

running head: Estesó et al, Sperm freezability in stags

Functional significance of the sperm head morphometric size and shape for determining freezability in Iberian red deer (*Cervus elaphus hispanicus*) epididymal sperm samples

MILAGROS C. ESTESÓ,<sup>\*†</sup> ANA J. SOLER,<sup>\*</sup> MARÍA R. FERNÁNDEZ-SANTOS,<sup>\*</sup> ARMANDO A. QUINTERO-MORENO,<sup>‡</sup> AND JOSE J. GARDE<sup>\*†</sup>

From the \* Biology of Reproduction Group, Department of Game Resources (IDR), Castilla-La Mancha University (UCLM), Albacete, Spain; † National Wildlife Research Institute (IREC), UCLM-CSIC-JCCM, Albacete, Spain; and ‡ Animal Production Research Unity, Faculty of Veterinary Medicine, Zulia University, Maracaibo, Venezuela.

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Correspondence to: Dr. José Julián Garde, IDR, Sección de Recursos Cinegéticos y Ganaderos (IDR). Campus Universitario, 02071. Albacete, Spain. (e-mail: Julian.Garde@uclm.es).

## Abstract

In the present study, computer-automated sperm-head morphometry of epididymal samples was used to determine if sperm-head area and shape are useful measurements for separating “good” and “bad” Iberian red deer freezers. A microscope slide was prepared from single diluted sperm fresh samples collected from 38 mature stags. Slides were air dried and stained with Hemacolor. The sperm-head area and shape (length/width) for a minimum of 145 sperm heads were determined for each male by means of the Sperm-Class Analyser (SCA). The remainder of each sample was frozen. After thawing, sperm cryosurvival was *in vitro* judged by microscopic assessments of individual sperm motility and of plasma membrane and acrosome integrities. All sperm parameters evaluated at thawing, were placed in a statistical database and a multivariate cluster analysis performed. Mean sperm parameters of the two clusters generated (“bad” and “good” freezers) were compared by ANOVA. Our results show that sperm quality at thawing for all sperm parameters evaluated was significantly higher ( $P < 0.01$ ) for “good” freezers than for the “bad” ones (Sperm motility index:  $67.4 \pm 2.0$  vs.  $57.1 \pm 2.8$ ; NAR:  $67.1 \pm 2.5$  vs.  $54.5 \pm 3.5$ ; Viability:  $68.8 \pm 2.0$  vs.  $60.1 \pm 2.8$ ; HOST:  $71.3 \pm 2.2$  vs.  $63.1 \pm 3.1$ ). Additionally, differences ( $P < 0.01$ ) in epididymal sperm-head area and shape were found between “good” and “bad” freezers before freezing, with the smallest overall sperm head dimensions found in the “good” freezers group (Area:  $32.04 \mu\text{m}^2$  vs.  $34.42 \mu\text{m}^2$ ). Thus, the lower the sperm head area in the fresh samples, the greater the sperm cryoresistance. Our results show that the 2 groups of males also differ in sperm head shape in fresh samples (Good: 1.96, vs. Poor: 1.72;  $P < 0.01$ ). It is possible that sperm head area and shape influence total sperm volume, thus causing differences in heat exchange as well as in movements of water, ions and cryoprotectants and, in turn, on sperm freezability.

**Keywords:** Red deer, Cryopreservation, Morphology, Freezability, Postmortem recovery

## **Introduction**

The interest in preserving germplasm of wild deer species has resulted in rather recent attention to the possible recovery, evaluation and cryopreservation of sperm from the epididymides of dead animals (Zomborszky et al, 1999; Comizzoli et al, 2001a,b; Hishinuma et al, 2003; Soler and Garde, 2003; Soler et al, 2003, 2005). However, little information is published on freezing and thawing methods for epididymal spermatozoa of this species, although there have been many studies of freezing of ejaculated semen from stags (Asher et al, 2000). Most procedures used to cryopreserve epididymal spermatozoa from red deer have been modified from those developed for ejaculated semen. This approach would not seem to be very appropriate since it is well known that there are important differences in the physiological characteristics of epididymal vs. ejaculated spermatozoa, specially in their membranes properties, that affect the sperm cell survival after freezing.

Variations between individuals in the extent to which their spermatozoa are damaged by freeze-thawing has been widely reported in several species. These differences in the resistance to thawing of the spermatozoa of different individuals have been observed for spermatozoa of domestic (see Curry, 2000) and wild (see Leibo and Songsasen, 2002) species. Within this context, semen donors have routinely been categorized as “good” or “bad freezers”. The mechanisms underlying differences in cryosensitivity between different individuals have yet to be elucidated, but there is some evidence for physiological differences between spermatozoa from individuals of the same species. Recently, it has been demonstrated that these consistent inter-individual variations in sperm freezability are genetically determined (Thurston et al, 2002).

Appropriate, sensitive, and rapid methods of assessment are necessary for adequate evaluation of sperm function. Conventional sperm parameters are not sufficient to identify animals known as “poor freezers”, whose sperm quality is greatly impaired by cryopreservation. The routine evaluation of semen, including normal sperm morphology assessment, has long been employed to evaluate the effects of freezing-thawing procedures on sperm cryosurvival. Poor semen morphology is an important indicator of decreased fertility in men (Kruger et al, 1993); stallions (Jasko et al, 1990), bulls (Sekoni and Gustafsson, 1987), and goats (Chandler et al, 1988). Although normal sperm morphology may be an indicator of the fertility potential of a given male, correlations have been based on subjectively performed analyses. However, large variations between technicians and laboratories in the subjective evaluation of semen characteristics are known to exist (Saacke, 1982; Baker and Clarke, 1987), making accurate interpretation of the resulting data difficult. The need for accurate objective assessment of sperm morphology has led to the development of computer-assisted sperm head morphometry analysis, ASMA, (Katz et al, 1986; Davis et al, 1992; Pérez-Sánchez et al, 1994). The precision of the ASMA system has been utilized to detect morphometric differences in sperm head dimensions of fertile and subfertile males (Casey et al, 1997) as well as subtle changes in human sperm head morphometry due to toxicant exposure when no morphological differences were detected by manual assessment (Davis et al, 1993).

Previous studies utilizing ASMA have also demonstrated that cryopreservation affects head morphometry of bull (Gravance et al, 1998), human (Thompson et al, 1994), stallion (Arruda et al, 2002), dog (Rijsselaere et al, 2004) and red deer (Esteso et al, 2006) spermatozoa. Recent evidences suggest that in boar semen, the sperm head dimensions of individual sperm

samples, may be an indicator of sperm cryosurvival (Thurston et al, 2001; Peña et al, 2005), while no data exist regarding sperm morphometry and freezability in deer species.

Thus, adopting methods previously utilized in other species (Sancho et al, 1998; Buendía et al, 2002) and in red deer by us (Esteso et al, 2003; 2006), our objective was to determine if sperm-head area and shape are useful measurements for predicting freezability in Iberian red deer epididymal sperm samples. To achieve this goal, firstly frozen-thawed spermatozoa from a single sample collected from 38 stags were evaluated for sperm motility (SMI), and for acrosome (NAR) and membrane integrities (HOST and viability). All data generated were used for a multivariate cluster analysis which objectively classified all sperm samples (stags) within a data set in to one of two groups, categorized as “good” or “bad” according with their freezability. In a second place, we retrospectively compared the routine sperm parameters and the sperm head size and shape in freshly epididymal sperm samples between the two defined groups.

## **Methods**

With the exception of DPX (Fluka, Madrid, Spain) all other chemicals were of reagent grade and were purchased from Sigma or Merck (both of Madrid, Spain).

### *Preparation of testes and collection of epididymal spermatozoa*

For this study, we used spermatozoa recovered from the epididymides of 38 mature stags (age > 4 years, weight > 140 kg) that were legally culled and hunted in their natural habitat during the rutting season (September-November). The hunting of stags was performed in accordance with the

harvest plan of each game reserve. The harvest plans were made following Spanish Harvest Regulation, Law 2/93 of Castilla-La Mancha, which conforms to European Union Regulation. Immediately upon removal, the testes with attached epididymides were placed into plastic bags and transported to the laboratory at room temperature (approximately 20°C) within 2 h after being removed. Samples were processed as soon as they arrived at the laboratory. For collection of epididymal spermatozoa, testes and epididymides were removed from the scrotal sac. Cauda epididymides, including about 5-10 cm of the proximal ductus deferens, were separated and transferred to 35 mm plastic dishes.

Spermatozoa were collected from the distal portion of the epididymis by repeated longitudinal and transverse cuts with a surgical scalpel and collecting the oozing sperm mass and placing it in 0.5 ml of a Salamon's modified solution (Fernandez-Santos et al, 2005). In particular, the base solution (Fraction A) containing Tris (2.70%, w/v), fructose (1%), citric acid (1.4%), glycerol (0%, v/v) and egg yolk (20%, v/v), with pH = 6.8 and osmolality of 300 mOsm. Epididymal contents from both testicles of an individual male were pooled for processing, because our previous observations showed that no differences appeared to exist in cell quality between testes belonging to the same individual (Garde et al, 1998).

#### *Semen processing and cryopreservation*

Then, the sperm mass was again diluted at room temperature to a final sperm concentration of  $\sim 400 \times 10^6$  sperm/ml with the Fraction A of the extender. Sperm dilution was performed in a two-step procedure at room temperature, first adding base extender up two times the final desired sperm concentration and then a second extender (Fraction B) at a 1:1 ratio to achieve a final

concentration  $\sim 200 \times 10^6$  spermatozoa/ml. Fraction B differed from the base diluent in the replacement of water (12%, v/v) with the same volume of glycerol (final concentration = 6%, v/v). This two-step procedure was employed for sperm dilution to obtain the same final concentration of glycerol for each stag. At this point, sub-samples were taken for sperm head morphometric dimensions evaluation. Then, the sperm diluted was placed in a 15-ml centrifuge tube (Iwaki, Japan) and was slowly cooled to 5°C. For it, the tubes were placed in a beaker with water (75 ml at room temperature) and transferred to a refrigerator at 5°C. Cooling down to 5°C lasted for about 1 h and then extended samples were held for equilibration at that temperature for 2 h more. After the equilibration of the diluted sperm samples, the extended sperm was loaded into 0.25 ml plastic straws. Immediately, they were frozen at 4 cm in nitrogen vapor (-120 °C) for 10 min before being immersed into liquid nitrogen (-196 °C) for storage. For freezing, straws were placed horizontally on a metal rack that was positioned into a freezing close container (Cryo Diffusion CD-45). The frozen straws remained for a minimum period of one month in liquid nitrogen before thawing was carried out.

Frozen semen was thawed in a water bath (37°C) for 20 s and the content of the straws poured into a glass tube. Samples were evaluated for motility, viability, and acrosome and membrane integrities after 5 min of incubation at 37°C, using the methods described below. At thawing, sub-samples were also taken for sperm head morphometric dimensions evaluation.

### *Semen evaluation*

Sperm concentration and subjective scores of motility were assessed shortly after collection. Sperm concentrations of the original suspensions were determined using a hemacytometer.

Percentage of individual motile sperm (motility) was noted and quality of motility was assessed using a scale of 0, lowest, to 5, highest. A Sperm Motility Index (SMI) was calculated = [% individual motility + (quality of motility x 20)] x 0.5. The sperm suspension was also used to assess acrosome integrity and viability. Acrosomal integrity was evaluated after a 1:1 dilution in 2% glutaraldehyde in 0.165 M cacodylate/HCl buffer (pH 7.3). The percentage of spermatozoa with intact acrosomes (% NAR) was assessed by phase-contrast microscopy.

In addition, samples were taken to assess the membrane integrity by means of the hypo-osmotic swelling (HOS) test. Plasma membrane functionality was assessed using an HOS test as described by Garde et al (1998). The osmotic swelling technique consisted of mixing 0.01 ml of diluted sperm samples with 0.1 ml of hypo-osmotic solution (100 mOsmol/kg) and incubating the mixture at room temperature for 30 min. The samples were then fixed in 2% glutaraldehyde buffered solution and evaluated by phase-contrast microscopy at X400. The sperm membrane was considered functional in cases where the sperm tail was coiled and the result was expressed as HOS (%).

Membrane integrity (viability) was also evaluated by using a nigrosin-eosin stain (NE). The NE stain was prepared according to the method by Soler et al (2005). Sperm (5 $\mu$ l) was mixed with the NE stain (10  $\mu$ l) at 37°C, incubated for 30 s, smeared and dried on a warm plate at 37°C. The samples were evaluated using bright field microscopy at X400. Live spermatozoa remained unstained, while dead cells were dull pink. The percentage of live spermatozoa was expressed as viability (%).

For sperm assessments, 100-200 spermatozoa were counted in each preparation. Additionally, slides of extended sperm samples were prepared from each sample for sperm head morphometric characterization.

For each sperm parameter evaluated, a ratio was calculated to assess the cryoprotective abilities of the different individual samples. They were calculated as follows:

$$\text{Cryoresistance ratio} = \left( \frac{\text{Value after thawing}}{\text{value in fresh sperm}} \right) \times 100$$

#### *Morphometric analysis of sperm heads*

Microscopic slides were prepared from each fresh (upon dilution) sample by placing 5  $\mu$ l of the sperm samples on the clear end of a frosted slide and dragging the drop across the slide. Semen smears were air dried and stained using a Hemacolor (Merck) procedure, originally described for staining of ram (Sancho et al, 1998) and alpaca (Buendía et al, 2002) sperm heads; and recently adapted by our group to red deer spermatozoa (Esteso et al, 2003). Stained sperm samples were permanently mounted to the slide with a coverslip and dibutyl phthalate xylene (DPX).

Stained slides were used to perform ASMA using the morphometry module of a commercially available system (Sperm-Class Analyzer, SCA, version 99 CASMA system. Microptic, Barcelona, Spain). The machine was equipped with a Nikon (Labophot-2, Tokyo, Japan) microscope with a x 60 bright-field objective and a Sony video camera (CCD AVC-D7CE, Sony Corporation, Tokyo, Japan) connected to a Pentium 950 MHz processor. The illumination source was centred and the intensity of the bulb and the gain and offset of the camera

were standardized for all samples. The configuration of the computer system included a PIP-1024 B video digitiser board (Matrox Electronic Systems Ltd., Quebec, Canada), the sperm image analysis software and a high-resolution assistant monitor Sony Triniton PVM-1443MD (Sony Corporation, Tokyo, Japan). The array size of the video frame recorder was 512 x 512 x 8 bits, digitised images were made up of 262,144 pixels (picture elements) and 256 grey levels. Resolution of images was 0.15 and 0.11  $\mu\text{m}$  per pixel in the horizontal and vertical axes, respectively.

The morphometric dimensions for area (A), and shape factor (length / width) were acquired for 150-160 images. Acquiring 150-160 images assures that a minimum of 145 properly measured sperm heads are analysed after improperly measured sperm heads are deleted from the analysis. The measurements of each individual sperm head from each stag and sperm treatment were saved in an Excel (Microsoft Corporation, Redmond, Washington, USA)-compatible database by the software for further analysis.

### *Statistical analysis*

Statistical analyses were performed using SPSS for Windows, version 11.5 (SPSS Inc, Chicago I11, USA). The effects of cryopreservation on sperm motility, viability, and acrosome and membrane integrities were compared across stags by General Linear Models analysis of variance (GLM-ANOVA) using a split plot design. Stags served as the main plot and sperm cryopreservation step (pre-freezing or post-thawing) served as the subplot. Group differences were compared by Fisher's Least Significant Differences test. Effects were considered significant at  $P < 0.05$ .

Data of post-thaw sperm motility, sperm viability and acrosome and sperm membrane integrities obtained from each of the 38 epididymal sperm samples were used to build a single data set. The data set was subjected to a multivariate analysis as described Estes et al. (2003) for head sperm morphometric patterns. This procedure uses the cluster analysis to classify the sperm samples, using all measured sperm variables within the data set, on a small number of groups. Two groups were finally obtained from the non- hierarchical clustering (K-means clustering) of the 38 sperm samples (e.g. stags) tested. The statistics of each sperm variables of the two groups of males were calculated, and compared using General Linear Models analysis of variance (GLM-ANOVA). Percentage data of sperm fresh parameters and sperm head area and shape in fresh samples from sperm samples clustered following the multivariate analysis were also compared using ANOVA. Data that did not follow a normal distribution were transformed. A probability of  $P < 0.05$  was considered to be statistically significant.

## **Results**

### *Effects of cryopreservation*

In this work, epididymal spermatozoa from 38 stags were frozen and thawed. Because we used epididymal spermatozoa for this experiment, it was only possible to make single observations for each stag. After freezing and thawing, a decrease ( $P < 0.0001$ ) in all routine sperm parameters was observed (Table 1). Thus, sperm motility index decreased from  $83.16 \pm 1.7\%$  in extended fresh samples to  $63.91 \pm 1.7\%$  in thawed samples. The results found for the others seminal parameters evaluated are similar if not identical (Table 1). Our results also showed differences among males for most of seminal parameters evaluated immediately after thawing (Figures 1 and 2).

### *Male subpopulations analysis*

After the cluster analysis of frozen-thawed sperm quality, two groups of sperm samples (stags) were clearly identified. Those samples with best frozen-thawed sperm characteristics were identified as “good” whereas the others represented samples showing considerably reduced frozen-thawed sperm characteristics and were considered as “bad” following a standard freezing protocols. Twenty-five samples were identified as “good” and 13 as “bad”. Summary statistic for these two groups are shown in Table 2. Note the significant differences ( $P < 0.01$ ) for all sperm parameter evaluated.

### *Sperm quality before freezing*

There was no significant variation between groups before freezing for all the sperm parameter evaluated (Table 3). Sperm samples from the two groups consisted of spermatozoa with high quality and would be expected to maintain this quality after cryopreservation.

### *Sperm cryoresistance*

As a consequence that differences in fresh sperm quality between stags exist, we also determined the cryoresistance ratios for all the sperm parameters evaluated (see Materials and Methods). The use of rates, rather than absolute values, allows for a direct comparison between males that differ

in fresh semen parameters (Table 4). Note the significant differences present between groups ( $P < 0.05$ - $0.0001$ ) for all the cryosurvival ratios. These results demonstrate that the resistance to the cryopreservation was different between “good” and “bad” freezers. Good freezers had the highest ratios (range: 70%-83%) in all the parameters evaluated (sperm motility index, % endosmosis and % intact acrosomes) with results that, in all cases, were significantly different from those seen in the “bad” group (range: 58%-70%).

#### Sperm head size and shape in fresh samples

Data of sperm head area and shape from stags classified as “good” and “bad” freezers, evaluating 3,406 and 1,821 properly digitised spermatozoa respectively, revealed that the frozen-thawed sperm characteristics (base for the discrimination into two groups) significantly were affected by the sperm head size and shape (Table 5). There were no differences in the percentage of properly analysed sperm heads between “good” and “bad” freezers (data not shown). Our results showed that the smallest overall sperm head dimensions in fresh samples were found in the “good” freezers group (Area:  $32.04 \mu\text{m}^2$  vs.  $34.42 \mu\text{m}^2$ ). Therefore, the lower the sperm head area in the fresh samples, the greater the sperm cryoresistance. Our results also show that the 2 groups of males differ in sperm head shape (Good: 1.96, vs. Bad: 1.72;  $P < 0.01$ ). Thus, the sperm heads in the fresh samples from the “good” freezers were more elongated and narrow than those from the “bad” group. In this sense, the sperm head length in the fresh samples from “good” freezers was approximately two times higher than the width. Figure 3 shows the sperm head area for each stag categorized as “good” or “bad” freezers.

## **Discussion**

Our previous studies utilizing ASMA have reported the normal sperm head dimensions for fresh and thawed red deer epididymal spermatozoa (Esteso et al, 2003; 2006). In the present study, the ASMA protocol used was useful to detect differences in sperm head area and shape between “good” and “bad” freezers before freezing. Thus, the lower the sperm head area in the fresh samples, the greater the sperm cryoresistance. Differences in the resistance to thawing of the spermatozoa of different individuals have been observed for spermatozoa of other domestic (see Curry, 2000) and wild (see Leibo and Songsasen, 2002) species. Within this context, semen donors have routinely been categorized as “good” or “bad freezers”. Although similar experiences have been reported for several species, no explanations for these differences have been substantiated. The mechanisms underlying differences in cryosensitivity between different individuals have yet to be elucidated, but there is some evidence for physiological differences between spermatozoa from individuals of the same species (see Leibo and Bradley, 1999).

On the other hand, authors have been very interested in determining the relationship between single fresh semen quality characteristics such as motility, viability, morphology or acrosome status and the freezability. In any case, single fresh semen quality could show positive relationship with freezability, but the relationship was either of low significance or measurable in some males but not in others. Inclusion of several sperm variables, either measured by functional methods, or by the combination of significant outcomes on a multivariate regression analysis has been regarded as more discriminative, and in some cases even of predictive value. For more details see Rodriguez-Martinez (2003). Bearing this in mind, the present work aimed to group a series of post-thaw sperm variables defining the survival status of red deer spermatozoa subjected

to a conventional freezing protocol and, by using a multivariate cluster analysis as described by Estes et al (2003) and used already by others authors to classify subpopulations of spermatozoa (Quintero-Moreno et al, 2003), defined two groups of stags as “good” and “bad” according with their freezability.

In this study, out of the 38 males, 25 were identified as “good” and 13 as “bad”. To explore the differences between the two groups, conventional ANOVA analysis was undertaken and significant differences were found for all parameters. The differences between stag groups were found for all quality parameters of thawed spermatozoa as well as the degree of declined cellular integrity by cryopreservation. However, there were no significant differences in routine semen quality between groups before freezing. On the other hand, differences in epididymal sperm-head area and shape were found between “good” and “bad” freezers before freezing, with the smallest overall sperm head dimensions found in the “good” freezers group. Besides, the sperm heads in the fresh samples from the “good” freezers were more elongated and narrow than those from the “bad” group. In this sense, the sperm head length in the fresh samples from “good” freezers was approximately two times higher than the width. It is noteworthy that when comparisons are made among species for their ability to sustain cold shock, clear sperm differences are evident; the spermatozoa of those species less sensitive to cryopreservation are smaller (Garde et al, 2003). Obviously, many others factors are involved in cryoresistance, but we hypothesized that sperm head area and shape cause differences in heat exchange as well as in movements of water and ions. It is, therefore, plausible to think that spermatozoa may vary in their physical properties depending on their area and shape and that these variations are, at least partially, responsible of the inter-individual resistance to the cryopreservation process.

This relationship between sperm head dimensions in fresh ejaculates and sperm freezability has been previously supported for boar spermatozoa (Thurston et al, 2001; Peña et al, 2005). The study of Thurston et al, (2001) reported that the percentage of spermatozoa in the fresh ejaculates with slightly tapering heads was positively correlated with the sperm quality after thawing. Thurston et al (2001) support the hypothesis that the inter-individual variations in sperm morphology are genetically determined and therefore, these differences in sperm morphology in fresh samples, that affect sperm freezability, are indicative of the genetic variations responsible of the relative ability of spermatozoa to withstand freezing procedures. Here, we present the hypothesis that these variations in sperm morphology, in addition to being indirect estimation of that sperm freezability is genetically determined (Thurston et al, 2002), can be also influence *per se* biophysical characteristics of the spermatozoa that are essential for successful cryopreservation. Taken together, our results revealed that fresh spermatozoa from the “good” freezers had small and elongated (higher shape) sperm heads, whereas spermatozoa from “bad” freezers had large and wide (lower shape) sperm heads. However, there was no significant variation between stag groups before freezing for the routine semen parameters evaluated. Therefore, these two groups of males only differ in the dimensions of their fresh sperm heads. So, it might be assumed that the sperm head size and shape in spermatozoa of fresh samples were good indicators of freezability. However, these results must be carefully interpreted since there were sperm samples classified as “good” showing large sperm heads ( $>34 \mu\text{m}^2$ ) and sperm samples categorized as “bad” with small sperm heads in the fresh samples (Figure 3). It seems reasonable then, based on our results, that sperm head area and shape of fresh spermatozoa discriminated between two clear cut populations whose sperm freezability is different. However, such relationship, albeit existing is not enough to accurately estimate, and less predict, the sperm freezability of individual samples. More studies are needed in order to develop procedures for prospectively selecting sperm samples for cryopreservation.

The observation that sperm head size and shape are highly indicative of stags sperm survival after cryopreservation is important for two reasons. First, it is possible that sperm head area and shape influence total sperm volume, thus causing differences in heat exchange as well as in movements of water, ions and cryoprotectants and, in turn, on sperm freezability (Curry, 2000). Second, it suggests that sperm survival from individuals considered as “bad freezers” (with large and wide sperm heads) can be optimised by modifying either the cryoprotectants concentration or the cooling rates. Thus, previous works have reported that sperm characteristics as surface area and volume, have important implications for determining optimum cooling and warming procedures for sperm cryopreservation (Curry et al, 1996). In this sense, improvements in boar sperm cryosurvival have been reported when spermatozoa were frozen at faster rates (Fiser and Fairfull, 1990). When cell suspensions are frozen, they are cooled at finite rates often referred as slow or fast. A rate that is slow for one cell type may be rapid for a second type. The optimum cooling rate is the one at which maximum survival is observed. Thus, the optimum cooling rates for freezing human, boar and ram spermatozoa are different (see Leibo and Bradley, 1999). Besides, sperm area and volume have important implications for determining optimum cooling procedures for cryopreservation (Curry et al, 1996). These sperm characteristics influence the rate at which the cell can lose water; and the rate at which a cell can lose water is a principal determinant of its optimum cooling rate. For reasons briefly described above, efforts to improve sperm freezing protocols have increased substantially during the past years. In this sense, we propose that sperm samples from the different stags require different cooling rates for optimal cryosurvival. Thus, the results of the present study suggest that the freezability of the spermatozoa from individuals considered as “bad freezers” (with large and wide sperm heads) could be increased in the future using different cooling rates than those used in this work

(20°C/min). We based this hypothesis in the fact that cellular area and volume have important implications for determining optimum cooling rates for cryopreservation (Curry et al, 1996; Leibo and Bradley, 1999).

In summary, the results of the present study showed that although a multivariate pattern analysed based on frozen-thawed red deer sperm quality attributes was capable to separate “good” and “bad” freezers and further that these populations could relate to sperm head area and shape in fresh samples; there were sperm samples outside this pattern. Such situation confirms the need for caution when aiming at estimating the potential freezability of a given sperm sample by solely assessing phenotypic variables of the spermatozoa. Future work will utilize ASMA to identify sperm morphometric subpopulations in epididymal fresh sperm samples and their possible relationships with freezability.

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## Legends to figures

Figure 1. Sperm motility index (SMI) of thawed red deer spermatozoa collected post-mortem from epididymides of different stags.

Figure 2. Individual characteristics of thawed red deer spermatozoa collected post-mortem from epididymides of different stags: **(A)** percentage of sperm cells with intact acrosomes (NAR), and **(B)** percentage of spermatozoa with intact plasma membrane (HOS).

Figure 3. Sperm head area ( $\mu\text{m}^2$ ) for individual “good” (●) and “bad” freezers (○).

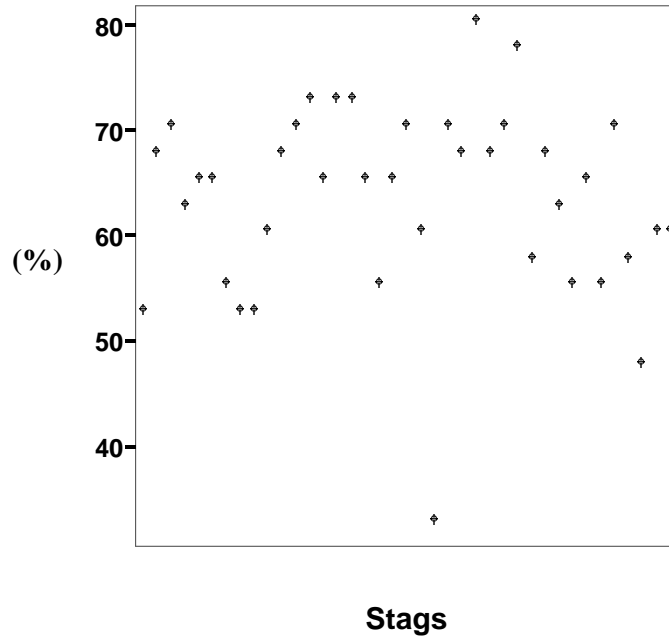


Figure 1. Estes et al

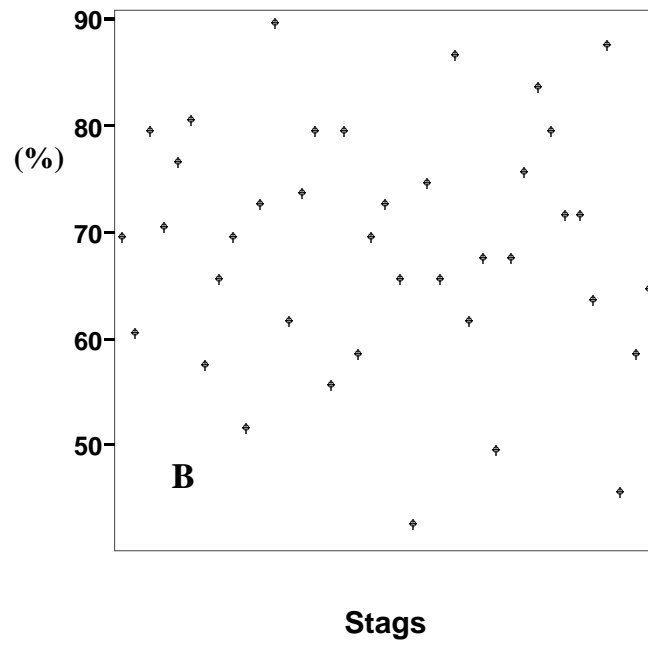
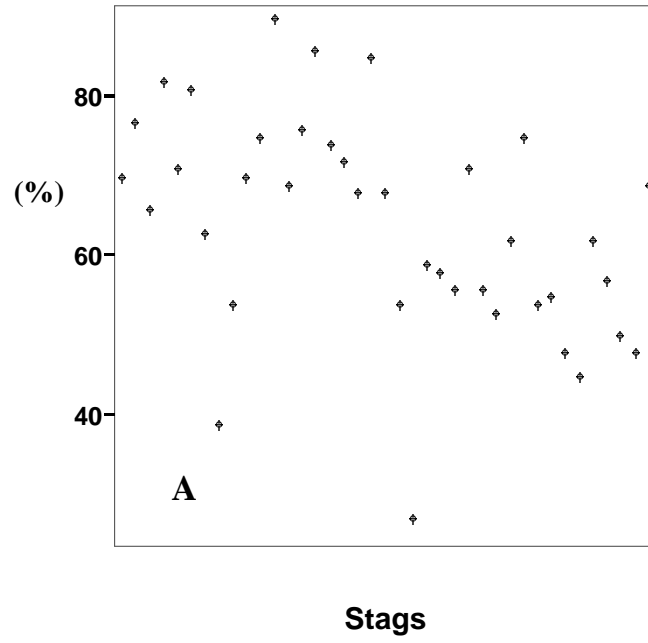


Figure 2. Estes et al

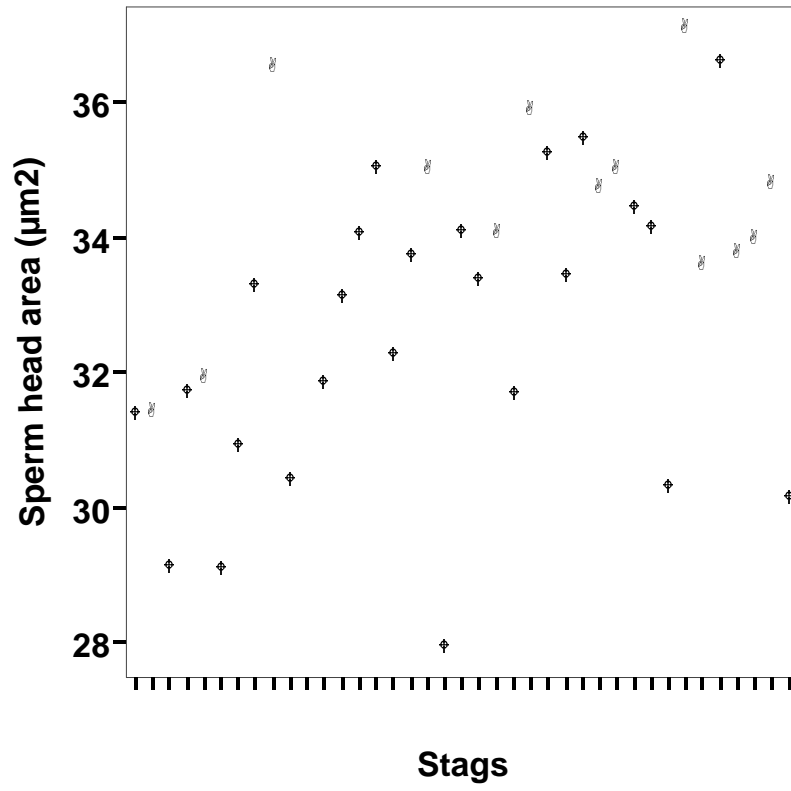


Figure 3. Estes et al

Table 1. Effects of cryopreservation on characteristics of red deer epididymal spermatozoa\*.

Sample	Assessed Parameters †			
	‡SMI (%)	‡NAR (%)	‡HOS (%)	‡Viability (%)
FRESH	83.16±1.7 <sup>a</sup>	94.59±1.6 <sup>a</sup>	89.61±1.6 <sup>a</sup>	89.43±1.4 <sup>a</sup>
CRYO	63.91±1.7 <sup>b</sup>	62.97±1.6 <sup>b</sup>	68.69±1.6 <sup>b</sup>	66.15±1.4 <sup>b</sup>

FRESH, unfrozen sperm sample; CRYO, cryopreserved sperm sample

\* Values are least-squares means ± SEM.

† Values with different superscripts in the same column were significantly different ( $P < 0.0001$ ).

‡ See Material and Methods for details of spermatozoa assessment.

Table 2. Descriptors of frozen-thawed sperm characteristics in the two groups (“good” vs. “bad”) of single stag sperm samples defined after multivariate cluster analysis\*.

Stag group	Assessed Parameters †				
	No. stags	‡SMI (%)	‡NAR (%)	‡HOS (%)	‡Viability (%)
GOOD	25	67.40±2.0 <sup>a</sup>	67.08±2.5 <sup>a</sup>	71.28±2.2 <sup>a</sup>	68.80±2.0 <sup>a</sup>
BAD	13	57.11±2.8 <sup>b</sup>	54.54±3.5 <sup>b</sup>	63.15±3.1 <sup>b</sup>	60.15±2.8 <sup>b</sup>

\* Values are least-squares means ± SEM.

† Values with different superscripts in the same column were significantly different (P < 0.01).

‡ See Material and Methods for details of spermatozoa assessment.

Table 3. Descriptors of fresh sperm characteristics in the two groups (“good” vs. “bad”) of single stag sperm samples defined after multivariate cluster analysis\*.

Stag group	Assessed Parameters †				
	No. stags	‡SMI (%)	‡NAR (%)	‡HOS (%)	‡Viability (%)
GOOD	25	81.40±2.0 <sup>a</sup>	95.52±1.6 <sup>a</sup>	88.88±1.7 <sup>a</sup>	90.24±1.7 <sup>a</sup>
BAD	13	86.23±2.7 <sup>a</sup>	93.00±1.9 <sup>a</sup>	90.61±2.4 <sup>a</sup>	87.61±1.2 <sup>a</sup>

\* Values are least-squares means ± SEM.

† Values with different superscripts in the same column were significantly different (P < 0.05).

‡ See Material and Methods for details of spermatozoa assessment.

Table 4. Sperm cryosurvival in the two groups (“good” vs. “bad”) of single stag sperm samples defined after multivariate cluster analysis\*.

Stag group	Assessed Parameters †				
	No. stags	‡SMI (%)	‡NAR (%)	‡HOS (%)	‡Viability (%)
GOOD	25	83.58±2.3 <sup>a</sup>	70.17±2.5 <sup>c</sup>	80.69±2.5 <sup>c</sup>	76.42±2.3 <sup>e</sup>
BAD	13	66.00±3.2 <sup>b</sup>	58.55±3.5 <sup>d</sup>	70.04±3.5 <sup>d</sup>	68.41±3.2 <sup>f</sup>

\* Values are least-squares means ± SEM.

† Values with different superscripts in the same column were significantly different: <sup>a,b</sup>(P < 0.0001); <sup>c,d</sup>(P < 0.01); <sup>e,f</sup>(P < 0.05).

‡ See Material and Methods for details of cryoresistance rates determination.

Table 5. Sperm head area and shape factor for fresh sperm samples in the two groups (“good” vs. “bad”) of single stags defined after multivariate cluster analysis\*†.

Stag group	No. stags	No. Spermatozoa	‡ Area ( $\mu\text{m}^2$ )	‡ Shape Factor (length / width)
GOOD	25	3,406	32.04 (9.8) <sup>a</sup>	1.96 (8.65) <sup>a</sup>
BAD	13	1,821	34.42 (9.2) <sup>b</sup>	1.72 (7.72) <sup>b</sup>

\* Values are least-squares means. Coefficients of variation (% CV) between stags are shown in parentheses.

† Values with different superscripts in the same column were significantly different ( $P < 0.01$ ).

‡ See Material and Methods for details of spermatozoa assessment.