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Boar sperm cryosurvival is better after exposure to seminal plasma from selected fractions than to those from entire ejaculate

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**Abstract**

Boar bulk ejaculates are now being collected instead of usual sperm-rich fractions (SRF) for artificial insemination purpose. The present study evaluated the influence of holding boar sperm samples before freezing surrounded in their own seminal plasma (SP), from either fractions/portions or the entire ejaculate, on post-thawing sperm quality and functionality. Ejaculates collected as bulk (BE) or as separate (first 10 mL of SRF [P1] and rest of SRF [P2]) from 10 boars were held 24 h at 15-17°C and then frozen. Some bulk ejaculate samples were frozen immediately after collections as Control. In addition, epididymal sperm samples from the same 10 boars were collected post-mortem and extended in SP from P1 (EP1), P2 (EP2) and post SRF (EP3), and also held 24 h before freezing for a better understanding of the influence of SP on boar sperm cryopreservation. The sperm quality (motility, evaluated by CASA, and viability, evaluated by flow cytometry) and functionality (flow cytometry assessment of plasma membrane fluidity, mitochondrial membrane potential and intracellular generation of reactive oxygen species [ROS] in viable sperm) were evaluated at 30, 150 and 300 min post-thaw. Post-thawing sperm quality and functionality of P1 and P2 were similar but higher (p < 0.01) than BE samples. Control samples showed higher (p < 0.01) post-thaw sperm quality and functionality than BE samples. Post-thawing sperm quality and functionality of EP1 and EP2 were similar but higher (p < 0.05) than EP3. These results showed that boar sperm from BE are more cryosensitive than those from the SRF, particularly when held 24 h before freezing, which would be attributable to the cryonegative effects exerted by the SP from post SRF.

Keywords: Ejaculate, epididymis, sperm, seminal plasma, porcine.
Introduction

Frozen-thawed (FT) boar sperm, other than the routinely used liquid-stored (LS), should be used in artificial insemination (AI) programs, given their additional benefits regarding biosecurity, international exchange and genetic improvement [29]. Despite these primary compelling benefits, FT-sperm is not yet widely used owing to its still notorious low cryosurvival and the short lifespan depicted by the sperm cryosurviving [26]. Therefore, research is still pursuing alternative ways to increase sperm cryosurvival.

Customary removal of the seminal plasma (SP) before freezing, in order to concentrate sperm for further extension in cooling and freezing extenders, has motivated alternative studies aiming to elucidate the role of native SP on boar sperm freezability. Despite many studies has attempted to tackle its role over the recent years, the topic is still pending, since results had been inconclusive [8]. It is, however, widely accepted that holding sperm suspended in their own native SP prior to cooling improves sperm resistance to cold shock [18,24]. Consequently, such procedure is praxis in conventional cryopreservation protocols, albeit there are substantial differences regarding either the duration of the SP-exposure [11,15,31,35,36] or the evidence of its effectiveness on sperm freezability. Some authors reported that SP-exposure improves it [37], while others found that its influence is inconsequent or even detrimental [12,21]. One explanation for these divergent results would be the lack of agreement among studies for the origin and characteristics of the SP surrounding the sperm used for the testing. The boar ejaculate is expelled in rather easily identifiable fractions, each one of these containing varying sperm numbers (or even their absence) but, as importantly, different composition of the SP. The SP composition and volume follows a differential secretion of accessory glands, and the differential presence of epididymal fluid [29]. Thus, unless the ejaculate is collected in one single container, marking its classical large volume and low sperm numbers per mL;
when fractions are collected they show clear differences among a sperm-rich fraction (SRF) and the following sperm-poor fraction, the so-called post-sperm-rich fraction (PSRF). With practice during sperm collection, portions of these can also be collected as for instance the sperm-peak portion [29]. While sperm numbers present defines these fractions, differences are also clear between fractions for the SP, particularly regarding its protein contents [30].

The traditional glove hand method used for manual ejaculate collection in pigs allows for easy sampling of different fractions or even portions within fractions. The SRF is usually the only fraction collected for freezing [9,17,18], although some cryopreservation protocols contemplate the entire ejaculation [4,14,27]. In contrast, other studies suggest only freezing the first 10 mL of the SRF, the sperm-peak portion because the SP of this portion seems to have a greater cryoprotective effect [29,32]. Although these results would prove to the SP surrounding sperm at ejaculation influences boar sperm freezability, it remains unclear the magnitude of this influence, which could depend of fractions/portions of the ejaculate. The main purpose of this study is clarifying this issue. Clarifying this issue is particularly relevant today because, for productivity and hygienic reasons, semi-automatic systems, such as Collectis® [2], are successfully replacing the usual method of the gloved hand for collection of boar ejaculates. These semi-automatic systems require collecting the bulk ejaculate, increasing the proportion of SP in the collected semen samples, and therefore, the putative influence of SP on the capability of boar spermatozoa to withstand the cryopreservation process may be more critical. In addition, it is likely that boar sperm will be held some time surrounded in SP before beginning the freezing process because AI centers are typically located far from the freezing facilities and the time required for ejaculate transport varies. The purpose of this experimental study was to evaluate the influence of native SP, from fractions/portions or
the entire ejaculate, on boar sperm freezability. Additionally, epididymal sperm were held in SP before freezing for a better understanding of the influence of SP.

**Material and methods**

**Reagents and Media**

Unless otherwise stated, all of the chemicals used in the experiments were analytical grade and purchased from Sigma-Aldrich Co. (St. Louis, MO, USA). The basic medium used for semen extension was Beltsville Thawing Solution (BTS: composed of 205 mM glucose, 20.4 mM sodium citrate, 10.0 mM KCl, 15.0 mM NaHCO$_3$, and 3.6 mM EDTA, pH 7.2, and 290-300 mOsmol/kg) supplemented with kanamycin sulfate (0.05 mM). EDTA-free phosphate-buffered saline (PBS; 137 mM NaCl, 2.7 mM KCl, 1.5 mM KH$_2$PO$_4$, 8.1 mM Na$_2$HPO$_4$, pH 6.8, and 280-300 mOsmol/kg) was used to dilute fluorochromes and extend sperm samples for flow-cytometric analysis. Sperm were frozen using a basic freezing medium (FE) containing 80% (v/v) Tris-citric acid-glucose extender (111 mM Trizma Base, 31.4 mM monohydrate citric acid, 185 mM glucose) and 20% (v/v) egg yolk, supplemented with 100 µg/mL kanamycin sulfate (pH 7.2; 295-300 mOsmol/kg).

**Experimental design**

The experimental design is shown in Figure 1. All of the procedures that involved animals were performed according to international guidelines and were approved by the Bioethics Committee of Murcia University (research code: 639/2012). The boars (Large-White) were three years old, healthy and with a proven history of fertility after conventional AI with liquid semen. At the moment when experiments were drawn, semen was regularly
collected (twice week) for commercial production of AI-semen doses in a commercial insemination center (AIM Iberica, Calasparra, Murcia, Spain). The boars were, 3-5 days after the month-long experimentation with ejaculates ended, slaughtered at a local slaughterhouse (Mercamurcia, Murcia, Spain). The testes and epididymides from each boar were removed immediately after slaughter and transported in insulated containers at 20-23 °C to the Andrology Laboratory of Veterinary Teaching Hospital of University of Murcia (VTH), arriving within 30 min after collection.

Ejaculates, bulk or in portions, were collected during the month prior to slaughter using the gloved-hand method and following the standard operating procedure. The portions of ejaculate collected separately were the first 10 mL of SRF (so-called P1), the remaining SRF (so-called P2) and the PSRF (only as a source of SP as it contained too few sperm). All semen samples from the bulk ejaculate or ejaculate portions met the following criteria: 70% total motile sperm (subjectively evaluated using light microscopy) and 80% sperm with normal morphology and intact acrosome ridges (evaluated using phase contrast microscopy of sperm samples fixed in buffered 2% glutaraldehyde solution). Semen samples were split in two aliquots immediately after collection; one was used for harvesting SP (see below) and the other was extended in pre-warmed (35 °C) BTS (2:1, v/v). Thereafter, semen and SP-samples were transported at 20-22 °C to the Andrology Laboratory of VTH, arriving within 2 h of collection. Once in the laboratory, the semen samples from bulk ejaculates were split into two aliquots and one was frozen immediately (as Control) whereas the other was stored at 17°C during 24 h before freezing (BE sample). P1- and P2-semen samples were also stored at 17°C during 24 h before freezing. A holding time as long as 24 h was chosen because it allows any AI center, even those located in remote locations, to ship semen samples overnight to freezing facilities. Also, to hold the semen samples 24 h before freezing has resulted in high sperm cryosurvival
rates and excellent fertility outcomes when SRF were frozen using the cryopreservation protocol used in this experiment [16,28]. The SP from each one of SP-sources (first 10 mL of SRF [SP\textsubscript{1}], rest of SRF [SP\textsubscript{2}] and PSRF [SP\textsubscript{3}]) was harvested via double centrifugation (Rotofix 32 A, Hettich Zentrifugen\textsuperscript{®}, Germany) at 1,500 x g for 10 min at rt. The SP was collected by aspirating fluid above the sperm pellet, preventing disturbance of the pellet. The SP samples were stored at -80 °C until used.

Epididymal samples were collected following the procedure described by Martinez-Pastor et al. [20] with slight modifications. Briefly, the epididymides were removed and dissected avoiding blood contamination. Cauda epididymal luminal fluid was collected by retrograde washing from the vas deferens using a syringe loaded with 1 mL BTS, injecting air afterwards until all the fluid was flushed out. The harvested caudal epididymal fluids, from both epididymides of the same boar, were collected and mixed in a sterile Petri dish, and the sperm concentration (using an SP-100 NucleoCounter; ChemoMetec A/S, Allerød, Denmark) and motility (see above) was then evaluated. All epididymal samples showed a total sperm motility above 70%. Epididymal samples were split into three aliquots, extended (to 300 x 10\textsuperscript{6} sperm/mL) with each one of the SP-sources, SP\textsubscript{1} (EP1 sample), SP\textsubscript{2} (EP2 sample) and SP\textsubscript{3} (EP3 sample); and stored at 17 °C during 24 h before freezing.

**Sperm cryopreservation**

The semen samples were centrifuged (Megafuge 1.0 R, Heraeus, Hanau, Germany) at 17 °C for 3 min at 2400 x g, and the sperm pellets were frozen using the straw freezing procedure described by Hernandez et al. [16]. Briefly, sperm pellets were extended in FE to a concentration of 1.5 x 10\textsuperscript{9} sperm/mL. After cooling to 5 °C for 150 min, the sperm
were re-extended with FE-glycerol-Equex extender (89.5% FE + 1.5% Equex STM (v/v)) (Nova Chemical Sales, Scituate, MA, USA) and 9% glycerol (v/v); pH 6.2; 1700-1730 mOsmol/kg) to a final concentration of 1.0 x 10^9 sperm/mL. The sperm were thereafter packed into 0.5 ml polyvinyl chloride (PVC) French straws (Minitüb, Tiefenbach, Germany) and frozen using a controlled-rate-freezing machine (IceCube 1810, Minitüb, Germany). The straws remained in liquid nitrogen for at least 1 week before thawing, which was performed in a circulating water bath at 37 °C for 20 s. The thawed sperm samples were extended in BTS (1:1 v/v) and incubated at 37 °C for up to 300 min.

Assessment of sperm quality and functionality
The sperm were assessed according to conventional quality parameters (total and progressive motility and viability) and functionality at 30, 150 and 300 min after thawing in semen samples stored in dark at 37 °C. Functionality was evaluated in terms of plasma membrane fluidity, intracellular H_2O_2 (hydrogen peroxide) generation and mitochondrial membrane potential. All of these assessments, except motility, which was evaluated using a computer-assisted sperm analyzer (CASA), were performed using flow cytometry. Flow cytometric analyses were carried out using a BD FACSCanto II (Becton Dickinson Co, Franklin Lakes, NJ, USA). The optical configuration includes one octagon and two trigon detector arrays. The octagon contains five photomultiplier tubes (PMT) and detects light from the 488-nm (blue) laser. Both trigons contain two PMTs. One detects light from the 633-nm (red) laser and the other detects light from the 633-nm (violet) laser. Data collection and compensation for spectral overlap were performed using BD FACSDiva Software (Becton Dickinson Co). Events were triggered by forward scatter (FSC) and side scatter (SSC) and non-sperm events were gated out based on Hoechst 33342 (H-42) fluorescence (DNA content). Acquisition was stopped after 10000 H-42
positive events were recorded. The fluorescence spectrum of H-42 was detected using a 450/50 nm band-pass (BP) filter. The fluorescence spectra of propidium iodide (PI) and fluorescein-conjugated peanut agglutinin (PNA-FITC) were detected using a 670 nm long-pass (LP) filter and a 530/30 nm BP filter, respectively. The fluorescence spectrum of Merocyanine 540 (M-540) was detected using a 585/42 nm BP filter, and Yo-Pro-1 was detected using a 530/30 nm BP filter. The fluorescence spectrum of dichlorofluorescein (DCF) was detected using a 530/30 nm BP following excitation at 488 nm. The fluorescence spectrum of Mitotracker Deep Red was detected using a 660/20 BP filter. All fluorescence parameters were log transformed, while FSC and SSC signal were processed in lineal mode.

*Sperm motility*

Sperm motility was objectively evaluated using ISASV1® CASA (Proiser R+D, Valencia, Spain) operating up to 100 videoframes/s and following a modification of the procedure described by Cremades et al. [6]. For each evaluation, a 5 µL sperm sample (30 x 10^6 spermatozoa/mL) was placed in a pre-warmed (38 °C) Makler counting chamber (Sefi Medical Instruments, Haifa, Israel), and at least five fields were analyzed to evaluate a minimum of 400 sperm. Sperm motility was recorded as the percentage of total motile sperm (average path velocity ≥ 20 µm/s) and the percentage of motile sperm showing rapid and progressive movement (straightness of the average path ≥ 40 %).

*Sperm viability*

Sperm viability was evaluated by simultaneously assessing the plasma membrane and acrosome integrity using a triple-fluorescence procedure [19]. Briefly, 100 µL aliquots containing 3 x 10^6 spermatozoa were transferred into culture tubes containing 3 µL of H-
42 (0.05 mg/mL in PBS), 2 µL of PI (0.5 mg/mL stock solution in PBS), and 2 µL of PNA-FITC (100 µg/mL stock solution in PBS). Samples were mixed and incubated at 37 ºC for 10 min in the dark. Immediately before analysis, 400 µL of PBS was added to each sample. The percentage of viable sperm population exhibiting intact plasma and acrosomal membranes (PI negative and PNA-FITC negative) is shown in the Results section.

**Plasma membrane fluidity**

The fluidity of the plasma membrane was evaluated by staining sperm with H-42, M-540 and Yo-Pro®-1 (Molecular Probes Europe BV, Leiden, The Netherlands). Aliquots of 50 µL of BTS-extended spermatozoa (30 x 10^6 spermatozoa/mL) were extended in 950 µL of PBS containing 1.5 µL of H-42 (0.05 mg/ml in PBS) and 1 µL of Yo-Pro-1 (25 µM in DMSO) and incubated at 38 ºC for 8 min in the dark. Then, 2.6 µL of M-540 (1 mM in DMSO) was added to each sample, and the samples were incubated for another 2 min under the same conditions before flow cytometric analysis. The sperm Yo-Pro-1 negative and M-540 positive were considered as viable with high plasma membrane fluidity.

**Intracellular ROS generation**

The intracellular generation of ROS in viable sperm was measured in terms of hydrogen peroxide (H_2O_2) using 5-(and-6) chloromethyl-20,70-dichlorodihydro-fluorescein diacetate acetyl ester (CM-H_2DCFDA) following the procedure described by Guthrie and Welch [13]. For each sperm sample, a 50 µL aliquot of BTS-diluted sperm (30 x 10^6 sperm/mL) was diluted in 950 µL of PBS containing 1.5 µL of H-42 (0.05 mg/mL in PBS), 1 µL of PI (1 mg/mL in PBS), and 1 µL of H_2DCFDA (1 mM in DMSO) and incubated at 38 ºC for 30 min in the dark prior to flow cytometric analysis. H_2DCF is
oxidized by H$_2$O$_2$ into DCF. The mean fluorescence intensity of DCF in viable sperm (PI negative) was expressed as fluorescence units (FU) per 10$^8$ live sperm.

**Mitochondrial membrane potential**

The mitochondrial membrane potential was evaluated with Mitotracker Deep Red 633 (Mitotracker, M-22426, Molecular Probes Europe BV) using a variant of protocol described by Domínguez-Rebolledo et al. [10]. Briefly, a 100 µL sperm sample (30 x 10$^6$ sperm/mL in PBS) was transferred to culture tubes containing 3 µL H-42 (0.05 mg/mL in PBS), 2 µL PI (0.5 mg/mL in PBS) and 0.5 µL Mitotraker (20 µM in PBS of a stock solution of 1 mM in DMSO). The samples were mixed and incubated at 38 °C in the dark for 15 min. Immediately before analysis by flow cytometry, 400 µL PBS was added to each sample, and the samples were mixed. The percentage of viable sperm (PI negative) with high mitochondria membrane potential (Mitrotraker positive) was reported in the Results section.

**Statistical analysis**

Statistical analyses were performed using the SPSS 19 Statistics package (SPSS Inc., Chicago, IL, USA) as follows. The residual data of each sperm quality and functionality variable was evaluated using the Kolmogorov-Smirnov test to confirm the assumption of normality and non-normally distributed data were arcsine transformed before statistical analysis. Data were analyzed using ANOVA procedures with either semen sample or SP-source and post-thaw incubation time (30, 150 and 300 min) as the main effects plus their interactions. When the interaction was not significant, it was omitted from the model. Whenever the main effects were significant, means were compared using Bonferroni test. Means of sperm quality and functionality post-thawing of semen and epididymal samples
were compared by unpaired Student’s t test corrected for inequality of variances (Levene’s test). Statistical significance was defined as \( p < 0.05 \). Data are shown as the means ± SEMs.

**Results**

*Effect of holding time before freezing on the freezability of semen samples*

Both semen sample and incubation time after thawing influenced (\( p < 0.01 \)) post-thaw sperm quality. Sperm quality decreased as post-thaw incubation time increased (\( p < 0.01 \)). The semen sample x incubation time interaction was significant (\( p < 0.01 \)) for total and progressive sperm motility, albeit the four semen samples showed the same pattern in the three post-thaw incubation times (Table 1). Total and progressive sperm motility of P1 and P2 samples were similar and always higher (\( p < 0.01 \)) than BE samples, regardless of post-thaw incubation time. In addition, sperm motility parameters of P1 were higher (\( p < 0.01 \)) than Control at 150 and 300 min post-thawing. Sperm viability of P1 and P2 was higher (\( p < 0.01 \)) than Control and BE samples, showing the last one the lower (\( p < 0.01 \)) percentages (Table 1).

The proportion of viable sperm with high plasma membrane fluidity was influenced (\( p < 0.01 \)) by semen sample and post-thaw incubation time, whereas the interaction was not significant. The P1 and BE samples showed the lowest (\( p < 0.01 \)) and highest (\( p < 0.01 \)) proportions, respectively (Table 1). Semen sample influenced the intracellular generation of \( \text{H}_2\text{O}_2 \) (\( p < 0.01 \)), whereas incubation time did not. The \( \text{H}_2\text{O}_2 \) levels generated by P1 and P2 samples were similar than Control samples and lower (\( p < 0.05 \)) than BE samples, regardless of the post-thaw incubation time (Table 1). Percentage of viable sperm with high mitochondrial membrane potential was not influenced by semen sample. However, it was influenced by incubation time post-thawing (\( p < 0.01 \)), decreasing the percentages
as post-thaw incubation time increased, regardless of the semen samples considered (Table 1).

*Influence of SP-source on freezability of epididymal sperm*

SP-source influenced (p < 0.01) total sperm motility and viability after thawing. Post-thaw incubation time negatively affected (p < 0.05) motility variables and sperm viability. SP-source x post-thaw incubation time interaction was not significant for any sperm quality parameter. The EP1 and EP2 samples showed higher (p < 0.001) percentages of total motility than EP3 samples. The EP1 and EP3 samples showed the highest (p < 0.01) and lowest (p < 0.01) percentages of viable sperm (Table 2). SP-source did not influence the proportion of viable sperm with both high plasma membrane fluidity and high mitochondrial membrane potential, whereas their post-thaw incubation time did (p < 0.01). SP-source x post-thaw incubation time interaction was not significant for both functionality parameters. The percentages of viable sperm with high plasma membrane fluidity increased whereas those with high mitochondrial membrane potential decreased with incubation time (Table 2). The intracellular generation of H$_2$O$_2$ in viable sperm was influenced (p < 0.05) by SP-source and incubation time, being not significant the interaction. The generation of H$_2$O$_2$ increased during incubation in all epididymal sperm samples, showing EP1 and EP2 samples lower (p < 0.05) generation than EP3 samples (Table 2).

*Sperm freezability differences between semen and epididymal samples*

Post-thaw sperm quality, in terms of motility parameters and viability, was lower (p < 0.05) in ejaculates than in epididymal samples throughout the three post-thawing incubation times (Table 3). Regarding post-thaw functionality of viable sperm (Table 3),
the percentage of those cells with high plasma membrane fluidity just differed between BE and EP3 samples, showing BE samples the highest (p < 0.05) percentage in all post-thaw incubation times. The intracellular H$_2$O$_2$ generation was higher (p < 0.05) in semen than in epididymal samples at 30 min; and only in BE samples than EP3 samples at 150 min. The percentage of sperm with high mitochondrial membrane potential was lower (p < 0.05) in semen compared to epididymal samples throughout all post-thaw incubation times.

**Discussion**

From a practical point of view, semen samples for freezing should withstand a holding time of several hours at cooling temperatures before freezing because AI-centers, where ejaculates are collected, are often away from freezing facilities. In this sense, 24 h seems a holding time sufficiently long for any AI centers, even those located in remote locations, to ship semen samples for freezing. This long holding time, already incorporated in some cryopreservation protocols, has proven to be effective since it allows achieving acceptable cryosurvival rates and fertility outcomes when held semen samples were of SRF origin [12]. Results from our laboratory, achieving cryosurvival rates above 80% [18] and fertility outcomes as high as 85.6 % of farrowing rates and 12.6 piglets born per litter [28], support the effectiveness of 24h of holding time to cryopreserve semen samples from SRF. Recently has been corroborated the suitability of a holding time of 24 h before freezing to improve cryotolerance of boar SRF-sperm [37].

However, the scenario we are confronted now is whether this long holding time before freezing would also be effective to cryopreserve spermatozoa collected from the bulk ejaculate, instead than being those fortuitously present in the SRF. To contemplate this
scenario is now mandatory since AI-centers are incorporating semiautomatic systems for ejaculate collection that do not allow to collect separate ejaculate fractions. The results of the present study clearly demonstrated that a holding time before freezing as long as 24 h impairs sperm freezability of spermatozoa retrieved from a bulk ejaculate. Moreover, this study also shows that sperm from the SRF withstands cryopreservation better than those from the bulk ejaculate when semen samples were held 24 h before freezing. Putative freezability differences between sperm from the SRF or the BE have not been comparatively evaluated previously. Instead, some previous studies have focused on evaluating differences between first 10 mL of SRF and rest of ejaculate, showing that sperm from the first withstands cryopreservation better than sperm from latter fractions [22,23,32]. These studies, that contemplate a holding time before freezing of only 2 h, found that the differences were particularly evident for sperm motility and not in all boars. Aware of these studies, semen samples from the first 10 ml of SRF were included in the present experiment and they were also held during 24 h before freezing. Cryosurvival rates and functionality of alive sperm from the first 10 ml of the SRF were similar to those from of rest of SRF, which was in agreement with previous results [34], but, as noticeably and in contrast to the above studies, clearly better of those from the BE. So, it seems that extending the holding time before freezing from 2 to 24 h makes differences in sperm freezability between SRF (including the first 10 mL) and bulk ejaculate become more apparent. The differences, evaluated in FT-sperm incubated over 300 min, were not only clearly evident in classical quality parameters, such as motility and viability, but also in some functionality sperm parameters, specifically plasma membrane fluidity and intracellular generation of H$_2$O$_2$, both being higher in the viable FT-sperm from the bulk ejaculate than those from both first 10 mL of SRF and rest SRF. This functional differences are particularly noticeable because a greater fluidity of plasma membrane
together with a higher endogenous generation of H$_2$O$_2$, the primary ROS generated by boar spermatozoa [1], are events exhibited by boar sperm exposed to capacitation conditions [19]. Thus, it seems that FT-sperm of semen samples held 24 h before freezing from bulk ejaculate are more sensitive to premature capacitation than those from SRF, including its first 10 mL (peak-sperm portion). The differences in sperm cryosurvival and functionality evidenced between SRF (including the peak-sperm portion) and BE should be attributed to the composition of SP surrounding the sperm during the holding time.

Certainly, differences in either SP proteins or bicarbonate concentration among ejaculate portions have been shown responsible for differences in post-thaw sperm functionality [32]. From the present results, it seems that the SP from the entire ejaculate is less cryoprotective than that from the SRF alone (including the peak-portion). Moreover, semen samples held long time before freezing surrounded in the SP of the entire ejaculate deteriorate during cryosurvival, perhaps due to a negative influence of the PSRF that counteracts the cryoprotective effects of SP from SRF (including first 10 mL).

To test the above hypothesis, samples collected from the epididymides of the same boars were held 24 h extended in SP from the three portions of the ejaculate (peak-sperm portion, rest of the SRF and from the PSRF). Cryosurvival rates of epididymal samples held in SP either the sperm-peak portion or the rest of the SRF were similar, mimicking results achieved with comparative ejaculated semen samples, but were better than sperm exposed to PSRF. In addition, sperm held in SP either the sperm-peak portion or the rest of the SRF generate less H$_2$O$_2$ than those exposed to PSRF. Overall, the results show that indeed the SP from PSRF is less sperm-cryoprotective than SP from SRF, which might be attributable to its higher levels of bicarbonate (and a more alkaline pH) [33], low concentrations of chemicals with antioxidant capacity [3] and highest content in HBPs proteins that under in vitro conditions failed to preserve the sperm viability and
functionality [5]. However, given these peculiarities in the composition of SP of PSRF and looking at the above results achieved with ejaculated sperm, a greater loss in post-thaw quality and functionality in the epididymal sperm held diluted in SP from PSRF was expected. The differences in both the quality and the post-thaw sperm function observed between the epididymis and ejaculated samples could give an explanation for this unexpected finding. Since we did not test sperm from PSRF (owing to the very low sperm numbers present), we compared the epididymal sperm held diluted in SP from PSRF and those exposed to that from entire ejaculate. Epididymal sperm always showed better post-thaw quality and functionality than those ejaculated, regardless of the ejaculate portion from which they came or the SP-source used to held epididymal sperm before freezing. Then, it is clear that epididymal sperm are more cryoresilient that ejaculated sperm, which was in agreement with previous report [25], which is attributed the protective effect of epididymal fluid, containing low bicarbonate levels and absence of relative high levels of proteins [7 and references therein,29]. Probably this greater protection of epididymal fluid explains why epididymal sperm withstood cryopreservation better irrespective of the exposure to PS from either PSRF or entire ejaculate.

In conclusion, our results showed that boar sperm from bulk ejaculate are more cryosensitive than those from the sperm rich ejaculate fraction, particularly when held 24 h before freezing surrounded by their own seminal plasma, which would be attributable to the effects exerted by the seminal plasma on sperm during the holding time. This evidence challenges the suitability of the collection of bulk ejaculates instead the hitherto usual sperm rich ejaculate fraction for cryopreservation, especially if semen samples must be hold some time before freezing to be shipped from a place to another for freezing. Although further fertility trials are needed to confirm in the farm the practical relevance of these results, they seem conclusive enough to recommend to AI-centers to continue
collecting rich ejaculate fraction instead of bulk ejaculate for sperm cryopreservation. Otherwise, the sperm cryopreservation procedure should be modified if the collection of bulk ejaculates is mandatory.

Conflict of interest

The authors declare that there is no conflict of interest that could prejudice the impartiality of this paper.

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Figure legend:

**Figure 1.** Schematic view of the experimental design showing how the semen samples from ejaculates and epididymides were handled.

Tables:

**Table 1.** Differences (mean ± SEM) in freezability among semen samples (n = 10) from bulk ejaculate (BE), the first 10 mL of sperm rich ejaculate fraction (P1) and the rest of sperm rich ejaculate fraction (P2) held at 15-17°C during 24 h before freezing surrounded in their own seminal plasma. Semen samples from entire ejaculate frozen immediately after collection were used as Control.

<table>
<thead>
<tr>
<th>Sperm parameters (%)</th>
<th>Semen Samples</th>
<th>Post-thawing incubation time (min)</th>
<th>30</th>
<th>150</th>
<th>300</th>
<th>mean</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Control</td>
<td>BE</td>
<td>P1</td>
<td>P2</td>
</tr>
<tr>
<td><strong>Total sperm motility</strong></td>
<td></td>
<td></td>
<td>Control</td>
<td>BE</td>
<td>P1</td>
<td>P2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>55.1 ± 2.0(^a)</td>
<td>42.8 ± 2.1(^b)</td>
<td>61.6 ± 1.7(^a)</td>
<td>57.7 ± 1.9(^a)</td>
</tr>
<tr>
<td></td>
<td>mean</td>
<td></td>
<td>56.1 ± 1.3</td>
<td>39.4 ± 1.3</td>
<td>10.5 ± 0.8</td>
<td></td>
</tr>
<tr>
<td><strong>Progressive sperm motility</strong></td>
<td></td>
<td></td>
<td>Control</td>
<td>BE</td>
<td>P1</td>
<td>P2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>41.5 ± 0.5(^a)</td>
<td>25.9 ± 1.0(^b)</td>
<td>43.8 ± 1.5(^a)</td>
<td>41.8 ± 1.5(^a)</td>
</tr>
<tr>
<td></td>
<td>mean</td>
<td></td>
<td>39.9 ± 1.1</td>
<td>29.6 ± 1.1</td>
<td>7.2 ± 0.5</td>
<td></td>
</tr>
<tr>
<td><strong>Sperm viability</strong></td>
<td></td>
<td></td>
<td>Control</td>
<td>BE</td>
<td>P1</td>
<td>P2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>50.3 ± 1.2</td>
<td>40.8 ± 2.1</td>
<td>61.0 ± 2.0</td>
<td>57.6 ± 1.7</td>
</tr>
<tr>
<td></td>
<td>mean</td>
<td></td>
<td>54.7 ± 1.3(^a)</td>
<td>48.3 ± 1.1(^b)</td>
<td>38.9 ± 1.0(^c)</td>
<td></td>
</tr>
<tr>
<td><strong>High plasma membrane</strong></td>
<td></td>
<td></td>
<td>Control</td>
<td>BE</td>
<td>P1</td>
<td>P2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>4.6 ± 0.3</td>
<td>8.2 ± 0.5</td>
<td>6.7 ± 0.4</td>
<td>10.3 ± 0.6</td>
</tr>
</tbody>
</table>

\(^a\) vs. Control; \(^b\) vs. BE; \(^c\) vs. P1; \(^d\) vs. P2
Table 2. Differences (mean ± SEM) in freezability among epididymal sperm samples extended in seminal plasma collected from either the first 10 mL of sperm rich ejaculate fraction (EP1), the rest of sperm rich ejaculate fraction (EP2) or the post sperm rich ejaculate fraction (EP3), and held at 15-17°C during 24 h before freezing.

<table>
<thead>
<tr>
<th>Sperm parameters (%)</th>
<th>Epiddidymal Samples</th>
<th>Post-thawing incubation time (min)</th>
<th>mean</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>30</td>
<td>150</td>
</tr>
<tr>
<td>Total sperm motility</td>
<td>EP1</td>
<td>72.1 ± 1.3</td>
<td>57.0 ± 2.2</td>
</tr>
<tr>
<td></td>
<td>EP2</td>
<td>70.8 ± 1.8</td>
<td>56.8 ± 2.0</td>
</tr>
<tr>
<td></td>
<td>EP3</td>
<td>66.1 ± 1.6</td>
<td>55.2 ± 1.5</td>
</tr>
<tr>
<td></td>
<td>mean</td>
<td>69.2 ± 1.0&lt;sup&gt;a&lt;/sup&gt;</td>
<td>56.3 ± 1.1&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Progressive sperm motility</td>
<td>EP1</td>
<td>51.4 ± 1.5</td>
<td>43.2 ± 1.0</td>
</tr>
<tr>
<td></td>
<td>EP2</td>
<td>52.1 ± 1.8</td>
<td>41.6 ± 0.7</td>
</tr>
<tr>
<td></td>
<td>EP3</td>
<td>48.9 ± 1.7</td>
<td>42.4 ± 1.4</td>
</tr>
<tr>
<td></td>
<td>mean</td>
<td>50.8 ± 1.0&lt;sup&gt;a&lt;/sup&gt;</td>
<td>42.4 ± 0.6&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Sperm viability</td>
<td>EP1</td>
<td>66.2 ± 1.0</td>
<td>57.5 ± 1.8</td>
</tr>
<tr>
<td></td>
<td>EP2</td>
<td>62.3 ± 1.1</td>
<td>56.6 ± 1.9</td>
</tr>
<tr>
<td></td>
<td>EP3</td>
<td>59.6 ± 1.8</td>
<td>53.9 ± 2.1</td>
</tr>
<tr>
<td></td>
<td>mean</td>
<td>62.7 ± 0.9&lt;sup&gt;a&lt;/sup&gt;</td>
<td>56.0 ± 1.1&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>High plasma membrane fluidity</td>
<td>EP1</td>
<td>3.6 ± 0.4</td>
<td>6.9 ± 0.7</td>
</tr>
<tr>
<td></td>
<td>EP2</td>
<td>3.6 ± 0.4</td>
<td>6.4 ± 0.7</td>
</tr>
<tr>
<td></td>
<td>EP3</td>
<td>4.1 ± 0.4</td>
<td>6.6 ± 0.5</td>
</tr>
<tr>
<td></td>
<td>mean</td>
<td>3.8 ± 0.2&lt;sup&gt;a&lt;/sup&gt;</td>
<td>6.6 ± 0.4&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Intracellular H₂O₂ generation (FU x 10⁸/sperm)</td>
<td>EP1</td>
<td>18.9 ± 1.1</td>
<td>23.9 ± 1.4</td>
</tr>
<tr>
<td></td>
<td>EP2</td>
<td>18.8 ± 1.5</td>
<td>23.9 ± 2.0</td>
</tr>
<tr>
<td></td>
<td>EP3</td>
<td>21.3 ± 1.4</td>
<td>24.9 ± 1.6</td>
</tr>
<tr>
<td></td>
<td>mean</td>
<td>19.6 ± 0.8&lt;sup&gt;a&lt;/sup&gt;</td>
<td>24.2 ± 0.9&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>High mitochondrial membrane potential</td>
<td>EP1</td>
<td>54.4 ± 0.4</td>
<td>50.7 ± 1.3</td>
</tr>
<tr>
<td></td>
<td>EP2</td>
<td>53.1 ± 0.7</td>
<td>49.4 ± 1.3</td>
</tr>
</tbody>
</table>

FU: fluorescence units. a-c: indicate differences (p < 0.05) among semen samples (rows); α, β, γ: indicate differences (p < 0.05) among incubation times (columns).
**Table 3:** Differences in sperm freezability in terms of quality and functionality between semen and epididymal samples at 30, 150 and 300 min post-thawing.

<table>
<thead>
<tr>
<th>Incubation time (min)</th>
<th>SP source</th>
<th>Sperm source</th>
<th>Total motility (%)</th>
<th>Progressive motility (%)</th>
<th>Viability (%)</th>
<th>High plasma membrane fluidity (%)</th>
<th>H$_2$O$_2$ generation (FU x 10$^8$)</th>
<th>High mitochondrial membrane potential (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>30</td>
<td>SP$_1$</td>
<td>Ejaculate</td>
<td>61.4 ± 2.4$^a$</td>
<td>43.8 ± 2.1$^a$</td>
<td>60.9 ± 2.0$^a$</td>
<td>2.6 ± 0.3</td>
<td>25.4 ± 1.3$^a$</td>
<td>50.9 ± 0.7$^a$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Epididymis</td>
<td>71.1 ± 1.9$^a$</td>
<td>51.0 ± 1.7$^a$</td>
<td>66.2 ± 1.0$^a$</td>
<td>3.6 ± 0.4</td>
<td>19.3 ± 1.3$^a$</td>
<td>54.4 ± 0.4$^a$</td>
</tr>
<tr>
<td>150</td>
<td>SP$_2$</td>
<td>Ejaculate</td>
<td>57.7 ± 2.4$^a$</td>
<td>41.7 ± 2.1$^a$</td>
<td>57.6 ± 1.7$^a$</td>
<td>3.8 ± 0.3</td>
<td>25.7 ± 1.4$^a$</td>
<td>49.3 ± 0.5$^a$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Epididymis</td>
<td>71.0 ± 2.2$^a$</td>
<td>52.1 ± 2.5$^a$</td>
<td>62.4 ± 1.2$^a$</td>
<td>3.6 ± 0.4</td>
<td>18.8 ± 1.5$^a$</td>
<td>53.1 ± 0.7$^a$</td>
</tr>
<tr>
<td>300</td>
<td>SP$_3$</td>
<td>Ejaculate</td>
<td>42.8 ± 3.1$^b$</td>
<td>25.9 ± 1.1$^b$</td>
<td>40.8 ± 2.1$^b$</td>
<td>8.2 ± 0.5$^b$</td>
<td>30.9 ± 1.9$^b$</td>
<td>49.3 ± 0.8$^b$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Epididymis</td>
<td>66.1 ± 2.5$^a$</td>
<td>48.9 ± 1.7$^b$</td>
<td>59.6 ± 1.8$^b$</td>
<td>4.1 ± 0.4$^b$</td>
<td>21.3 ± 1.4$^b$</td>
<td>55.4 ± 0.4$^b$</td>
</tr>
<tr>
<td></td>
<td>SP$_1$</td>
<td>Ejaculate</td>
<td>44.9 ± 2.2$^b$</td>
<td>34.4 ± 2.0$^b$</td>
<td>52.2 ± 2.1$^a$</td>
<td>5.9 ± 0.4</td>
<td>27.2 ± 2.1$^a$</td>
<td>46.4 ± 0.8$^a$</td>
</tr>
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<td>Epididymis</td>
<td>57.0 ± 2.7$^b$</td>
<td>43.1 ± 1.7$^b$</td>
<td>57.5 ± 1.8$^b$</td>
<td>6.9 ± 0.7</td>
<td>24.4 ± 1.7$^b$</td>
<td>50.7 ± 1.3$^b$</td>
</tr>
<tr>
<td></td>
<td>SP$_2$</td>
<td>Ejaculate</td>
<td>42.0 ± 1.8$^b$</td>
<td>30.9 ± 1.6$^b$</td>
<td>50.3 ± 1.4$^b$</td>
<td>6.2 ± 0.4</td>
<td>25.3 ± 1.4$^b$</td>
<td>45.4 ± 0.9$^b$</td>
</tr>
<tr>
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<td>Epididymis</td>
<td>56.8 ± 2.9$^b$</td>
<td>41.6 ± 1.0$^b$</td>
<td>56.6 ± 1.9$^b$</td>
<td>6.4 ± 0.7</td>
<td>23.9 ± 2.0$^b$</td>
<td>49.4 ± 1.3$^b$</td>
</tr>
<tr>
<td></td>
<td>SP$_3$</td>
<td>Ejaculate</td>
<td>26.6 ± 1.9$^b$</td>
<td>19.0 ± 1.6$^b$</td>
<td>37.9 ± 1.9$^b$</td>
<td>10.3 ± 0.6$^b$</td>
<td>31.4 ± 1.3$^a$</td>
<td>48.7 ± 0.5$^a$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Epididymis</td>
<td>55.2 ± 2.1$^b$</td>
<td>42.4 ± 1.8$^b$</td>
<td>53.9 ± 2.1$^b$</td>
<td>6.6 ± 0.5$^b$</td>
<td>24.9 ± 1.6$^b$</td>
<td>48.7 ± 1.8$^a$</td>
</tr>
<tr>
<td></td>
<td>SP$_1$</td>
<td>Ejaculate</td>
<td>13.7 ± 1.9$^a$</td>
<td>9.4 ± 1.4$^a$</td>
<td>42.0 ± 1.8$^a$</td>
<td>8.8 ± 0.6</td>
<td>27.3 ± 1.3$^a$</td>
<td>31.8 ± 1.3$^a$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Epididymis</td>
<td>44.1 ± 2.7$^a$</td>
<td>32.1 ± 1.4$^a$</td>
<td>51.2 ± 1.3$^a$</td>
<td>9.4 ± 0.9</td>
<td>27.3 ± 1.7$^a$</td>
<td>42.7 ± 0.7$^a$</td>
</tr>
<tr>
<td></td>
<td>SP$_2$</td>
<td>Ejaculate</td>
<td>10.5 ± 1.3$^b$</td>
<td>7.6 ± 1.1$^b$</td>
<td>40.4 ± 1.4$^b$</td>
<td>10.2 ± 0.5</td>
<td>25.7 ± 1.1$^b$</td>
<td>32.5 ± 1.9$^b$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Epididymis</td>
<td>42.5 ± 2.3$^b$</td>
<td>31.5 ± 1.6$^b$</td>
<td>48.5 ± 0.8$^b$</td>
<td>8.9 ± 0.6</td>
<td>27.4 ± 1.7$^b$</td>
<td>42.8 ± 0.9$^b$</td>
</tr>
<tr>
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<td>SP$_3$</td>
<td>Ejaculate</td>
<td>5.5 ± 0.9$^b$</td>
<td>3.7 ± 0.5$^b$</td>
<td>30.2 ± 1.7$^b$</td>
<td>13.7 ± 0.6$^b$</td>
<td>30.9 ± 1.2$^b$</td>
<td>33.3 ± 2.3$^b$</td>
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<td></td>
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<td>Epididymis</td>
<td>40.7 ± 3.1$^b$</td>
<td>29.5 ± 2.3$^b$</td>
<td>44.8 ± 1.8$^b$</td>
<td>9.6 ± 0.9$^b$</td>
<td>29.6 ± 2.4$^b$</td>
<td>41.7 ± 1.1$^b$</td>
</tr>
</tbody>
</table>

FU: fluorescence units. a-b: indicate differences (p < 0.05) between sperm source within each SP source and incubation time.
Boars (n=10)

Ejaculates

Seminal Plasma (SP) sources
- First 10 mL of SRF = SP₁
- Rest of SRF = SP₂
- Post SRF = SP₃

Semen samples
- Bulk ejaculate - 0h* = Control
- Bulk ejaculate = BE
- First 10 mL of SRF = P₁
- Rest of SRF = P₂

Epididymides

Epididymal samples diluted in:
- SP₁ = EP₁
- SP₂ = EP₂
- SP₃ = EP₃

Frozen

>1 week in LN₂

Thawing (37 °C/20s) and dilution in BTS (1:1 vol/vol)

Incubation at 37 °C during 300 min

Sperm quality and functionality evaluated at 30, 150 and 300 min

*Holding time before freezing