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Technical note

Quality of frozen-thawed semen in brown bear is not affected by timing of glycerol addition

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Abstract

We have tested several freezing protocols for brown bear semen, modifying the time when glycerol was added (before and after cooling to 5 °C). No differences were found among protocols, indicating a good tolerance of brown bear semen to glycerol. This finding indicates that freezing protocols for brown bear semen could be modified to fit practical solutions which would facilitate preparation of the seminal samples in the field with the addition of glycerol at ambient temperature.

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1. Introduction

Genome Resource Banks (GRB) are a valuable support for wild species conservation, but their application requires the optimization of cooling and freezing protocols. We are currently working on the conservation of Cantabrian brown bear (*Ursus arctos*) in Spain, which is seriously endangered due to habitat loss and population fragmentation, and is a suitable target for the application of GRBs [1].

One of the critical steps in the cryopreservation of a semen sample is the choice of an adequate cryoprotectant, its optimal concentration and the method of adding it during the freezing protocol to minimize the osmotic and toxic effects induced by cryoprotectant on

the cells. Glycerol has been one of the most widely applied cryoprotectant additives for mammalian spermatozoa and it has been used successfully to freeze bear spermatozoa (giant panda [2], Hokkaido brown bears [3] and Japanese black bears [4-6]). Different protocols of glycerol addition have been reported in these studies. Thus, giant panda spermatozoa were cryopreserved adding glycerol at room temperature, immediately before slow cooling [2]. Ishikawa et al [3] reported freezing semen from Hokkaido brown bears using serial dilutions, adding 1/10, 2/10, 3/10 and 4/10 of 14% extender at 4 °C, achieving a final concentration of 4.7%. Okano et al [4-6] cryopreserved semen from Japanese black bears at different final concentrations, diluting extended semen at 4 °C, adding the same volume of extender with twice the final glycerol concentration. However, these results do not show which is the most effective method of addition in the manage-

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ment of bear semen. To our knowledge, there are no previous studies on when or the method to be used for adding glycerol to the extended semen of brown bear.

In other species, many studies have been carried out on the protocol of addition of glycerol, some concluding that glycerol should be added after cooling (4–5 °C), for example in sheep [7], pig [8] and dog [9], whereas others recommend adding it at room temperature, before cooling the samples (stallion [10], red deer [11] and human [12]). Other authors have obtained the same quality regardless of when glycerol was added (stallion [13], red deer [14] and dog: [15]).

Work with endangered population often has to be carried out in the field. This includes collecting and freezing semen, therefore, the protocols are conditioned by this need. Thus, the aim of the present study was to find the most suitable protocol of glycerol addition at a final concentration of 8% for freezing brown bear semen to facilitate the preparation of the semen in the field, keeping the semen quality in these difficult conditions.

We tested three variants: 1) reaching half of the final concentration at ambient temperature, completing to final concentration after cooling to 5 °C; 2) adding all the glycerol at ambient temperature, reaching the final concentration [2]; and 3) adding all the glycerol after cooling to 5 °C [3–6]. Our starting hypothesis was that if the method of glycerol addition does not affect the sperm quality, it could be added at ambient temperature, thus making preparation of the seminal sample in the field easier.

2. Materials and methods

All the chemicals were obtained from Sigma (Madrid, Spain), except Equex STM Paste (Minitüb, Tiefenbach, Germany). Animal manipulations were performed in accordance with Spanish Animal Protection Regulation RD223/1998 (European Union Regulation 86/609).

2.1. Animals and sample collection

We used 17 fresh semen samples, collected from 16 adult brown bears (between 7 and 20 years old) in seventeen electroejaculation sessions, during the breeding season (late April to early July). The animals were housed in a half-freedom regimen in Cabárceno Park (Cantabria, Spain; 43° 21′ N, 3° 50′ W, altitude: 142 metres), and fed with a varied diet.

The animals were immobilized by intramuscular administration of Zolacepan HCl, Tiletamine HCl

(Zoletil1001; Virbac, Carros, France) 7 mg/kg and ketamine (Imalgene 10001; Rhone-Mérieux, Lyon, France) 2 mg/kg applied by teleanaesthesia and were monitored in order to control their general status under anaesthesia. The pubic region and the penis were cleaned, the rectum was emptied of stools and the bladder was emptied by catheterization and electroejaculation was carried out with a PT Electronics electroejaculator (PT Electronics, Boring, OR, USA) using a transrectal probe 320 mm in length and 26 mm in diameter. Electric stimuli were given until ejaculation (10 V and 250 mA, in average) and ejaculates were collected in 15-mL glass tubes at 30–32 °C.

2.2. Experimental design and semen evaluation

Samples obtained by electroejaculation were centrifuged immediately to remove seminal plasma (600×g, 6 min). The pellet was divided in three aliquots depending on the moment of glycerol addition: PRE (before cooling to 5 °C), POST (after cooling to 5 °C) and HALF (half of the final concentration at ambient temperature, completing to final concentration after cooling to 5 °C). First, each aliquot was diluted with the same volume of TTF extender (TES-Tris-Fructose 300 mOsm/kg, pH 7.1, with 20% egg yolk, 2% EDTA and 1% Equex STM paste) at ambient temperature with 16% glycerol (PRE), 8% glycerol (HALF) and 0% glycerol (POST) according to treatment. The tubes with extended sample were put in glass vessels containing 100 mL of water at ambient temperature and transferred to a 5 °C refrigerator. After reaching 5 °C, the same volume of extender was added to the POST and HALF tubes (16% and 8% glycerol, respectively) to reach the final glycerol concentration (8%). Final sperm concentration was achieved by adding the appropriate volume of extender to achieve a final glycerol content of 8%). After packaging into 0.25 mL plastic straws, the sample is immediately cooled for 1 h at 5 °C, samples were frozen in a programmable biofreezer (Kryo 560-16; PLANER plc, Sunbury, UK) at -20 °C/min down to -100 °C, and then transferred to liquid nitrogen containers. The cryopreserved samples remained in liquid nitrogen for a minimum of 1 week. Thawing was performed by plunging the straws in water at 65 °C for 6 s. After thawing, the samples were subjected to 4 h incubation at 37 °C. The semen samples were evaluated before freezing and after thawing (at 0 h and 4 h post-incubation to check for latent damage to the sperm).

Motility parameters were evaluated using a com-

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