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Cryopreservation of caprine ovarian tissue using glycerol and ethylene glycol

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Abstract

Cryopreservation of ovarian tissue may be a potential alternative for the conservation of genetically superior animals, including high milk- and meat-producing goat breeds. However, until now, no information was available concerning the cryopreservation of preantral follicles (PF) enclosed in caprine ovarian tissue. The objective of the present study was to evaluate the structural and ultrastructural characteristics of caprine PF after exposure to and cryopreservation of ovarian tissue in 1.5 and 3 M glycerol (GLY) and ethylene glycol (EG). At the slaughterhouse, each ovarian pair from five adult mixed breed goats was divided into nine fragments and randomly distributed into treatment groups. One fragment was immediately fixed for histological examination and ultrastructural analysis, after slaughter (control). Four of the ovarian fragments were equilibrated at 20 °C for 20 min in 1.8 ml of MEM containing 1.5 or 3 M GLY or EG for a toxicity test and the final four fragments were slowly frozen using these cryoprotectants at the concentrations above. After toxicity testing and freezing/thawing, the ovarian fragments were fixed for histological examination. Histological analysis showed that after toxicity testing and cryopreservation of the ovarian tissue in GLY or EG at both concentrations, the percentage of normal PF was significantly lower than controls. Ultrastructural analysis of PF frozen in 1.5 and 3 M GLY, as well as 3 M EG demonstrated that these follicles remained morphologically normal. In conclusion, we demonstrated cryopreservation of caprine PF in ovarian tissue.

Keywords: Preantral follicles; Cryopreservation; Cryoprotectants; Caprine; Ovarian tissue

1. Introduction

In an attempt to protect biodiversity and to assure the preservation of a species, cryotechnology has been extensively used for male and female gamete conservation; this has allowed the development of germplasm banks of valuable animal stocks and endangered species. Several studies have reported successful cryopreservation of metaphase II oocytes [1], [2], [3] and [4]. However, it is well known that in metaphase II oocytes, the chromosomes are aligned along a delicate spindle of microtubules that are sensitive to cooling and the cryoprotective solutions [5] usually used in cryopreservation protocols.
It has been suggested that preservation of female gametes may be better achieved by storing pieces of ovarian tissue, containing numerous immature small oocytes enclosed in preantral follicles (PF) [6]. These oocytes are less differentiated, possess fewer organelles, lack a zona pellucida and cortical granules, and are less active metabolically. All these characteristics are potentially beneficial to the freezing/thawing process [7], [8] and [9]. Gosden et al. [10] demonstrated that the fertility of ovariectomized sheep can be restored after cryopreservation and transplantation of ovarian fragments containing immature oocytes enclosed in preantral follicles. Although there has been some success in ovarian tissue cryopreservation, several studies [11], [12], [13], [14] and [15] indicated that follicular damage occurs after freezing and thawing. Such damage can occur during any step of the cryopreservation process (exposure of the tissue to the cryoprotectant, freezing or thawing). To improve the efficiency of this procedure, it is necessary to identify and to quantify oocyte losses at different stages of cryopreservation.

Most protocols for ovarian tissue cryopreservation utilize permeable cryoprotectants at a concentration of 1.5 M and do not report the effect of other concentrations of cryoprotectant [15], [16] and [17]. Some authors [12] and [18] affirm that GLY is less permeable than other cryoprotecants such as EG, and consequently, not suitable for the cryopreservation of ovarian tissue. However, none of the cryoprotectants noted have been tested on caprine ovarian tissue. Moreover, most reports are based on histological findings [17], [19], [20], [21], [22], [23] and [24], that are less reliable than transmission electron microscopy (TEM). Therefore, the objective of this study was to evaluate the structural and ultrastructural characteristics of caprine preantral follicles after exposure and cryopreservation of the ovarian tissue in GLY and EG at concentrations of 1.5 and 3 M.

2. Materials and methods

2.1. Source and preparation of ovarian tissue

Caprine ovaries (n=10) from adult mixed breed goats (n=5) were obtained at a local abbatoir. The ovaries were trimmed, and washed in 70% alcohol and then in phosphate buffered saline (PBS). Subsequently, the ovarian pair from each animal was divided into nine fragments of approximately 3-mm thickness (Fig. 1). For fresh controls (Treatment 1: T1), one ovarian fragment was taken randomly and immediately fixed for 24 h in Trump McDowell fixative solution (TMFS) for histological examination. From this fragment, a small piece was taken and fixed in a paraformaldehyde/glutaraldehyde solution (2% paraformaldehyde and 2.5%
glutaraldehye in 0.1 M cacodylate buffer, pH 7.2) for TEM. The other eight fragments were transported to the laboratory within 1 h in PBS at 20 °C and submitted to toxicity testing (n=4) or freezing–thawing (n=4), as described in the following section.

![Fig. 1. Experimental design for toxicity testing and cryopreservation of goat preantral follicles enclosed in ovarian tissue.](image)

### 2.2. Toxicity test

The toxicity test was performed to evaluate the effect of the exposure of ovarian tissue to glycerol (GLY) or ethylene glycol (EG), without freezing, on the percentage of normal follicles. For this test, ovarian fragments (n=4) were exposed for 20 min at 20 °C (equilibration period) in 1.8 ml of MEM with GLY or EG, both at 1.5 or 3 M (T2–T5). After the equilibration period, cryoprotectants were immediately removed from the tissue according to the method described by Candy et al. [12]. The fragments were washed three times for 5 min in minimum essential medium (MEM) supplemented with 10% fetal bovine serum (FBS-MEM+) at room temperature. Finally, the ovarian fragments were fixed in TMFS for histological examination.

### 2.3. Freezing and thawing procedures

For freezing, ovarian fragments (n=4) were placed individually in 2.0 ml vials and equilibrated in 1.8 ml MEM+ with cryoprotectants (GLY or EG) at the concentrations and equilibration period used in the toxicity test (T6–T9). After the equilibration period, the vials containing the ovarian tissue were transferred to a biological programmable freezer (Freeze...
Control, CryoLogic Pty Ltd., Waverley, Australia) at 20 °C. The vials were cooled at 2 °C/min to −7 °C and ice crystal formation (seeding) was induced manually by touching the vials with forceps pre-chilled in liquid nitrogen. The specimens were held at this temperature for 15 min, then cooled at 0.3 °C/min to −30 °C and thereafter at 0.15 °C/min to −33 °C. The vials were plunged directly into liquid nitrogen (−196 °C) and stored for up to 5 days before thawing. The freezing curve used in this study was chosen because most procedures currently used to cryopreserve oocytes [25], [26], [27], [28], [29] and [30] and ovarian tissue [10], [11], [12], [14], [15], [18], [24], [31], [32] and [33] stipulate a cooling rate of 0.3–0.5 °C/min from the seeding temperature (usually −5 to −9 °C) to a lower temperature, usually between −30 and −40 °C.

For thawing, the vials were taken from the liquid nitrogen, warmed rapidly at room temperature for 1 min and immersed in water bath at 37 °C until the ice melted. The cryoprotectant was then removed as described above for toxicity testing and fixed for histological examination and TEM. Each treatment was repeated five times.

2.4. Histological analysis

Ovarian tissue from all treatments was fixed in TMFS for 24 h to evaluate the quality of goat preantral follicles inside the ovarian tissue. After fixation, fragments were dehydrated, diafanized and embedded in paraffin wax. Serial sections (7 μm) of ovarian tissue were cut and every fifth section was mounted on glass slides and stained with periodic acid Schiff (PAS)–hematoxylin. All sections were examined using a light microscope at magnifications 20× and 40×. Thirty preantral follicles in each replicate were evaluated per treatment (T1–T9). Preantral follicles were classified as either an oocyte surrounded by one flattened and/or cuboidal layer or several layers of only cuboidal granulosa cells. To avoid counting a follicle more than once, preantral follicles were counted only in the sections where the oocyte nuclei were seen. Follicular quality was evaluated based on morphological features such as integrity of the oocyte, granulosa cells and basement membrane. Preantral follicles were classified as morphologically normal (preantral follicles containing an intact oocyte and granulosa cells without pyknotic nuclei), degenerated Grade 1 follicles (preantral follicles characterized by pyknosis of the oocyte nucleus) and degenerated Grade 2 (preantral follicles characterized by oocyte nucleus pyknosis plus oocyte shrinkage, with or without detachment of granulosa cells from the basement membrane).
2.5. Ultrastructural analysis

For a better evaluation of follicular morphology, ultrastructural analysis was performed using ovarian fragments from control groups and cryopreservation treatments. Tissue fragments with a maximum dimension of 1 mm3 were fixed in 2% paraformaldehyde and 2% glutaraldehyde in 0.1 M sodium cacodylate buffer (pH 7.2) for 3 h. After fixation and several washes, specimens were post-fixed in 1% osmium tetroxide, 0.8% potassium ferricyanide and 5 mM calcium chloride in 0.1 M sodium cacodylate buffer for 1 h. Subsequently, the samples were dehydrated through a gradient of acetone solutions and the tissues were embedded in Spurr. Semi-thin sections (3 μm) were stained with toluidine blue. The ultra-thin sections (60–70 nm) were cut and stained with uranyl acetate and lead citrate, and examined under a Jeol JEM 100 C transmission electron microscope.

2.6. Statistical analysis

The effect of type (GLY and EG) and concentration (1.5 and 3 M) of cryoprotectant on the percentage of normal and degenerated Grades 1 and 2 preantral follicles were analyzed by Chi-square test, using data from five replicates. Values were considered statistically significant when P<0.05.

3. Results

3.1. Morphological aspects of normal and degenerated Grades 1 and 2 preantral follicles

A total of 1350 preantral follicles were examined in the five replicates. Histological analysis of the ovarian tissue indicated that normal (Fig. 2A) and degenerated Grade 1 (Fig. 2B) and Grade 2 follicles (Fig. 2C) were found in controls as well as after toxicity testing and cryopreservation. Combined for all data, the distribution of normal and degenerated Grades 1 and 2 preantral follicles was 89, 4, and 7% for control, 52, 6 and 42% for toxicity test, and 39, 15 and 46% for cryopreservation, respectively. Of the total degenerated follicles (especially related to the oocyte compartment), 14% had only nuclear pyknosis and 16% had oocyte shrinkage associated with nuclear pyknosis. Considering both compartments (oocyte and granulosa cells), 63% of the follicles had pyknosis in the oocyte nucleus plus oocyte shrinkage.
and detachment of granulosa cells from the basement membrane, 2% detachment of granulosa cells from the basement membrane accompanied by oocyte shrinkage, and 5% detachment of granulosa cells from the basement membrane, accompanied by pyknosis in the oocyte nucleus. The major finding observed by ultrastructural analysis of degenerated follicles was an intense vacuolization in the oocyte cytoplasm and loss of granulosa cell contents.

Fig. 2. Histological sections of goat normal (A) and degenerated Grade 1 (B) and degenerated Grade 2 (C) preantral follicles after cryopreservation (magnification: 400x). O: normal oocyte; NU: nucleus of normal oocyte; GC: granulosa cells; RO: retraced oocyte; PN: pycnotic nucleus; DGC: disorganized granulosa cells.
3.2. Percentage of normal preantral follicles after toxicity testing and cryopreservation

The percentage of normal preantral follicles analyzed by histological analysis in controls as well as after toxicity testing and cryopreservation is shown in Table 1. Exposure to GLY or EG in both concentrations (toxicity testing) reduced (P<0.05) the percentage of normal preantral follicles compared to controls. For both cryoprotectants, the percentage of normal follicles was lower (P<0.05) for hen 3 M versus 1.5 M. When the cryoprotectants were compared to each other at the same concentration, the percentage of normal follicles was higher (P<0.05) in EG than GLY for both concentrations. When both cryoprotectants were compared between the different concentrations, the percentage of normal preantral follicles was higher (P<0.05) in 1.5 M EG compared to 3 M GLY. However, in 3 M EG, the percentage of normal preantral follicles was similar to that observed in 1.5 M GLY (Table 2).

Table 1
Normal preantral follicles (%) in control, after toxicity testing and cryopreservation, as determined by histological evaluation

<table>
<thead>
<tr>
<th>Cryoprotectant</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>GLY 1.5 M</td>
<td>51±a,b,d (76/150)</td>
</tr>
<tr>
<td>GLY 3 M</td>
<td>33±b,d (50/150)</td>
</tr>
<tr>
<td>EG 1.5 M</td>
<td>75±a,d (77/150)</td>
</tr>
<tr>
<td>EG 3 M</td>
<td>51±a,b,d (77/150)</td>
</tr>
</tbody>
</table>

Values within rows (a, b, c) are different (P<0.05).
Values within columns (d, e) are different (P<0.05). *
P<0.05, significantly differs from control

Table 2.
Grades 1 and 2 preantral follicles (%) in controls, after toxicity testing and cryopreservation

<table>
<thead>
<tr>
<th>Control</th>
<th>Toxicity testing</th>
<th>Cryopreservation</th>
</tr>
</thead>
<tbody>
<tr>
<td>GLY 1.5 M</td>
<td>4a (61/150)</td>
<td>13a,b,c (99/150)</td>
</tr>
<tr>
<td>GLY 3 M</td>
<td>5a,b (95/150)</td>
<td>22a,b (119/150)</td>
</tr>
<tr>
<td>EG 1.5 M</td>
<td>4a,b,c (67/150)</td>
<td>36a,b,c (125/150)</td>
</tr>
<tr>
<td>EG 3 M</td>
<td>35a,b,c (85/150)</td>
<td>47a,b,c (117/150)</td>
</tr>
</tbody>
</table>

Different superscripts (a, b, c) differ significantly at the same cryoprotectant and same concentration.
Different superscripts (d, e) differ significantly between Grades 1 and 2 degeneration.
Different superscripts (f, g) differ significantly between toxicity test and cryopreservation. *
P<0.05, significantly differs from control

After freezing and thawing, the percentage of normal follicles was lower (P<0.05) than in fresh ovarian tissue (controls). After cryopreservation, the percentage of normal preantral...
follicles did not differ among treatments, except for 3 M EG, which had a significantly higher percentage of normal follicles than 1.5 M EG.

When comparisons were made between toxicity testing and cryopreservation using the same cryoprotectants and concentrations, in 1.5 M EG the percentage of normal preantral follicles was lower (P<0.05) after freezing and thawing when compared to toxicity testing.

3.3. Distribution of follicular degeneration grade in controls, after toxicity testing and cryopreservation

Fig. 3 shows the distribution of degenerated Grades 1 and 2 preantral follicles in controls and after toxicity testing and cryopreservation. The percentage of degenerated Grade 1 follicles in controls was lower (P<0.05) than in 3 M EG after toxicity testing and 1.5 M GLY and EG after cryopreservation. After toxicity testing and cryopreservation, the percentage of degenerated Grade 2 follicles treatments was higher (P<0.05) than the controls. There was no significant difference between percentages of degenerated Grades 1 and 2 follicles in controls.

Fig. 3. Electron micrograph of a follicle from control group. O: oocyte, GC: granulosa cells, Nu: nucleus, m: mitochondria, er: endoplasmic reticulum, v: vesicles (magnification: 3400×).

Exposure of ovarian tissue to 1.5 M EG (toxicity testing) had a lower percentage (P<0.05) of degenerated Grades 1 and 2 follicles when compared to 3 M. The percentage of Grade 2 degeneration was higher (P<0.05) in GLY than in EG at both concentrations. With regard to the distribution of follicular degeneration grades for each treatment, for both cryoprotectants and concentrations, the percentage of degenerated Grade 2 follicles was higher (P<0.05) than degenerated Grade 1 follicles.

In contrast to observations in the toxicity test, after cryopreservation with 1.5 M EG, a greater percentage (P<0.05) of degenerated Grade 1 follicles was found in 3 M EG, as well as in
1.5 M GLY. Regarding Grade 2 degeneration, when the concentrations were compared to each other in the cryoprotectant, no significant difference was observed. The percentage of Grade 2 follicles was higher (P<0.05) when compared to Grade 1 follicles in all treatments for both cryoprotectants and concentrations, except for 1.5 M EG.

When comparisons were made between toxicity testing and cryopreservation in the same cryoprotectant and concentration, in 1.5 M GLY and EG, the percentage of degenerated Grade 1 follicles in the toxicity test was lower (P<0.05) than after cryopreservation. Similar results were also observed in degenerated Grade 2 follicles in 1.5 and 3 M EG.

3.4. Ultrastructural analysis of the ovarian tissue

Only preantral follicles from controls (Fig. 3) and those frozen in 1.5 and 3 M GLY (Fig. 4A and B), as well as 3 M EG (Fig. 4D) appeared morphologically normal after ultrastructural analysis. These follicles exhibited a large number of mitochondria and some vesicles distributed through the ooplasm. Transmission electron microscopy revealed that follicles from ovarian tissue frozen at 1.5 M EG considered normal at histological analysis (Fig. 4C) were extremely vacuolated. In some cases, the fusion of these vacuoles produced an empty area. Sometimes this alteration was associated with the loss of granulosa cell content and, in some cases, with oocyte nucleus shrinkage (data not shown). In order to verify that the deleterious effect of 1.5 M EG occurred in toxicity testing, ultrastructural analysis was also performed in the ovarian tissue from this treatment. The ultrastructure of normal preantral follicles in the ovarian tissue exposed to 1.5 M EG was similar to controls, indicating that the ultrastructural change occurred during the cryopreservation process.
The present study was the first report of toxicity testing and cryopreservation in GLY and EG of caprine preantral follicles enclosed in ovarian tissue. Although degenerated follicles were observed during exposure to cryoprotectants and after freezing and thawing, a normal morphologic appearance was maintained in 31% (EG: 1.5 M) to 47% (EG: 3 M) of the preantral follicle population after freezing and thawing. Normal follicles obtained after freezing/thawing indicated that the morphology was similar to that observed in vivo. In addition, our preliminary results demonstrated the ability of cryopreserved follicles to survive after in vitro culture.

Toxicity testing revealed that all treatments had a toxic effect on caprine preantral follicles; there was a significant reduction in morphologically normal follicles. Cryoprotectants can chemically interact with important biomolecules in cells or tissues and exert toxic effects
if they are used in high or inadequate concentrations. Besides concentration, temperature and duration of exposure to a certain cryoprotectant can also influence its toxicity. Therefore, we suggest that the temperature (20 °C) and the time (20 min) of exposure used in the present study may not be appropriate for GLY and EG at the tested concentrations, since the percentage of morphologically normal preantral follicles was lower than for controls. The percentage of normal preantral follicles after cryopreservation was also significantly lower than in controls. However, that reduction had already been observed immediately after exposure to the cryoprotectant, indicating that cryopreservation was not the only factor responsible for the loss of follicular viability. Contrary to our findings, Hovatta et al. [14] observed no difference in the percentage of normal and atretic preantral follicles in human ovarian tissue before and after freezing. These results may be due to the differences between species, as well as to the cryopreservation protocol (type and time of exposure to cryoprotectant, slower freezing curve and a multistep process removal of cryoprotectant) used. The successful cryopreservation of oocytes has also been previously reported. Candy et al. [25], working with mouse oocytes, showed that the rate of maturation to metaphase II was not significantly different between frozen and fresh oocytes (83% versus 74%). Additionally, the in vitro fertilization rate was similar for frozen and fresh oocytes (70% versus 81%). In other studies, there was successful cryopreservation of oocytes of several species, with live births reported for mice [36], rabbits [37], and cows [1]. The success obtained in these experiments may be due to the cryopreservation protocols being more appropriate for the developmental stage of the oocytes and the preservation forms used (oocytes without cumulus cells), as well as to intrinsic characteristics of each species.

With regard to the toxicity test, 3 M GLY and EG were more toxic to ovarian tissue than 1.5 M. However, after cryopreservation, 3 M EG was more efficient at maintaining follicle integrity than 1.5 M EG, suggesting that 1.5 M EG provided less protection of caprine ovarian tissue against cryoinjury or thawing. A concentration of 3 M EG is probably more effective than 1.5 M EG at reducing intracellular ice formation (IIF). The cryoprotectants diffuse across cell membranes in exchange for cell water. This displacement of water by cryoprotectants, in addition to freezing point depression, decreases the possibility of IIF and maintains cell volume during freezing, avoiding damage [38]. For caprine ovarian tissue, we speculate that a higher concentration of EG may be necessary. Toxicity testing also demonstrated that 1.5 and 3 M EG were more efficient than GLY (at the same concentrations) for maintaining normal morphology of the preantral follicles. However, after cryopreservation, the efficiency of these two cryoprotectants was similar. The present study demonstrated the importance of the toxicity test; it showed that exposure to cryoprotectants can have deleterious effects on ovarian
tissues that can compromise the cryopreservation procedure. However, it also demonstrated that toxicity testing could not predict the ideal cryoprotectant and concentration for cryopreservation. Toxicity testing verified that 1.5 M EG was better than 3 M EG or 1.5 and 3 M GLY. After cryopreservation, alterations were intensified during the other phases of the cryopreservation process, but only at concentrations of 1.5 M EG; this may be due to the inadequacy of time (20 min), and temperature (20 °C) of exposure to 1.5 M EG, or the combination of those factors with the cooling rate used.

In all treatments, from the toxicity test to after cryopreservation, the percentage of degenerated Grade 2 follicles was greater (P<0.05) than that of controls, demonstrating that the toxic effect of the cryoprotectant can be detrimental for both follicular compartments (oocyte and granulosa cells); this may be due to osmotic damage caused by GLY and EG. During exposure to cryoprotectants, cells initially lose water and shrink to compensate for the high extracellular osmolarity caused by the cryoprotectants [39]. Considering these normal events and the present results, we suggest that the concentrations of cryoprotectants, temperature and time of exposure tested were not optimal for caprine preantral follicles.

When comparisons were performed between the cryoprotectants at both concentrations (1.5 M GLY×3 M EG and 3 M GLY×1.5 M EG), after toxicity testing the percentage of Grade 1 preantral follicles was significantly superior in 3 M GLY and EG when compared, respectively, to 1.5 M EG and GLY. On the contrary, after cryopreservation, 1.5 M GLY and EG were, respectively, more harmful than 3 M EG and GLY, indicating that for ovarian tissue needs a higher concentration of each cryoprotectant for cryopreservation, probably due to the occurrence of IIF. According to Mazur [40], the two main causes of cellular death during freezing and thawing are solution effects and IIF.

Regarding the percentage of Grade 2 degeneration after toxicity testing, the rate was significantly higher for 1.5 and 3 M GLY when compared respectively, to 3 and 1.5 M EG. After cryopreservation, the percentage of degenerated Grade 2 follicles was also significantly greater for 3 M versus 1.5 M EG. On the other hand, the percentage of follicles with degeneration Grade 2 in 3 M EG was similar to that found for ovarian tissue cryopreserved in 1.5 M GLY. Therefore, GLY was more toxic for ovarian tissue than EG. This may have been due to the low permeability of GLY; this could have increased osmotic stress during addition, thawing and removal of the cryoprotectant, since the influx of water is faster than the efflux of GLY from the cells [12] and [41].

The cryopreservation of ovarian tissue in 1.5 M GLY and EG resulted in a significant increase in the percentage of preantral follicles with Grade 1 degeneration when compared to the toxicity test. Therefore, to obtain a lower percentage of degenerated Grade 1 follicles, we
could utilize a higher cryoprotectant concentration, although the percentage of normal preantral follicles, for both 1.5 and 3 M GLY, was similar. Similar results were observed for follicles with Grade 2 degeneration, in the presence of 1.5 and 3 M EG, for both toxicity testing and cryopreservation. These results may be due to ice crystal formation between follicular cells and within follicular cells and the oocyte. Sugimoto et al. [13] demonstrated that the formation of ice in the intercellular space could easily damage the cells and destroy cell–cell communication, which is essential to maintain tissue integrity. Ovarian tissue, like other tissues, is a complex system of organized cells that requires coordinated interactions between the cells and the extracellular matrix to maintain function [42]. Addition of cryoprotectant prior to cooling and its removal after warming involves coupled membrane transport of water and the cryoprotectant; if performed improperly, this can result in deleterious cell volume changes [43]. Some cell types are damaged by the cell shrinkage induced by cryoprotectant addition. However, cell swelling during removal of cryoprotectant is more likely to be deleterious [44].

The follicular degeneration (nuclear pyknosis and oocyte shrinkage) detected in the present study were also described by other authors after cryopreservation of human [14] and non-human primate ovarian tissue [11]. Additionally, the degeneration in the granulosa cell layer (detachment and disorganization) observed in our study has also been detected after vitrification of murine [13] and slow freezing of bovine [15] ovarian tissue. The system used to classify degenerated follicles (Grades 1 and 2) was similar to that described by Paynter et al. [15] for bovine degenerated follicles in cryopreserved ovarian tissue. However, there was an important difference between the classifications. In the present study, alterations in the granulosa cells of the main follicular category used (i.e. primordial follicles) were always accompanied by oocyte shrinkage and/or pyknosis. In contrast, Paynter et al. [15] used primary and secondary bovine follicles. This result may be due to the differences between species, as well as to the developmental stage of the follicles used in the cryopreservation procedure.

Histology did not reveal qualitative differences between different cryoprotectants and concentrations in preantral follicles (soon after cryopreservation), except between 1.5 and 3 M EG. However, TEM revealed that follicles cryopreserved in 1.5 M EG had many signs of ultrastructural abnormality, such as the presence of numerous vacuoles in the oocyte cytoplasm. Hay et al. [45] and Assey et al. [46] demonstrated that cytoplasmic vacuoles are also a characteristic sign of degeneration in both granulosa and cumulus cells during degeneration in vivo. According to Fuku et al. [47], in bovine cryopreserved oocytes, these vacuoles may represent altered mitochondria. The ultrastructural analysis also demonstrated
that the ultrastructural morphology of cryopreserved follicles in 3 M EG was better preserved than that of follicles cryopreserved in 1.5 M EG and GLY at both concentrations. Hovatta et al. [14] reported that histology is relatively uninformative if performed immediately after the thawing; sudden changes in the organelles can appear even several hours after thawing. Therefore, when TEM is not available, it is advisable to maintain the ovarian tissue in an incubator prior to fixation [14]. In addition, using different cryopreservation protocols from that used in the present study, others have demonstrated the inefficiency of GLY in the cryopreservation of female gametes [48] and [49]. Parrot [48] showed that 80% of the oocytes present in mice ovarian tissue were destroyed after exposure to 1.6 M GLY for 1.5 h at room temperature. In nonhuman primates (Rhesus and Cynomolgus monkeys) the deleterious effect of GLY was also demonstrated in cooling of immature oocytes. In these species, the actin filament system was distorted in oocytes exposed to 1.0 and 2.0 M GLY at 23 °C [49]. Previous work with mature caprine oocytes indicated that the permeability of GLY was highly dependent on temperature, and at 20 °C [4] and 4 °C [16], this cryoprotectant has lower permeability than EG. Newton et al. [18] compared the efficiency of four cryoprotectants (DMSO, propanediol, EG and GLY) in the cryopreservation of human ovarian tissue; they showed that EG had better results because it penetrated more quickly in the tissue, due probably to its lower molecular weight [50]. Based on the literature [12] and [16], EG was more effective at protecting cells during cryopreservation than GLY, perhaps due to its low toxicity and efficient permeability [51]. Although some authors [12], [16] and [18] have shown that GLY is inferior to the other cryoprotectants, in our study the percentage of normal preantral follicles after cryopreservation in GLY was similar to EG. It is noteworthy that we used a different slow freezing curve and a removal procedure for GLY similar to the employed by Candy et al. [12]. However, results obtained by Candy et al. [12] using GLY (4.3–28.3 and 16.3–32%: survival of primordial and primary follicles, respectively) were comparable to those for caprine preantral follicles in our experimental conditions. Another report in human ovarian tissue [18] using a freezing protocol similar to Candy et al. [12], i.e. slow freezing curve and fast removal of GLY without sucrose, obtained only 10% viable preantral follicles after freezing and thawing. Similarly to Candy et al. [12] and Newton et al. [18], in the present study we used an initial cooling rate of 2 °C/min to the seeding temperature and a cooling rate of 0.3 °C/min from the seeding temperature (−7 °C in this experiment) to a lower temperature. However, unlike previous studies in which the ovary was cooled to −140 °C [18] or −150 °C [12] before storage in liquid nitrogen, in our study the ovarian tissue was cooled to −33 °C. According to Shaw et al. [52], although cells cooled in this way still contain some water when they are plunged into liquid nitrogen, there is not enough intracellular water to cause damage
(providing that appropriate rapid warming protocols are used). However, the cryopreservation procedure used in this work might be improved by testing different freezing curves, removal procedures of cryoprotectant, and equilibration periods.

In conclusion, we demonstrated that caprine preantral follicles in ovarian tissue can be cryopreserved using GLY and EG. Based on these results, cryopreservation of caprine ovarian tissue can be a feasible alternative for preservation of female genetic material, with the opportunity to subsequently restore ovarian activity via transplant and/or in vitro maturation of preantral follicles enclosed in the ovarian tissue.

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