LOW TAURINE CONCENTRATIONS POSSITIVELY AFFECT RABBIT SPERMATOZOA PROPERTIES IN LATER TIME INTERVALS

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INTRODUCTION

The artificial insemination is a routine technique in rabbit breeding. Insemination must take place within 24-48 hours after semen collection due to decreased viability induced by production of free radicals (Carlucio et al., 2004). Reactive oxygen species (ROS) have a beneficial role in spermatozoa functioning enabling spermatozoa to maturate, capacitate, hyperactivate, undergo the acrosome reaction and to fuse with oocyte. However, long time storage of ejaculated semen results in further production of ROS and its accumulation (Kothari et al., 2010). The excessive free radicals production causes the decrease in semen quality resulting from oxidative damage of biomolecules. Large amount of polysaturated fatty acids contained in plasma membrane along with low concentration of scavenging enzymes in cytoplasm make spermatozoa vulnerable to ROS (Lucio et al., 2016; Tvrda et al., 2016). The survival of the spermatozoa depends not only on the storage temperature but also on the interaction of temperature and the composition of storage medium (Carlucio et al., 2004). Taurine is identified as a non-permeating spermatozoa cryoprotectant which minimizes the cellular damage (Chhillar et al., 2012). Intracellular taurine is present in cells at high amounts and its role is determined by the cell type. The use of taurine in reproduction comes out of ability to sustain the spermatozoa motility and to take part in spermatozoa capacitation (Tirpak et al., 2015). The addition of anti-oxidants, especially those which are contained in epididymal and oviductal fluids (e.g., taurine), to semen extenders protects the spermatozoa against the ROS generated during freezing and thawing process (Bucak and Tekin, 2007; Sariözkan et al., 2009). Our study was aimed on the improvement of spermatozoa properties during long time storage. Previous studies on the influence of taurine on bovine (Chen et al., 1993; Chhillar et al., 2012; Sariözkan et al., 2009), ram (Bucak and Tekin, 2007), boar (Jang et al., 2006), goat (Bucak et al., 2009), dog (Michael et al., 2007) and fish (Martinez-Páramo et al., 2013) spermatozoa reported enhanced parameters associated with successful fertilization. The aim of this study was to examine the effects of taurine on selected rabbit spermatozoa motility and viability parameters during in vitro incubation at 39°C.

MATERIAL AND METHODS

Semen collection and processing

Ejaculates were obtained from sexually mature New Zealand White rabbits (n=9), bred at experimental farm of Animal Production Research Centre in Lužianky. Semen collection was accomplished by the use of pre-warmed artificial vagina following the sexual stimulation by a rabbit doe. Consequenly, semen was stored at 5°C. Fresh ejaculates (30 µl) were diluted in ratio 1:7 with physiological solution (NaCl 0.9% Braun, Braun Melsungen AG, Germany) for the control group (C). Experimental samples were prepared according to the same dilution rate using three concentrations of taurine (Taurine≥99%, Sigma Aldrich, Bratislava, Slovakia) dissolved in the physiological solution. Taurine solutions TAU3 (3.125 nM), TAU6 (6.250 mM) and TAU12 (12.5 mM). Ejaculates of nine adult rabbits were subjected to this study and observed at time intervals 0, 90, 180 and 270 minutes while cultivated at 39°C. For detection of viability the MTT test was used and motility parameters were carried out using CASA analysis. Experimental samples were compared against the control. Assessment of the mitochondrial toxicity revealed that the addition of taurine ensured the higher viability in supplemented samples however with no statistical significance. Monitored motility parameters (motility, progressive motility and velocity curved line of spermatozoa) showed that taurine treated spermatozoa have lower activity at the beginning of the study but along with increasing time the positive effect of taurine was induced. Significant (P<0.05) values were observed in velocity curved line in time intervals 0 (TAU3 – positive effect: 110.6±26.85 μm.s-1) and 180 (TAU6 – positive effect: 105.8±28.36 μm.s-1). Results indicate that taurine antioxidant properties maintain the spermatozoa viability and activity. In spite of the fact that taurine supplemented spermatozoa had lower activity in incentive time intervals, taurine may find its use as additive with ROS protective properties during long-term storage.

Keywords: taurine, CASA, spermatozoa, rabbit, MTT
The data were expressed in percentage – comparing to metabolic activity of sperm cells in the control group (Jambor et al., 2017; Tvrdá et al., 2016).

**Motility analyses**

Semen analyses were performed using the Computer assisted semen analysis (CASA) method with SpermVision software (Minitube, Tiefenbach, Germany) and the microscope Olympus BX 51 (Olympus, Japan). Semen samples were placed into Makler counting chamber (10 μm, Sfi-Medical Instruments, Germany). Measurements of spermatozoa motility were carried out at four time periods (0, 90, 180, 270 minutes). For the duration of the experiment, tested samples were stored in incubator at 39°C. The following spermatozoa characteristics were assessed: motility (MOT), progressive motility (PRO), velocity curve lined (VCL). Every single output of the CASA system is the result of examinations of 7 different areas of Makler Counting Chamber (Tírpák et al., 2016; Slanina et al., 2015).

**Statistical analyses**

For the comparison of the CASA and MTT results in certain time intervals with the focus on effect of additives on spermatozoa, ANOVA and Dunnett’s comparative test were applied using GraphPad Prism 5 (GraphPad Software Inc., USA). All statistical tests were carried out at levels of significance at P<0.05, P<0.01 and P<0.001 and results were interpreted as means and expressed with SD.

**RESULTS AND DISCUSSION**

Present study showed moderately increased spermatozoa viability and motility after extension of fresh rabbit ejaculates with various concentrations of taurine. This trend was manifested mainly in later time intervals while incubated at temperature (39°C) simulating the female reproductive tract temperature. Cell viability was assessed using MTT test which monitors the production of succinate dehydrogenase – mitochondrial enzyme involved in Krebs cycle. The cell viability of taurine treated semen did not statistically differ from the control group. Experimental group TAU3 produced 24.19% more formazan particles than control sample, but was overtaken by absorbance of colorimetric change in group TAU6 - 35.53% higher viability in comparison to the control group. After 270 minutes of incubation tested taurine concentration in sample TAU12 engendered the 11.87% higher cell viability than in the fresh rabbit semen (Figure 1). Asha and Devadasan (2013) suggest that taurine significantly (P<0.001) elevates the activity of the Krebs cycle enzymes after peroxidative intoxication. Assessment of mitochondrial activity, based on the production of the succinate dehydrogenase, indicates that supplementation of rabbit semen with taurine results in higher cell viability of our experimental samples. Bucak and Tekin (2007), who observed the effect of taurine on cooled ram semen, evaluated the spermatozoa viability using the cosin-nigrosin staining technique. The 100 mM taurine solution, which was the most effective, added to conventionally used extender showed the same percentage (78.3%) of intact cells as the control group. These findings correspond with Martínez-Páramo et al., (2013) fluorescent viability evaluation where 1 mM of taurine was applied to an extender of European sea bass semen. However, according to Chhillar et al., (2012) bull semen treated with 50 mM of taurine solution had a significantly (P<0.05) positive effect on viability as compared in its absence. Motility percentage found immediately after dilution was non-significantly lower in experimental samples. Motility percentage varied from 82.58±10.62% to 83.32±9.00% while the C group showed 85.63±7.76% motility. Similar tendency appeared after 90 minutes of cultivation when the semen diluted only with the physiological solution overcame the values of all experimental samples. As shown in Figure 2, 180 minutes of incubation resulted in elevated motility with absence of statistical significance in all samples supplemented with taurine. In the last time interval, semen extended with TAU12 solution reached the highest motility percentage (45.51±15.81%) of all tested samples. Other experimental samples (42.25±18.21%; 41.56±16.24%) were also higher than the control sample (39.27±19.00%).

Subjective assessments of spermatozoa motility conducted by Michael et al., (2007) on dog and by Chhillar et al., (2012) on cattle semen implicated enhanced motility in taurine enriched samples (2.4 mM and 50 mM) with statistical proof (P<0.05). Taurine addition to insemination doses prior to cryopreservation shows no difference in overall motility between 100 mM concentration of additive and conventional sample (Bucak and Tekin, 2007).

More objective computer assisted semen analysis was performed by other authors (Martínez-Páramo et al., 2013; Tírpák et al., 2015) whose results suggest that the most efficient taurine concentrations for the species used in their experiments (fish – 1 mM; bull – 200 mM) had a significantly higher motility after thawing (P<0.05). Even though slightly increased motility was detected in present study, in general it seems that spermatozoa benefit more from taurine protective properties (added in higher doses) than from its antioxidative properties (used in lower concentrations) when stored at simulated female intracorporal temperature. The highest progressive motility (68.00±12.83%), assessed in inceptive time interval, was detected in the group C. Continual proportion between raising taurine concentration and progressive motility increase was determined in all experimental samples of initial measurement. Slighter difference between experimental samples and the control was observed after 90 minutes of cultivation when all values were within the range of 60.69±17.73% and 63.60±18.38%. TAU12 was the most effective taurine solution for the maintenance of the highest possible progressive motility (35.43±24.43%) at 180 minute time interval. Very similar percentages of progressive motility were monitored after 270 minutes of incubation while the values ranged from 21.79±18.09% to 23.33±17.97%.

Tírpák et al., (2015) monitored the effect of five taurine concentrations on post thaw progressive motility of bull spermatozoa. The most effective concentration of taurine (200mM) significantly (P<0.001) leveled up the progressive motility in times 0, 30, 60 and 90 minutes. Negative effect of taurine on spermatozoa velocity was noted in inceptive time and represented by samples TAU12 and TAU3 (P<0.05) in Figure 4. The quickest spermatozoa movement was detected in sample TAU6. After 90 minutes of incubation the bar of control sample was the highest (117.40±27.63 μm.s⁻¹). Experimental samples presented to raising tendency dependent on the concentration of taurine. Positive effect of taurine was proved after 180 minutes of cultivation when the increased spermatozoa velocity was observed in all experimental samples compared to the control. Velocity of spermatozoa extended with TAU6 solution was even statistically significant (P<0.05). Measurements in the latest time interval showed the higher speed of rabbit spermatozoa in experimental samples rather than in control group. Values for the experimental groups TAU3 and TAU6 were fairly the same (76.61±29.06 μm.s⁻¹; 76.23±27.13 μm.s⁻¹) while TAU12 (68.61±26.33 μm.s⁻¹) was moderately higher than C group (68.23±25.05 μm.s⁻¹). Positive effect of taurine on motility has been shown after 180 minutes of incubation. Until then, spermatozoa motility in experimental samples was lower than in the control group.

CASA assessments revealed for Tírpák et al., (2015) that supplementation of bull insemination doses with taurine (50mM, 100mM, 200mM and 400mM) highly (P<0.01 and P<0.001)increases the VCL following the thawing process. The same analyzing system was used by Martínez-Páramo et al., (2013) for the analysis of the effect of taurine on cryopreserved sperm of Sea bass although the significant difference between the control sample and sample enriched with 1mM of taurine was not observed.

Cryopreservation is the biotechnological method successfully applied in domestic animals breeding worldwide; however, rabbit semen processing for cryo storage is still very complicated. The storage of cooled rabbit spermatozoa is the only effective technique for rabbit artificial insemination. Thus, the use of additives which may improve spermatozoa properties may be comprehended as a tool for improvement of spermatozoa motility (Marichal et al., 2017; Stephens et al., 2013). Therefore, synergic effect of beneficial bioactive compounds along with antioxidant activity of taurine might result in improved properties of spermatozoa (Tírpák et al., 2016).
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