Improvement of the Vitrification Method Suppressing the Disturbance of Meiotic Spindle and Chromosome Systems in Mature Oocytes

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ABSTRACT: Vitrification method is widely used in oocyte cryopreservation for IVF but the birth rates are lower than that of the fresh oocyte. One of the known main reasons is structural instability of meiotic spindle and chromosome systems of mature oocyte. To get the best way for keeping competence of matured oocytes, we studied the best conditions for vitrification focused on equilibration times. The mature oocytes were underwent vitrification with current popular method and analyzed the survival rates, microtubule stability and DNA integrity. The survival rates of recovered oocyte are almost same between groups and are more than 93%. The structural configuration of meiotic spindle was well kept in 10 min equilibration group and the stability rate was almost same with that of control. The chromosomal breakdown was observed in all experimental groups, but the chromosomal stability was higher in 10 min equilibration group than the other groups. The 10 min equilibration group showed best condition compared with the other groups. Based on these results, the equilibration time is one of the key factors in successful keeping for competence of mature oocyte. Although, more fine analysis about the effects of physical stress on oocyte during vitrification is needed to define the optimal condition, it is suggested that the optimal equilibration time to get competent oocyte in mouse is 10 min. Information acquired this study may provide insight into intracellular structural events occurring in human oocytes after vitrification and application for cryopreservation of human oocyte.

Key words: Cryopreservation, Vitrification, Chromosome, Meiotic spindle, Matured oocyte

INTRODUCTION

How maintain or improve the fertility of male or female is one of the main interesting in development and reproductive biology. The methods for preservation of sperm, oocyte, embryo and reproductive tissue has been developed recently. Oocyte cryopreservation has more wide clinical implications than embryos freezing procedure (Wennerholm, 2000) because women who have no partner or have the possibility to lose their ovarian function due to surgery, chemotherapy or radiotherapy can store their oocytes for future use. Cryopreservation of the excess oocytes can avert the repeated ovarian stimulation and oocyte collection in women undergoing in-vitro fertilization (IVF) or can be a source for oocyte donation. It also gives more chance for women to achieve their social goal and provide more
reproductive choices because oocyte storage allows women to preserve the fertility for the future. Oocyte cryopreservation therefore has the potential to be an important adjunct to ART in human and other mammals. However, there has been the relatively limited preservative of mature oocyte (Veeck, 2003; Borini et al., 2004) compared with that of embryos (Mandelbaum et al., 1998).

Mature oocytes stay in metaphase block during secondary meiotic division in mammals including human. At this stage, the chromatins are condensed and become chromosome, and made specific connection with the metaphase II (M II) spindles. A spindle apparatus is a dynamic conglomerate of microtubules and associated structural proteins, acting to coordinate cytokinetic and karyokinetic events essential for normal chromosome segregation. First meiotic division is completed by cooperation of cytoskeletons, especially microfilaments and microtubules. Cytoskeleton is a key factor to get the successfully competent oocytes. Until sperm entry, the matured oocytes stay in metaphase block. This block is accomplished by stable spindle and other regulators for cell cycle. As oocytes transit into metaphase, microtubules change from radial arrays to an organized barrel-shaped bipolar structure containing a blend of dense material at either pole known as microtubules organizing centers (MTOCs).

The most important things are keeping the structural stability of both spindle and chromosomes for developmental competence. Unfortunately, many reports, however, have demonstrated that cryopreservation of oocytes may cause depolymerization and disorganization of spindle microtubules (Pickering & Johnson, 1987; Vincent et al., 1989; Aman & Parks, 1994). Oocytes are exposed to many stresses including thermal, mechanical and chemical stress during process of cryopreservation and caused the lost of survivability (Meryman, 1971; Mazur et al., 1972). In fact, the process of cryopreservation of meiotic spindles is leading to impairment of fertilization of such oocytes and the growth of embryos (Aman & Parks, 1994; Eroglu et al., 1998).

There are two methods that are used to cryopreserve the mamalian oocytes, slow-freezing and vitrification (Bernard & Fuller, 1996). High survival rates of human oocytes following slow-freezing have been achieved with sodium depleted medium and/or elevated sucrose concentrations (Fabbri et al., 2001; Bianchi et al., 2005; Borini et al., 2006; Stachecki et al., 2006). However, despite the increase in survival rates, implantation rates are still low; between 12-14% per transferred embryo (Boldt et al., 2006; Oktay et al., 2006). Recently, vitrification has been widely used for cryopreservation of ovine, equine, murine, rabbit, bovine, and porcine embryos at all stage of embryonic development, including blastocyst stages, and oocyte. It is suggested that vitrification may be more effective than slow-freezing (Kuleshova & Lopata, 2002), resulting in improved oocyte survival and pregnancy rates (Yoon et al., 2003; Katayama et al., 2003; Lucena et al., 2006).

Vitrification is a process that produces a glasslike solidification of living cells that completely avoids ice crystal formation during cooling and warming (Al-Hasani et al., 1986; Diedrich et al., 1988). However, a major concern of vitrification relates to the use of higher concentration of membrane-permeable cryoprotectants (CPA) in the suspending solution, such as dimethyl sulfoxide (DMSO), 1,2-propanediol (PROH), and/or ethylene glycol (EG). It has been known that exposure time to the high concentration of cryoprotectants during vitrification is one of the main factors influencing the oocytes. Therefore in this study, the stability of meiotic organelles was examined to evaluate the vitrification influences on the survival of mature oocyte. In addition, cryoinjury of oocytes was detected at DNA level during vitrification.

MATERIALS AND METHOD
1. Mature oocytes collection

All animals involved in this study were approved by the Animal Care Committee and studies were conducted for the Care and Use of Laboratory. To get MII stage oocyte, the female mice were injected with 5 IU of pregnant mares serum gonadotropin (PMSG; Folligon, Intervet) followed by injection with 5 IU of human chrionic gonadotropin (hCG) in 48 hrs later (i.p.). After 15 h of hCG injection, the mice were sacrificed and the oviducts were dissected and placed into a dish containing Quinn's Advantage™ Medium with HEPES (Sage Biopharma, San Clemente, CA; cat. no. 1023) supplemented with 1.0 mg/mL bovine serum albumin (BSA) (Sigma, A8022). The cumulus-oocyte complexes (COC) were released by tearing the ampullae of the oviduct. The cumulus mass were removed using both hyaluronidase (80 IU/mL) (Sigma, H3506) and mechanical force using a fine glass pipette. The denuded oocytes were washed several times with Quinn's Advantage™ Medium with HEPES supplemented with 1.0 mg/mL BSA at RT and undergone for next examination.

2. Vitrification of mature oocyte

Oocytes were cryopreserved by vitrification methods as described by Kuwayama and his colleagues (2005) with minor modifications. The solution for vitrification and warming was PBS containing cryoprotectants and 10% serum protein substitute. The equilibration solution consisted of 7.5% ethylene glycol (EG; Sigma E9129) and 7.5% dimethyl sulfoxide (DMSO; Sigma D5879). The vitrification solution consisted of 15% EG and 15% DMSO, 0.5 M sucrose (S; Sigma S7903). The oocytes were placed in 7.5% ethylene glycol EG and 7.5% DMSO for 5, 10 and 15 min. Then those were transferred into the vitrification solution (15% EG + 15% DMSO + 0.5 M sucrose). After 40–60 sec of exposure in the vitrification solution, oocytes were loaded onto straw (Fig. 1) and directly plunged into liquid nitrogen (LN2).

3. Morphological assessment of oocytes after warming

Oocytes were defined as having morphologically survived if the oocytes possessed an intact zona pelludia and plasma membrane and refractive cytoplasm. They were counted and recorded.

4. Evaluation of meiotic spindle and chromosome in oocytes

Tubulin and chromatin were stained by immunofluorescent staining, as described previously (Jack et al., 2007). Briefly, after warming the oocytes were incubated for 1 hr at 37°C in 5.0% CO2 incubator. Then the warmed oocytes were fixed in 4% paraformaldehyde for 5 minutes. After fixation, the oocytes were washed extensively in PBS and blocked overnight at 4°C in blocking medium (PBS, supplemented with 0.02% NaN3, 0.2% non-fat dry milk, 2% goat serum, 2% BSA and 0.1 moL/L glycine). After rinsing in PBS, the oocytes were incubated with β-tubulin monoclonal antibody

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**Fig. 1. Device for vitrification of oocyte.** A. Cryo vessel (modifed straw). B. Photographs showing the loaded oocytes.
(Sigma F2043) diluted 1:150 in PBS at 4°C overnight. After then, oocyte washed several times with PBS and incubated with fluorescein isothiocyanate (FITC)- conjugated anti-mouse IgG (Sigma F5387) diluted 1:150 in PBS for 60 min at 37°C. After washing with PBS for several times, the samples were mounted onto a slide under a cover slip in the Vectashield mounting medium (Vector Laboratories, Burlingame, CA), containing 4,6-diamidino-2-phenylindole (DAPI).

5. Comet assay in vitrified oocytes
Morphologically intact oocytes underwent comet assay to detect cryoinjury at DNA level. The individual oocytes were employed for comet assay that was carried out as previously described (Singh et al., 1988). All products for comet assay were obtained from Trevigen, Inc (Gaithersburg, MD). Quantitative analysis of DNA damage included the counting the number of oocytes with the presence of a comet tail.

6. Statistics
The *t*-test was used to evaluate the difference between controls and experiment groups. Results were presented as MEAN ± SD. A *p*-value less than 0.05 were considered to be a significant difference.

**RESULTS**

1. Change the volume of oocyte during equilibration and vitrification
The volumetric changes were recorded during vitrification (Fig. 2). At the time of equilibration, within 1 min after expose to dehydration solution the oocytes were shrunk (Fig. 2B) and got equilibration from 10 min (Fig. 2C, D). After treatment with vitrification solution for 40 sec, there was dramatic shrinking of the equilibrated oocytes (Fig. 2E).

![Fig. 2. Photomicrograph of subjected oocytes during dehydration and equilibration periods. A, B, C and D showed 0min, 5 min, 10 min and 15 min oocytes after treatment of equilibration solution, respectively. E is shown the oocytes which is treated in vitrification solution. F is shown the volume changes of oocytes during equilibration time.](image)

2. Morphological assessment of oocyte survival after warming
A total 445 oocytes were vitrified. After warming and washing, oocytes were assessed for cryosurvival according to the criteria mentioned in Materials and Methods (Fig. 3). Survival rates of the recovered oocytes were higher than 90% and there was no difference between experimental groups. M II oocytes after vitrification and warming were morphologically indistinguishable compared to them before cryopreservation. The recovered oocytes had normal-appearing zona pellucidae, intact polar bodies, normal perivitelline spaces, intergral plasma membranes, and normal-appearing evenly granular cytoplasm. After 1 hr of culture, the survival rates were 97.1% (102/105) and 93.5% (174/186) and 95.5% (147/154) for 5, 10, and 15 min, respectively.
Vitrification Method for Matured Oocyte

Fig. 3. Photomicrograph of survived or damaged oocytes after vitrification-warming. A showed survived oocyte and B showed damaged oocytes.

Table 1. Effect of the pre-equilibration time on survival rates of recovered oocytes after vitrification and warming of mature oocytes

| Time (min) | No. of vitrified oocytes | Survival rate (%) |
|-----------|--------------------------|-------------------|
| 5         | 105                      | 102/105 (97.1%)   |
| 10        | 186                      | 174/186 (93.5%)   |
| 15        | 154                      | 147/154 (95.5%)   |

From those results, the device for vitrification and time point were good to keep the viability of oocytes.

3. Analysis of spindle and chromosome configuration

Morphological analysis is not enough to prove the stability of competency of matured oocytes. So we analyzed the stability of spindle and chromosome. The spindle morphology and chromosomal patterns of vitrified oocytes of 1 hr incubation in 5% CO₂, in air at 37°C. The majority of the recovered oocytes maintained the normal meiotic spindle morphology and chromosome alignment (Fig. 4).

The rates of oocyte having normal spindle and chromosomal configuration were 74.3%, 81.1% and 66.3% in 5 min group, 10 min group, and 15 min group, respectively. In control group which were collected after superovulation, the rate of normal spindle and chromosomal arrangement was 87%. Compared to the control group, there were no significant difference in 5 min and 10 min group, but there was a significant decrease in 15 min group (Table 2).

Table 2. Morphological stability of meiotic spindle and chromosome in mature oocytes after vitrification-warming

| Equilibration time | No. of oocyte | Spindle and chromosome configuration (%) |
|--------------------|---------------|------------------------------------------|
|                    |               | Normal | Abnormal |
| Control            | 69            | 60 (87.0) | 9 (13.0) |
| 5 min              | 77            | 57 (74.3)  | 20 (25.7) |
| 10 min             | 90            | 73 (81.1)  | 17 (18.9) |
| 15 min             | 83            | 55 (66.3)  | 28 (33.7) |

a: p<0.05 control vs. experimental group
b: p<0.05 5 and 10 groups vs. 15 groups

The 10 min equilibration gave more stability significantly than the other groups. Compared with control and 10 min group, the stability did not change (Table 2). However, the stability of spindle and chromosomal configuration was significantly decreased in 5 min and 15 min group compared to 10 min group.

4. DNA fragmentation of vitrified and warmed oocyte

One of the well established method for the analysis of the chromosomal stability is comet method and it was applied after morphological analysis (Fig. 5). In control group, the oocytes which had comet tail were 15.3% but
Fig. 5. DNA fragmentation analysis of the mouse oocytes based on the comet tail. DNA of the Oocyte shows no fragmentation (A) and the fragmented DNA of the oocyte has migrated out from the membrane (B). The direction of electrophoresis is from right to left and the comet tail containing the DNA fragments that was stained by SYBR green.

Table 3. Analysis of DNA fragmentation after vitrification by comet assay

| Equilibration time | No. of oocytes | No. of oocytes with comet tail (%) |
|--------------------|----------------|----------------------------------|
| Control 13         | 13             | 2 (15.3)                         |
| 5 min 14           |                | 6 (42.9) a                       |
| 10 min 16          |                | 7 (43.8) a                       |
| 15 min 15          |                | 12 (73.3) a, b                   |

a: $p<0.05$ control vs. experimental group
b: $p<0.05$ 5 and 10 groups vs. 15 groups

As shown in Table 3, the percentage of oocytes with comet tails was significantly higher ($P<0.05$) in 15 min group compared to 5 min exposure group. However, there were no difference in the percentage of oocytes with comet tails between 5 min exposure group oocytes and 10 min exposure group.

DISCUSSION

A major advantage of vitrification is the elimination of mechanical damage caused by intra- or extra-cellular ice crystals and potential reduction of chilling damage by shortening the exposure to suboptimal temperatures. However, vitrification is a result of high cooling rates associated with high concentrations of cryoprotectants. If used incorrectly, these cryoprotectants can be toxic. Therefore, vitrification necessitated the reduced and precise exposure times and rapid movement of cells through solution.

An important problem with oocyte cryopreservation is the variable survival rate (Chen, 1986; Porcu et al., 1997; Boldt et al., 2003; Fosas et al., 2003). Modified slow-freezing methods have improved survival rates (Fabbri et al., 2001; Fosas et al., 2003; Bianchi et al., 2005; Borini et al., 2006; Stachecki et al., 2006), but as yet it is still unclear as to which method provides the best overall success. However, it is suggested that vitrification has advantages in various points compared with slow-freezing.

In this study, the survival rates of recovered oocytes are almost same between groups and are higher than 93%. The survival rate was higher previously reports which employed vitrification and other freezing methods (Huang et al., 2008).

At the time of ovulation, the oocyte is arrested at MII stage of meiosis with the chromosomes aligned at the equatorial plate of the meiotic spindle and the first polar body extruded. The completion of meiosis depends on the presence of intact spindle microtubules to achieve normal segregation of the chromatids. Spindle microtubules consist of polymerized tubulin, the major component of spindle
microtubules, in equilibrium with the free tubulin pool within the oolemma (Vincent & Johnson, 1992). The dynamic equilibrium between polymerized and free tubulins in mammalian oocytes is extremely sensitive to temperature change (Zenses et al., 2001). Cooling of mouse oocytes causes tubulin to undergo depolymerization and results in the disappearance of microtubule organizing centers (Magistrini & Szollosi, 1980; Webb et al., 1986; Pickering & Johnson, 1987). In addition, exposure of human and animal oocytes to cryoprotectant can induce spindle alterations and chromosomal anomaly (Van Elast et al., 1988; Sathananthan et al., 1988; Pickering, 1990). Based on the morphological analysis of spindle microtubules and chromosome, normal rate of spindle and chromosome configuration was significantly high in 10 min equilibration time group compared with the other groups.

Surprisingly the chromosomal breakdown was severe in the recovered oocytes in all experimental groups. The chromosomal stability was kept well in 10 min equilibration group compared with the other groups. The 10 min equilibration group showed best condition compared with the other groups, but the rate was less than 56%. Therefore, to improve the stability of chromosome, more fine analysis about the effects of physical stress on oocyte during vitrification is needed to define the optimal condition, it is suggested that the optimal equilibration time to get competent oocyte in mouse is 10 min.

The survival rate was higher than those of the previous reports independently to the equilibration times. In addition, the normal rate of spindle and chromosome configuration was improved by changing the equilibration time. However, the stability of DNA was not dramatically improved. Based on these results, it is suggested that the equilibration time is one of the key factors in successful keeping for competence of mature oocyte. Information acquired this study with mouse oocyte may provide insight into intracellular structural events occurring in human oocytes after vitrification and application for cryopreservation of human or other mammals oocytes.

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