added and the slides were then incubated for 30 min at 4 °C, followed by the addition of 2.5 ml of 4 mM LIS (lithium diiodosalicylate) and incubation for 90 min at room temperature. The slides were then kept in an alkaline solution (> pH 13) for 20 min at room temperature in the darkness. The slides were subjected to electrophoresis in 1XTBE (Tris-borate EDTA) buffer at 25 V for 10 min, stained with SYBR Green (Trevigen), and analyzed under a microscope (IX-70, Olympus Co., Tokyo, Japan). DNA damage of freeze-dried spermatozoa was assessed by Comet assay twice per experimental group. Comet tail length is the distance the damaged DNA has migrated from the sperm head and gives an estimate of the extent of the damage.

Preparation of oocytes

 Mature B6C3F1 females were induced to superovulate by i.p. injections of 5 IU equine chorionic gonadotropin (Serotrophin, Teikokuzoki Co., Tokyo, Japan) and 5 IU human chorionic gonadotropin (hCG; Puberogen, Sankyo Co., Tokyo, Japan) 48 h later. Freshly ovulated oocytes were collected from oviducts 15–16 h after being injected with hCG. The oocytes were treated with 0.1% hyaluronidase (280 units/mg; H-3506, SIGMA Chemical Co.) in Whitten’s medium (Whitten 1971) supplemented with 100 µM ethylene diamine tetraacetic acid disodium salt (EDTA; Abramczuk et al. 1977) to remove cumulus cells.

Intracytoplasmic sperm injection

After rehydration of the freeze-dried spermatozoa as described above, one part of the sperm suspension was thoroughly mixed with nine parts 0.9% NaCl solution (saline) containing 12% (w/v) polyvinyl pyrrolidone (PVP, No. 99219, Mt. 360 000, Irvine Scientific, Santa Ana, CA, USA). Two drops (~5 µl each) of 12% PVP saline and two drops of 20 mM Hepes-buffered Whitten’s medium containing 0.1% polyvinyl alcohol (PVA, P-8136, MW 30 000–70 000, Sigma) were linearly placed on the injection chamber (Kawase et al. 2001) and then covered with mineral oil (M-8410, embryo tested, Sigma). The first drop of 12% PVP saline was used to wash the injection pipette, and added to the second drop was 1–2 µl of the diluted sperm suspension. The first drop of the medium was used to remove spermatozoa that had attached to the surface of the injection pipette. The cumulus-free oocytes were placed in the second drop of Hepes-buffered Whitten’s medium. The injection chamber with the spermatozoa and oocytes was transferred onto the stage of an inverted microscope maintained at ~18 °C (MATS-555RSP, Tokai Hit, Shizuoka, Japan).

The procedure for micromanipulation of the sperm for ICSI was essentially the same as that described previously (Kimura & Yanagimachi 1995, Kawase et al. 2001). The sperm head was separated from the tail by applying three or four piezo pulses (controller setting: speed 2, intensity 2) to the head–tail junction of the spermatozoon. In the same manner, a total of 3–5
isolated sperm heads were lined up in the pipette. Several piezo pulses (controller setting: speed 2, intensity 2) were applied to advance the tip of the injection pipette to the surface of zona pellucida; the pipette was advanced mechanically while applying slightly negative pressure. The oolemma was punctured using 1–2 piezo pulses (controller setting: speed 1, intensity 1). A single sperm head was then expelled into the ooplasm accompanied with a minimum amount of medium. Following retrieval of as much as possible of the medium, the injection pipette was withdrawn while applying negative pressure to the pipette.

**Culture of oocytes and embryos transfer**

Sperm-injected oocytes were incubated and cultured in Whitten’s medium supplemented with 100 μM EDTA at 37.5 °C in 5% CO₂ and 95% air. After 6-h ICSI, live oocytes showing two distinct pronuclei and a second polar body were considered fertilized. The fertilized eggs were further cultured in Whitten’s medium supplemented with 100 μM EDTA for 96 h at 37.5 °C in 5% CO₂, 95% air. About 96 h after ICSI, the blastocysts were transferred into the uteri of pseudopregnant ICR recipients (CLEA Japan) 2.5 days postcoitum (dpc) using the embryo transfer method described by Suzuki et al. (1994). The recipient females were killed on 18.5 dpc to determine the number of implantation sites by macroscopic check and the number of term fetuses.

**Statistical analysis**

Data presented in this study were analyzed statistically by the χ²-test and Tukey’s test for non-parametric multiple comparisons (SAS version 6.12, SAS Institute, Cary, NC, USA). In all statistical tests, the difference was considered significant when P < 0.05.

**Results**

The developmental rates to the blastocyst stage of embryos from ICSI by freeze-dried spermatozoa without storage, with storage at 30 °C for 3 days, and at 4 °C for 6 months are shown in Tables 1–3. All the three storage conditions showed the highest rate of embryonic development when primary drying was performed at 0.37 mbar. In addition, the number of live-term fetuses produced was higher at 0.37 mbar than at other pressures for groups non-stored and stored at 4 °C for 6 months although the difference was not significant (Table 4). All fetuses were morphological normal – some of the mice have since been mated and delivered normal offspring. A numerical indicator for overall efficiency of mouse production after ICSI using freeze-dried mouse spermatozoa was determined by dividing the ratio of the number of embryos that developed blastocysts to the number of fertilized oocytes, by the ratio of the number of live-term fetuses to the number of blastocysts transferred (Fig. 1). With storage at 4 °C for 6 months, the overall efficiency from a pressure of 0.37 mbar (10.3) or 1.03 mbar (8.8) was significantly higher than 0.04 mbar (1.1). Moreover, with a pressure of 0.37 mbar (25.7) non-stored, the overall efficiency after ICSI was significantly higher than for 0.04 mbar (17.5) or 1.03 mbar (6.62). The results showed that blastocyst formation was significantly higher when primary drying was performed at 0.37 mbar than at 0.04 and 1.03 mbar irrespective of the storage conditions (non-stored, 30 °C for 3 days, or 4 °C for 6 months; Tables 1–3, Fig.1).

Percentages of spermatozoa with comet tails freeze-dried at 4 °C for 6 months were 100 and 96%, and stored at 30 °C for 3 days were 4 and 100% respectively. Although there was no significant difference between the percentages of spermatozoa with damaged DNA at 0.04 mbar stored at 30 °C for 3 days and of those without

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**Table 1** Effect of vacuum pressure at primary drying on the in vitro development of embryos generated by ICSI of freeze-dried, non-stored spermatozoa.

<table>
<thead>
<tr>
<th>Vacuum pressure (mbar)</th>
<th>No. of oocytes injected</th>
<th>No. (%) of oocytes survived</th>
<th>No. (%) of oocytes fertilized</th>
<th>No. (%) of embryos developed to two-cell</th>
<th>No. (%) of embryos developed to blastocyst</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.04</td>
<td>864</td>
<td>608 (70)a</td>
<td>576 (95)a</td>
<td>557 (97)a</td>
<td>337 (59)a</td>
</tr>
<tr>
<td>0.37</td>
<td>253</td>
<td>194 (77)b</td>
<td>187 (96)ab</td>
<td>181 (97)a</td>
<td>132 (71)b</td>
</tr>
<tr>
<td>1.03</td>
<td>404</td>
<td>317 (78)b</td>
<td>302 (95)a</td>
<td>288 (95)a</td>
<td>99 (33)c</td>
</tr>
</tbody>
</table>

Different superscript letters within a column indicate significantly different values (P < 0.05).

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**Table 2** Effect of vacuum pressure at primary drying on the in vitro development of embryos generated by ICSI of freeze-dried spermatozoa stored at 30 °C for 3 days.

<table>
<thead>
<tr>
<th>Vacuum pressure (mbar)</th>
<th>No. of oocytes injected</th>
<th>No. (%) of oocytes survived</th>
<th>No. (%) of oocytes fertilized</th>
<th>No. (%) of embryos developed to two-cell</th>
<th>No. (%) of embryos developed to blastocyst</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.04c</td>
<td>251</td>
<td>175 (70)a</td>
<td>170 (97)a</td>
<td>162 (95)a</td>
<td>34 (20)a</td>
</tr>
<tr>
<td>0.37</td>
<td>161</td>
<td>117 (73)a</td>
<td>113 (97)ab</td>
<td>111 (98)a</td>
<td>61 (54)b</td>
</tr>
<tr>
<td>1.03</td>
<td>187</td>
<td>145 (78)a</td>
<td>132 (91)b</td>
<td>129 (98)a</td>
<td>25 (19)b</td>
</tr>
</tbody>
</table>

Different superscript letters within a column indicate significantly different values (P < 0.05).

aPercentage of oocytes survived. bPercentage of oocytes fertilized. cData from Kawase et al. (2005).

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storage, developmental rates to the blastocyst stage after ICSI was significantly reduced with storage. In contrast, spermatozoa freeze-dried at 0.37 mbar and then assessed immediately (non-stored) or after storage at 4°C for 6 months did not have comet tails (Fig. 2, Table 5). At the primary drying pressure of 0.04 mbar, the average comet tail length of spermatozoa stored at 30°C for 3 days and at 4°C stored for 6 months was 0.7 ± 3.4 and 20.2 ± 5.0 μm (P < 0.05), and at 1.03 mbar, average tail length was 23.4 ± 6.2 and 23.0 ± 8.4 μm (P > 0.05) respectively (Table 5).

**Discussion**

Preservation of freeze-dried spermatozoa for extended periods of time, potential 100 years or more, by current methods would require storage temperatures lower than −80°C (Kawase et al. 2005). Successful long-term preservation of mouse spermatozoa at higher temperatures requires investigation and modification of the current freeze-drying method. Although the primary drying phase is one of the most important processes in the freeze-drying process of spermatozoa, there have been no studies on the primary drying pressure. Nail et al. (2002) reported that the pressure in the freeze-dryer has to be lower than the vapor pressure of ice at the temperature of the product, and pressures of 0.04, 0.37, and 1.03 mbar are appropriate for materials stored at −50, −30, and −20°C respectively. Thus, in this study, we selected primary drying pressures of 0.37 and 1.03 mbar, in addition to the commonly used 0.04 mbar. Primary drying is characterized by the specimen undergoing rapid shrinkage as the ice sublimes (ice forming water vapor and leaving the specimen). In this study, ~98% of the total volume of water (0.51 g wet weight) was lost. During this step, evaporative cooling keeps the temperatures low. When most ice has sublimed, heat is no longer lost by evaporative cooling and the temperature of the product usually increases sharply toward shelf temperature. Since the driving force for freeze-drying is the vapor pressure of the ice, from the standpoint of process efficiency, it is important to keep the product temperature as high as possible during primary drying (Nail et al. 2002). As we reported recently, freeze-dried sperm kept at a pressure of 0.04 mbar and then stored at 30°C for 3 days or at 4°C for 3 months had the same rate of blastocyst formation (Kawase et al. 2005). Although a pressure of 0.37 mbar gave the best results when...
compared with 0.04 and 1.03 mbar, a decrease in developmental rates to blastocysts and live born was seen after storage, compared with freshly freeze-dried samples (Tables 1–4). When comparing sperm dried using three pressures, it seems that sperm dried at 0.04 mbar deteriorate significantly during storage both at 30 and 4 °C (P < 0.05); sperm dried at 0.37 mbar lowers its potentials similarly during storage at 30 and 4 °C (P < 0.05); and sperm dried at 1.03 mbar deteriorate slightly at 30 °C (P < 0.05), but maintains its reproductive potentials when stored at 4 °C (P > 0.05). Although precise mechanisms affecting storage potential in pressure at primary drying of freeze-drying mouse spermatozoa was not clarified in this study, differences of an alteration of tertiary structure in seminal proteins by freeze-drying might influence their potential (Jeyendran et al. 1983). Sperm freeze-dried at 1.03 mbar yielded the same proportion of development to blastocysts and live born after transfer when injected immediately after preservation (no storage) and after 6 months storage (Tables 1, 3, and 4). However, overall efficiency at 1.03 mbar was not significantly different between the experimental groups.

Comet assay, the evaluation of the shape of the DNA ‘comet’ tail and migration pattern, is widely used to measure DNA damage in many different cell types, including sperm from laboratory animals and humans (Haines et al. 1998). In this study, we performed a comet assay using alkaline electrophoresis to measure single-stranded DNA breaks and alkali-labile sites to assess the DNA integrity of individual freeze-dried sperm. Assessment of DNA damage is determined by the percentage of spermatozoa with tails and the average comet tail length. Comet tails of substantial average length were detected by comet assay at 0.04 and 1.03 mbar stored at 4 °C for 6 months and at 1.03 mbar stored at 30 °C for 3 days (Table 5), suggesting that storage induces DNA damage in freeze-dried spermatozoa.

The overall efficiency determined from the data in this study clearly indicates that the pressure at primary drying is an important factor affecting the outcome after storage. Modifications to the freeze-drying method, with

<table>
<thead>
<tr>
<th>Primary pressure</th>
<th>0.04 mbar</th>
<th>0.37 mbar</th>
<th>1.03 mbar</th>
<th>Fresh</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of sperm with tail/total (%)</td>
<td>Average of tail length (µm)</td>
<td>No. of sperm with tail/total (%)</td>
<td>Average of tail length (µm)</td>
</tr>
<tr>
<td>Without storage</td>
<td>0/56 (0)</td>
<td>0</td>
<td>0/62 (0)</td>
<td>0</td>
</tr>
<tr>
<td>Stored at 30 °C for 3 days</td>
<td>2/55 (4)</td>
<td>0.7 ± 3.4a</td>
<td>0/54 (0)</td>
<td>0</td>
</tr>
<tr>
<td>Stored at 4 °C for 6 months</td>
<td>55/55 (100)b</td>
<td>20.2 ± 5.0b</td>
<td>0/55 (0)</td>
<td>0</td>
</tr>
<tr>
<td>Stored at 37 °C for over 1 week</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

Different superscript letters between same primary pressure indicate significantly different values (P < 0.05).
attention to the pressure at primary drying, will contribute to the success of permanent preservation of mammalian spermatozoa stored at higher temperatures.

Acknowledgements

We thank S Uchida and Y Nakajima for their technical assistance, and Ms F Ford for proofreading the manuscript. This study was supported in part by the Special Coordination Funds for Promoting Science and Technology from the Ministry of Education, Culture, Sports and Science, Japan. The authors declare that there is no conflict of interest that would prejudice the impartiality of this scientific work.

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Received 28 August 2006
First decision 18 September 2006
Revised manuscript received 29 December 2006
Accepted 9 January 2007