Computational flow cytometry reveals that cryopreservation induces spermtptosis but subpopulations of spermatozoa may experience capacitation-like changes


1Laboratory of Equine Reproduction and Equine Spermatology, Veterinary Teaching Hospital, University of Extremadura, Cáceres, Spain, 2Reproduction and Obstetrics Department of Animal Medicine and Surgery, and 3Department of Molecular Biology, University of León, León, Spain

Correspondence should be addressed to F J Peña; Email: fjuanpvega@unex.es

Abstract

The reduced lifespan of cryopreserved spermatozoa in the mare reproductive tract has been attributed to both capacitative and apoptotic changes. However, there is a lack of studies investigating both phenomena simultaneously. In order to improve our knowledge in this particular point, we studied in raw and frozen-thawed samples apoptotic and capacitative markers using a wide battery of test based in flow cytometry. Apoptotic markers evaluated were caspase 3 activity, externalization of phosphatidylserine (PS), and mitochondrial membrane potential. Markers of changes resembling capacitation were membrane fluidity, tyrosine phosphorylation, and intracellular sodium. Conventional and computational flow cytometry using nonlinear dimensionally reduction techniques (t-distributed stochastic neighbor embedding (t-SNE)) and automatic classification of cellular expression by nonlinear stochastic embedding (ACCENSE) were used. Most of the changes induced by cryopreservation were apoptotic, with increase in caspase 3 activation ($P < 0.01$), PS translocation to the outer membrane ($P < 0.001$), loss of mitochondrial membrane potential ($P < 0.05$), and increase in intracellular Na+ ($P < 0.01$). Average values of markers of capacitative changes were not affected by cryopreservation; however, the analysis of the phenotype of individual spermatozoa using computational flow cytometry revealed the presence of subpopulations of spermatozoa experiencing capacitative changes. For the first time advanced computational techniques were applied to the analysis of spermatozoa, and these techniques were able to disclose relevant information of the ejaculate that remained hidden using conventional flow cytometry.

Reproduction (2017) 153 293–304

Introduction

In spite of equine male gametes trade constituting a major component of the equine industry, cryopreservation of stallion spermatozoa, as an indispensable tool for the commerce of frozen-thawed semen doses, still has numerous drawbacks (Peña et al. 2011). To obtain acceptable fertility, intensive management of the mares is required to perform AI close enough to ovulation as the surviving spermatozoa have reduced lifespan in the mare’s reproductive tract. This intensive management increases the costs related to insemination precluding a wider use of frozen-thawed semen. The reduced lifespan of the thawed sperm has been attributed to either changes similar to capacitation (Cormier et al. 1997, Pons-Rejraji et al. 2009) or changes that resemble apoptosis (Ortega-Ferrusola et al. 2008), or spermtptosis as it has been recently termed (Gallardo Bolanos et al. 2014, Peña et al. 2016b). Capacitation is characterized by changes in sperm membrane fluidity, overall increase in tyrosine phosphorylation, and decrease in intracellular sodium (Escoffier et al. 2012), while spermtptosis implies caspase activation, loss of mitochondrial membrane potential, and transposition of phosphatidylserine (PS) to the outer leaflet of the membrane (Ortega-Ferrusola et al. 2008, Ortega Ferrusola et al. 2009, Gallardo Bolanos et al. 2014). However, there is a lack of detailed studies analyzing both types of changes simultaneously. We aimed to focus our study on the surviving population of spermatozoa, to identify how characteristics related to apoptosis or capacitation change during this process in the same ejaculates following a split sample design. Recent developments in flow cytometry applied to sperm analysis are increasing the number of parameters that can be assessed simultaneously in a single assay (Gallardo Bolanos et al. 2014, Peña et al. 2015, Peña 2016a), this therefore demands the
interpretation of the results due to the multidimensional nature of the data gathered, compared with the traditional bi-dimensional dot plots or uni-dimensional histograms. Moreover, the heterogeneous nature of the ejaculate is not disclosed in classical flow cytometry analysis based in manual gating. By using t-SNE data generated by multicolor flow cytometry can be represented in a two-dimensional (2D) plot, similar to biaxial plots, in which differences account for all the markers (Grimes et al. 2013). ACCENSE (Automatic Classification of Cellular Expression by Nonlinear Stochastic Embedding) is a tool that computes a 2D nonlinear distillation of the raw data, and automatically stratifies cells into phenotypic subpopulations based on their distribution of markers (Shekhar et al. 2014), avoiding the pitfalls derived from manual gating and classification of cells.

In view of these facts, we studied, in the same ejaculates following a split sample design, changes occurring after cryopreservation potentially related to capacitation (such as increase in tyrosine phosphorylation, increase in membrane fluidity, production of reactive oxygen species (ROS), and changes in intracellular Na+) and potentially related to apoptosis (such as mitochondrial membrane potential, activation of caspase 3, and transposition of PS). Moreover, we applied traditional and computational flow cytometry analysis to shed light on the changes that cryopreservation induces in the surviving spermatozoa population, and to disclose changes in the sperm subpopulation structure at the single cell level. We hypothesize that computational flow cytometry may improve our understanding of the changes induced by cryopreservation at the single cell level, disclosing changes in the sperm subpopulation structure of the ejaculates.

Materials and methods
Reagents and media
Merocyanine 540 (Excitation: 488 nm; Emission: 560 nm) (Ref: M24571); YO-PRO-1 (Excitation: 491 nm; Emission: 509 nm) (Ref: Y3603); Hoechst 33342 (Excitation: 350 nm; Emission: 461 nm) (Ref: H3570); 5,5′,6,6′-tetrachloro-1′,3′,3′-tetraethyl benzimidazoyl carbocyanine iodine (JC-1) (Excitation: 488 nm; Emission: 530 nm, monomer form) (Excitation: 561 nm; Emission: 591 nm, aggregate form) (Ref: T3168); Annexin V Pacific Blue™ conjugate (Excitation: 490 nm; Emission: 530 nm) (Ref: A35122); CellROX Deep Red Reagent (Excitation: 488 nm; Emission: 530 nm) (Ref: C10422); Cell Event Caspase-3/7 Green Detection Reagent (Excitation: 502 nm; Emission: 530 nm) (Ref: C10423); ethidium homodimer (Excitation: 528 nm; Emission: 617 nm) (Ref: E1169); LIVE/DEAD® Fixable Violet Dead Cell Stain Kit (Excitation: 405 nm; Emission: 451 nm) (Ref: L34955) and Sodium Green cell permeant Indicator (Ref S-6901) (Excitation: 488 nm; Emission: 532 nm) were purchased from ThermoFisher Scientific (Molecular Probes). Phospho-Tyrosine Mouse mAb (P-Tyr-100) (PE conjugate) (Ref: 14967) was acquired from Cell Signalling Technology.

Semen collection and processing
Three ejaculates per horse were collected from 7 fertile stallions, which were maintained according to institutional and European regulations (Law 6/2013 June 11th and European Directive 2010/63/EU). The samples were obtained on a regular basis (three collections/week) during the 2016 breeding season. The ejaculates were obtained using a pre-warmed (45°C), lubricated Missouri model artificial vagina with an inline filter to eliminate the gel fraction. The semen was immediately transported to the laboratory for evaluation and processing. The ejaculates were separated in two aliquots, diluted 1:2 in INRA96 (IMV, L’Aigle, France), and centrifuged at 600 g for 10 min at room temperature. One of the aliquots was further extended in INRA 96 after centrifugation to obtain a final concentration of 100 × 10⁶ spermatozoa/mL and was kept at room temperature (22°C) during 1 h for analysis as raw sperm controls. The other aliquot was diluted in the freezing medium Cáceres (University of Extremadura, Cáceres, Spain) containing 2% egg yolk, 1% glycerol, and 4% dimethylformamide to 100 × 10⁶ spermatozoa/mL. After loading the extended semen into 0.5-mL straws (IMV, L’Aigle, France), the straws were ultrasonically sealed with UltraSeal 21® (Miniture of America MOFA, Verona, WI, USA) and immediately placed in an IceCube 14S (SY-LAB, Neupurkersdorf, Austria) programmable freezer. The following freezing curve was used. Straws were kept for 15 min at 20°C, and they were then slowly cooled from 20°C to 5°C at a cooling rate of 0.1°C/min. Thereafter the freezing rate was increased to −40°C/min from 5°C to −140°C. The straws were then plunged into liquid nitrogen and stored until analysis. For the analysis, two straws per stallion and freezing operation were thawed in a water bath at 37°C for at least 30 s and diluted in pre-warmed INRA 96 extender to a final concentration of 50 × 10⁶ spermatozoa/mL. All analyses were conducted immediately after thawing.

Flow cytometry
Flow cytometry analyses were conducted using two equipments: a MACSQuant Analyzer 10 (Miltenyi Biotech) flow cytometer equipped with three lasers emitting at 405, 488, and 635 nm and 10 photomultiplier tubes (PMTs): V1 (excitation 405 nm, emission 450/50 nm), V2 (excitation 405 nm, emission 525/50 nm), B1 (excitation 488 nm, emission 525/50 nm), B2 (excitation 488 nm, emission 585/40 nm), B3 (excitation 488 nm, emission 655–730 nm (655LP+split 730)), B4 (excitation 488 nm, emission 750 LP), R1 (excitation 635 nm, emission 655–730 nm (655LP+split 730)), and R2 (excitation 635 nm, emission filter 750 LP); and a MACSQuant® VYB (Miltenyi Biotech) flow cytometer equipped with yellow, violet, and blue lasers (561, 405, 488 nm) with 10 optical channels for multiparameter flow cytometry. The system was controlled using MACS Quantify software. The quadrants or regions used to quantify the frequency of each sperm subpopulation depended on the particular assay. Forward and sideways light scatter was recorded for a total of 50,000 events.
per sample. Gating the sperm population after Hoechst 33342 staining eliminated non-sperm events. The instruments were calibrated daily using specific calibration beads provided by the manufacturer, and a compensation overlap was performed before each experiment. The data were analyzed using FlowJo V 10.2 Software (Ashland, OR, USA). Unstained, single stained, isotype, and Fluorescence Minus One (FMO) controls were used to determine compensations and positive and negative events, as well as to set regions of interest as described in previous publications from our laboratory (Pena et al. 2003, Gallardo Bolanos et al. 2014) (data not shown).

**Simultaneous flow cytometric assessment of caspases 3 and 7 activity, viability, and oxidative stress (ROS)**

CellEvent Caspase-3/7 Green Detection Reagent is a fluorogenic substrate for activated caspases 3 and 7. The reagent consists of a four-amino-acid peptide (DEVD) conjugated to a nucleic acid-binding dye. This cell permeant substrate is intrinsically non-fluorescent because the DEVD peptide inhibits the ability of the dye to bind to DNA. After activation of caspase 3 and caspase 7 in apoptotic cells, the DEVD peptide is cleaved, enabling the dye to bind to DNA and produce a bright, fluorogenic response with an absorption/emission maxima of 502/530 nm. Spermatozoa exhibiting oxidative stress emit fluorescence in the far-red spectrum, whereas Hoechst 33342-positive sperm emit blue fluorescence. The semen samples were diluted in PBS for staining to a final concentration 5 × 10³ spermatozoa/mL. Then, cells were stained with 1 µL of CellROX (5 µM), 1 µL CellEvent Caspase-3/7 Green Detection Reagent (2 mM), and 0.3 µL of Hoechst 33342 (0.5 µM). After thorough mixing, the sperm suspension was incubated at room temperature in the dark for 25 min, washed in PBS, and loaded with 0.3 µL ethidium homodimer (1.167 mM in DMSO) to each sample. After incubation for a further 5 min, the samples were immediately run on the flow cytometer. The controls consisted of unstained and single stained controls to properly set gates and compensations. The positive controls for oxidation and caspase activation were samples supplemented with 800 µM SO₄²⁻, 1.7 M H₂O₂, and 2.7 µM JC-1 (Molecular Probes) to induce the Fenton reaction (Gallardo Bolanos et al. 2014). The debris were gated out based on the DNA content of the events after Hoechst 33342 staining.

**Simultaneous assessment of mitochondrial activity and PS translocation**

The mitochondrial potential was assessed by 5,5′,6,6′-tetrachloro-1,1′,3,3′ tetraethyl benzimidazolyl carbocyanine iodine (JC-1). This probe forms multimeric aggregates in mitochondria with high membrane potential (active mitochondria) and these aggregates emit in the high orange wavelength of 595 nm when excited at 561 nm (yellow laser). In mitochondria with low membrane potential (inactive mitochondria), JC-1 forms monomers that emit in the green wavelength (530 nm) when excited at 488 nm. Apoptosis-like changes were detected in spermatozoa with the use of Annexin V, Pacific Blue™ conjugate (Dead Cell Apoptosis Kit, Molecular Probes), which detects the translocation of PS from the inner to the outer leaflet of the plasma membrane associated with membrane changes related to sperm processing. Apoptotic cells show violet fluorescence (455 nm) when excited with the violet laser (405 nm). Both stains were combined in a multiparametric test and evaluated by FC. A final concentration of 5 × 10⁴ spermatozoa/mL was obtained by adding 10 µL of diluted spermatozoa to 990 µL of PBS. The samples were loaded with 0.5 µL JC-1 (0.3 µM) and incubated at 37°C for 40 min. The samples were washed in PBS by a short centrifugation spin for 12 sec and suspended in 100 µL of Annexin binding buffer (10 mM HEPES, 140 mM NaCl, 2.5 mM CaCl₂, pH 7.4). To 100 µL of sample per assay, 5 µL of Annexin V, Pacific Blue™ were added. After 15 min of incubation in the dark at room temperature, 400 µL of 1× Annexin binding buffer were added. Cytometry analysis was then performed using a MACSQuant® VYB (Miltenyi Biotech) flow cytometer with the yellow and violet lasers (561, 405 nm).

**Evaluation of fluidity (lipid disorder) of the sperm membrane**

Fluidity of sperm membranes was assessed with Merocyanine 540 (488–560 nm) (blue laser, channel B2) (Molecular Probes Europe) and membrane changes were detected with YO-PRO-1 (491–509) (blue laser, channel B1) (Molecular Probes Europe). Hoechst 33342 (350–461 nm) (violet laser, channel V1) was also incorporated to gate out debris. FC analyses were carried out with a MACSQuant Analyzer 10 (Miltenyi Biotech, Pozuelo de Alarcón, Madrid, Spain). A suspension of 5 × 10⁴ spermatozoa was obtained by adding 10 µL of diluted spermatozoa to 990 µL of PBS. The samples were loaded with Merocyanine 540 at a final concentration of 2.7 µM, YO-PRO-1 25 nM, and 0.3 µM Hoechst 33342 and incubated in the dark for 30 min. The cells were washed in PBS by a short centrifugation spin for 12 sec, before reading in the FC. While Merocyanine 540 attaches to the sperm membrane and fluoresces in orange proportionally to the grade of membrane disorder present, YO-PRO-1 is a high-affinity nucleic acid stain that penetrates cells with compromised plasma membrane, fluorescing in green when is excited at 488 nm.

**Assessment of tyrosine phosphorylation**

The anti-phospho-Tyrosine Mouse mAb (P-Tyr-100) (PE conjugate) was used to detect the phosphorylation of tyrosine. The samples (1 mL) containing 10 × 10⁴ spermatozoa/mL in PBS were stained with 1 µL of LIVE/DEAD violet Fixable Dead Cell Stain Kit Solution. After thorough mixing, the samples were incubated at room temperature (22°C) for 30 min in the dark. Then, the spermatozoa were washed with PBS (1×) and fixed with 900 µL of 4% paraformaldehyde and incubated for 10 min at 37°C. Afterwards, samples were incubated for 1 min on ice. The aliquots were washed by centrifugation twice. For the permeabilization, pre-chilled cells were slowly resuspended in ice-cold 100% methanol to a final concentration of 90% methanol. The samples were incubated 30 min on ice. Then, the spermatozoa were washed in incubation buffer (0.5 g bovine serum albumin (BSA) in 100 mL 1× PBS) twice. The cells were loaded with 2 µL/mL Phospho-Tyrosine Mouse mAb (P-Tyr-100) (PE conjugate) (50 µg/mL # 14967), and incubated for 30 min at room temperature (22°C) in the dark.
after thorough mixing. Then, the spermatozoa were washed in 500µL PBS. Finally, the samples were washed again in PBS before reading in the flow cytometer.

**Determination of intracellular Na+**

The amount of intracellular Na+ in stallion spermatozoa was determined flow cytometrically after loading the cells with Sodium Green Permeant (5µM), and incubated at r.t. 30'. Finally, samples were washed in PBS and dead spermatozoa were excluded from the analysis after ethidium homodimer staining (0.35µM 5' at room temperature).

**Computational flow cytometry (t-SNE analysis and ACCENSE)**

Flow cytometry data are usually analyzed using a series of 2D plots and manual gating, however, the increase in the number of parameters measured increase the number of 2D plots to display for every marker combination. For example, a combination of four colors will require 30 2D plots. To overcome these problems computational methods to automatically identify populations in multidimensional flow cytometry data have been developed (Mair et al. 2016). Using FlowJo v 10.2 and ACCENSE software available at http://www.cellaccense.com, compensated data of each multiparametric assay described in the Materials and methods section were exported as FCS files from the flow cytometer, and loaded in FlowJo and ACCENSE for computational analysis. Data were concatenated and single cell events analyzed. Flow cytometry data were analyzed using nonlinear dimensionally reduction techniques (t-SNE) and ACCENSE. The Barnes–Hut implementation of t-SNE was used for low-dimensional embedding to perform dimensional reduction of cytometry data, classification of cells was based in K means techniques with the significance level set at $P=0.0001$, and data of all subpopulations generated were exported as FCS files. ACCENSE identifies clusters within multidimensional data without losing single cell resolution (Shekhar et al. 2014, Chester & Maeker 2015), allowing automatic gating of cells.

**Statistical analysis**

Three ejaculates were collected from each of the 7 individual stallions. All experiments were repeated at least three times with independent samples (three separate ejaculates from each of the seven stallions). The normality of the data was assessed using the Kolmogorov–Smirnov test. Since the data showed equivalence of variance, the results were analyzed by ANOVA followed by a Tukey post hoc test to perform pair-wise comparisons (SPSS 19.0 for Mac). Differences were considered significant when $P<0.05$, and are indicated as *$P<0.05$ and **$P<0.01$. Results are displayed as mean±S.E.M.

**Results**

**Changes induced by cryopreservation in caspase 3 activation, viability, and ROS production**

Representative cytograms and gating strategies of the assay are given in Figure 1. The conventional flow cytometry analysis applied made evident that cryopreservation induced an increase in the percentage of spermatozoa expressing active caspases 3 and 7 (Fig. 2A). Paradoxically, ROS production was reduced in thawed samples (Fig. 2B and C). However, this reduction was not observed in a population of caspase 3-positive spermatozoa (Fig. 2B). The embedding (t-SNE) and clustering steps in ACCENSE revealed a complex structure of the ejaculate identifying more than 100 sperm subpopulations both in raw sperm and after freezing and thawing (Fig 3A, B, C, and D, E, F). Visualization of the structure of the ejaculate allowed identifying easily changes in the expression of caspase 3 as a result of the cryopreservation (Fig. 3B and E). The algorithm also gave FCS files of every subpopulation in the complex ejaculate structure: Figure 3G and H shows representative overlays of characteristic subpopulations in fresh sperm (in general, low caspase 3 expression and high superoxide production) and Figure 3I and J shows representative overlays in frozen and thawed samples characterized by low superoxide production and high caspase 3 activity. Reduction of multidimensional data to 2D plots using t-SNE in FlowJo to visualize changes in the sperm subpopulation structure of viable spermatozoa after cryopreservation (Fig. 4) revealed interesting changes; virtually all spermatozoa experienced changes after cryopreservation, with increases in the caspase activation, reduction of live spermatozoa, and changes in the position of sperm subpopulations in the 2D structure as the most relevant.

**Changes in mitochondrial membrane potential ($\Delta \Psi m$) and PS transposition to the outer leaflet of the membrane (Annexin assay) after cryopreservation**

Cryopreservation induced a massive transposition of PS to the outer leaflet of the sperm membrane (Fig. 5I), and also reduced the membrane potential ($\Psi m$) of sperm mitochondria (Fig. 5I), with hierarchical gating showing that this effect occurred principally in Annexin + sperm mitochondria (Fig. 5G and H). When ACCENSE analysis was applied based in these markers, once again a complex subpopulation structure of the ejaculate was evident, with more than 100 distinct subpopulations revealed both in raw semen and after freezing and thawing (Fig. 5A, C, D, and F). Visualization of the t-SNE maps after ACCENSE gating and classification allowed identification of phenotypic changes in the presence of PS in the outer membrane (Fig. 6A and D) and in the mitochondrial membrane potential (Fig. 6B, E high $\Psi m$ and C, F low $\Psi m$). Subpopulations in fresh sperm with PS transposition were evident in raw sperm (Fig. 5B) that were not obvious in the conventional analysis, expression of PS in the outer membrane increased after cryopreservation (Fig. 6D). ACCENSE analysis also revealed changes in mitochondria after cryopreservation with a general reduction in mitochondrial membrane...
potential, although subpopulations of spermatozoa with high $\Psi m$ remained (Fig. 6B and E). ACCENSE analysis provided FCS files of all the subpopulations originated; representative overlay cytograms of subpopulations in fresh and frozen-thawed samples are given in Figure 6G and H subpopulations in raw sperm were characterized by high $\Psi m$, and low PS externalization (H) while subpopulations in thawed samples show low $\Psi m$ and PS externalization (I).

**Evaluation of changes in membrane fluidity induced by cryopreservation**

Cryopreservation reduced the fluidity of the sperm membranes (Fig. 7A, D, and E), however, also induced significant increases in Yo-Pro-1 uptake indicating increased membrane permeability and compromised membranes (Fig. 7B, C, D, and E).

The t-SNE analysis and visualization of the subpopulation structure revealed that cryopreservation induced dramatic changes in the structure of the ejaculate; downsampling and concatenation revealed that both raw sperm and frozen-thawed samples conserved two subpopulations of high and low merocyanine fluorescence (Fig. 8B and D). However, it was evident that in thawed samples the difference among these subpopulations was higher (Fig. 8D) than in raw sperm (Fig. 8B). The analysis of the phenotype of the spermatozoa using ACCENSE revealed that true subpopulations of high mero fluorescence were more common in fresh sperm than in thawed samples, where most of the spermatozoa showing high mero fluorescence also showed high Yo-Pro-1 uptake (Fig. 8A, B, D, and E). Overlay of sperm phenotypes in the t-SNE map revealed that the major change after freezing and thawing was an increase in Yo-Pro-1 uptake (green population in Figure 9C and F).

** Changes in tyrosine phosphorylation induced by freezing and thawing**

When conventional gating was applied, no major changes were observed in the intensity of the fluorescence of the phosphorylated proteins (Fig. 10C), the gating strategy to identify tyrosine phosphorylation
is depicted in Figure 10D, E, F, and G. But when computational cytometry techniques were applied to the analysis, the t-SNE map and automatic ACCENSE classification of spermatozoa revealed that overall intensity of phosphorylation in tyrosine of sperm proteins was higher in fresh samples, and was reduced after cryopreservation, with the exception of a subpopulation of spermatozoa showing high intensity in both raw spermatozoa and thawed samples (Fig. 10A and B).

Changes in intracellular Na+ after cryopreservation

We used flow cytometry to investigate how intracellular Na+ changes during cryopreservation; the population of dead spermatozoa was gated out from the analysis after PI staining (Fig. 11B). Cryopreservation resulted in an increase in the content of intracellular Na+ in the spermatozoa surviving cryopreservation (Fig. 11A and C).

Discussion

In this study we evaluated and compared changes induced by cryopreservation that may relate to apoptosis (spermptosis) or to capacitation (cryocapacitation) in the same ejaculates following a split sample design. Moreover, for the first time we applied to sperm analysis techniques of computational flow cytometry allowing automatic gating and the identification of phenotypically distinct sperm subpopulations after flow cytometry analysis. The heterogeneous nature of the mammalian ejaculate has been recognized; different subpopulations of spermatozoa can be distinguished based on, for example, morphometric and kinetic analysis (Aparicio et al. 2005, Nunez-Martinez et al. 2006, 2007, Pena et al. 2006, Ortega-Ferrusola et al. 2009). This complex and heterogeneous nature has been...
identified in many different biological systems (Newman & Weissman 2006, Newman et al. 2006) and its importance also stressed in different aspects of medicine (Sigal et al. 2006, Cohen et al. 2008, Saadatpour et al. 2015). The heterogeneous nature of the spermatozoa has led to the consensus of the limited value of the report of average values in the sperm analysis (Newton 1998). Here, for the first time we have disclosed sperm subpopulations using computational flow cytometry approaches, and these techniques are powerful tools to identify sperm subpopulations in the ejaculate. Most dramatic changes in the spermatozoa surviving cryopreservation were related to apoptotic changes, with intense caspase 3 activation as the most evident change of all the parameters studied using both conventional and computational flow cytometry. This finding confirms...

**Figure 5** ACCENSE applied to high-dimensional multicolor cytometry data. (A and D) The 2D t-SNE maps of stallion spermatozoa after multicolor staining (Anexin-V and JC-1) in fresh (A) and frozen-thawed spermatozoa (D). Each point represents a cell derived from downsampling from the original dataset. (B and E) Density map representing the intensity of PS externalization in raw (B) and FT samples (E) of stallion spermatozoa, PS externalization increased after cryopreservation, although subpopulations with increased PS externalization were already evident in raw semen (see heat map bar at the right of each t-SNE map). (C–F) Composite map depicting the complex structure of the stallion ejaculate; circles represent centers of phenotypically distinct subpopulations ($P<0.001$). (G) Representative dot plots of the JC-1 assay. (H) Representative overlay histograms of PS expression in raw semen (red) and thawed spermatozoa (blue). (I) Changes in the percentage of spermatozoa showing PS transposition and (J) changes in mitochondrial membrane potential induced by cryopreservation *$P<0.05$, **$P<0.001$.

**Figure 6** ACCENSE-derived t-SNE maps depicting channels for each phenotype marker in stallion spermatozoa in raw and thawed stallion spermatozoa. (A and D) Modifications in the externalization of PS after cryopreservation. (B–E) Changes in the mitochondria showing high $\psi_{m}$. (C–F) Changes in the mitochondria showing low $\psi_{m}$. (G and H) 20 representative subpopulations of spermatozoa in raw and frozen-thawed stallion spermatozoa, subpopulations in raw sperm were characterized by low PS and high $\psi_{m}$, while subpopulations in thawed spermatozoa were characterized by high PS externalization and low $\psi_{m}$.
previous reports indicating that cryopreservation activates apoptotic phenomena (Pena et al. 2009, Ortega Ferrusola et al. 2010, Gallardo Bolanos et al. 2012, Caselles et al. 2014, Gallardo Bolanos et al. 2014, Kopeika et al. 2015, Pena et al. 2015, Munoz et al. 2016). Other features typical of apoptosis were also evident in our study including PS transposition and decreases in the mitochondrial membrane potential. An apparently paradoxical finding occurred, cryopreservation reduced ROS production; however, similar findings have been recently reported (Gibb et al. 2014, Yeste et al. 2015, Munoz et al. 2016). This is explained based in two factors: on one hand, the probe used in our study is specially sensitive to the superoxide anion (O$_2^•$) (Plaza Davila et al. 2015) and, on the other hand, mitochondria are the main source of ROS in the spermatozoa (Plaza Davila et al. 2015), so it is not surprising that reduced mitochondrial activity in thawed sperm (Pena et al. 2015) leads to decreased production of O$_2^•$. Markers of capacitation were also evaluated, including the fluidity of the sperm membranes and tyrosine phosphorylation of sperm proteins. Conventional flow cytometry revealed a reduction in the percentage of spermatozoa showing low membrane fluidity in cryopreserved samples; however, increases in membrane fluidity were also accompanied by increased Yo-Pro-1 uptake, suggesting early damage in the membrane instead of capacitation. The t-SNE analysis revealed not only a subpopulation of spermatozoa with increased membrane fluidity, but also increased Yo-Pro-1 uptake after thawing. Since an overall increase in tyrosine phosphorylation of sperm proteins is considered a hallmark of capacitation, this was investigated in fresh and thawed samples. There was not a significant increase in tyrosine phosphorylation after thawing. On the other hand, ACCENSE analysis revealed that increased tyrosine phosphorylation occurred only in a subpopulation of spermatozoa. Finally, since it has been shown that capacitation is associated with reduced intracellular Na$^+$ (Escoffier et al. 2012), modifications in the sperm concentration of this ion were studied. We observed an increase in intracellular Na$^+$ after cryopreservation; disruption of the Na+/K+ pump has been linked to cell shrinkage associated with apoptosis in many cell systems (Bortner & Cidlowski 2003, Iurinskaia et al. 2010, Sapia et al. 2010, Orlov et al. 2013, Honisch et al. 2014). In spermatozoa, reduced intracellular ATP associated with mitochondrial dysfunction (Plaza Davila et al. 2015, Davila et al. 2016) may explain increased intracellular Na$^+$ in thawed samples.

Markers related with apoptosis were more evident in thawed samples; however, markers related to capacitation appeared only in specific subpopulations of thawed spermatozoa. Capacitation only occurs in a small percentage of cells (Bahat et al. 2003, Eisenbach 2003, Eisenbach & Giojalas 2006, Spiegel et al. 2006), this is in agreement with our findings. Capacitation is a
redox-regulated process, with increased ROS activating a soluble adenylyl cyclase that stimulates cAMP and PKA; the latter activates SRC kinases and suppresses an inhibitor of SRC. SRC phosphorylates and inactivates a protein phosphatase permitting a dramatic upregulation of tyrosine phosphorylation that characterizes capacitation (Salicioni et al. 2007, Buffone et al. 2014, Aitken et al. 2015, Alvau et al. 2016). Increased production of H₂O₂ leads to inhibition of tyrosine phosphatase activity, contributing also to increased tyrosine phosphorylation. Thus is not surprising that under conditions of reduced production of mitochondrial ROS, tyrosine phosphorylation is reduced, as seen in our experiments. It has been proposed that capacitation and apoptotic changes share a common

Figure 9 ACCENSE-derived t-SNE maps depicting channels for each phenotype marker in stallion spermatozoa in raw (A and D) and thawed (B and E) stallion spermatozoa. The intensity of Merocyanine 540 fluorescence and of the Yo-Pro1 uptake across the whole population of spermatozoa is depicted in a heat map. Comparing Merocyanine 540 fluorescence and Yo-Pro1 uptake allows to determine that in thawed samples the highest uptake of Yo-Pro1 corresponds also to high mero fluorescence (arrows). (C–F) t-SNE maps showing overlays of distinct sperm subpopulations in raw and frozen-thawed sperm, the most evident changes is an increase in the subpopulation of high Yo-Pro-1 uptake (in green) in thawed samples.

Figure 10 ACCENSE-derived t-SNE maps depicting channels for the intensity of tyrosine phosphorylation in the whole population of spermatozoa after downsampling and dimensional reduction. (A) Raw stallion spermatozoa. (B) Thawed stallion spermatozoa; although overall phosphotyrosine expression is reduced, a subpopulation with high expression appear in FT sperm. (C) Average fluorescence intensity in raw sperm and in frozen-thawed samples. (D–E) Gating strategy; D: fluorescence minus one (FMO) control and live a dead spermatozoa. (E–F) The region of live sperm phosphotyrosine-positive is gated to further analysis. (G) Histogram showing intensity of the fluorescence of phosphotyrosine-positive events.

Figure 11 Changes in the content of intracellular Na⁺ after freezing and thawing. (A) Relative fluorescence units in raw and thawed samples **P<0.05. (B) Representative cytogram of the assay, dead spermatozoa are excluded after ethidium homodimer staining, Na⁺ concentration is estimated with sodium green as indicated in materials and methods. (C) Overlay histograms representing intracellular Na⁺ in raw (red) and frozen-thawed spermatozoa (blue).
trigger in spermatozoa, oxidative stress, being opposite ends of a phenomena with a common triggering event (Aitken et al. 2015). Previously, we demonstrated that ROS imbalance is the key phenomena explaining apoptotic changes induced by cryopreservation (Ortega Ferrusola et al. 2009, Pena et al. 2015, Munoz et al. 2016). ROS imbalance leads to increased production of 4-hydroxynonenal (4-HNE) and caspase activation; however, traditional markers of oxidative stress can be reduced after cryopreservation (Martin Munoz et al. 2015). Our findings match with the hypothesis proposed by Aitken et al. (Aitken et al. 2015); cryopreservation leads to a transient burst of mitochondrial ROS, which initially induces capacitation in a subpopulation of spermatozoa; however, later, mitochondrial osmotic stress leads to reduced mitochondrial function with reduced ATP and stops mitochondrial ROS production. As the production of 4-HNE depends only on the initial oxidative stress and a continuous source of ω-6 fatty acids, which are very abundant in the membranes of stallion spermatozoa (Garcia et al. 2011, Macias Garcia et al. 2011), 4-HNE levels increase in spite of reduced mitochondrial ROS. Increased 4-HNE activates caspase 3 leading to sperm death in the presence of reduced ROS levels, explaining this apparent paradox.

Of special interest was the ability of the ACCENSE analysis to disclose hidden subpopulations in the ejaculate. In our study, we were able to identify subpopulations with high levels of PS externalization or groups of spermatozoa showing intense phosphorylation in tyrosine. These techniques have been used with success to identify rare subpopulations in CD8+ T cells in mouse (Shekhar et al. 2014) and similar techniques used to disclose intra-tumor heterogeneity in acute myeloid leukemia, linked to prognosis (Levine et al. 2015). Our results show that these techniques can be applied to the study of sperm biology, being able to provide relevant new information, thanks to the automatic gating and classification of spermatozoa into specific subpopulations, which are not distinguishable on biaxial plots of conventional markers.

In summary, changes induced by cryopreservation are apoptotic, with reduced mitochondrial functionality and caspase 3 activation and PS transposition as the more obvious, although a subpopulation of spermatozoa may experience capacitative changes. Moreover, we introduced for the first time computational flow cytometry tools in spermatology as being potent and promising for the study of the spermatozoa biology.

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

Acknowledgements

The authors received financial support for this study from the Ministerio de Economía y Competitividad-FEDER, Madrid, Spain, grant AGL2013-43211-R, Junta de Extremadura-FEDER (GR 15029). PMM is supported by a pre-doctoral grant from the Ministerio de Educación, Cultura y Deporte, Madrid Spain FPU13/03991. COF is supported by a post-doctoral grant from the Ministerio de Economía y Competitividad “Juan de la Cierva” IJCI-2014-21671.

References


Computational flow cytometry in sperm...


Received 3 October 2016
First decision 24 October 2016
Revised manuscript received 2 November 2016
Accepted 12 December 2016