Effect of Different Media on Short Term Storage of Leaping Mullet (Liza saliens (Risso, 1810)) Sperm

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A B S T R A C T

The present study aimed to evaluate the activation characteristics of the leaping mullet spermatozoa and diluted with four different media (D1, D2, D3, D4) for 250 h at 4±2°C. The semen was collected from nine wild fish in Homa Lagoon, İzmir, Turkey in July, during reproductive season. In each trial (n = 9), semen samples were collected and mixed to reduce sperm quality variation to obtain the required gametes for the experiment. The semen was activated with natural seawater (salinity of 34, pH 7.9 and 4±2°C) to determine: motility time, motility rate and viability of sperm. Dilution with inactivation media 1:3 increased motility time of diluted semen than control group. At the end of the experiment, no significant difference was found for duration of survival in the D2 and D4 groups. The highest survival rate was found in D2 group.

Introduction

In aquaculture, prolonged cryopreservation of fish semen using extender and cryoprotectants is limited to only 40 to 50% of cellular protection (Watson, 2000). Therefore, storage of fish sperm at temperatures above freezing point and without the addition of cryoprotectants has been successfully used for a long time (McNiven et al., 1993). Short-term preservation techniques are an easy technique that allows the use of sperm to fertilize the eggs at various time intervals and provide more productivity during reproduction time. It can also help improve the genetic quality of the resulting offspring, as well as allowing the examination of sperm properties by facilitating sperm use by more males per female (Ciereszko and Dabrowski, 1994). The cooling of semen to temperatures close to 4°C is one conservation technique, which prolongs temporal viability of undiluted semen from a few hours to days. This can be explained by reduction of spermatozoa’s metabolic activity at temperatures below the physiological range.

The leaping mullet belonging to the Mugilidae family is widespread in a wide area that includes the Pacific and Atlantic oceans. Third world countries, South East Asia, Saudi Arabia, Gulf region countries and Eastern and Southern Mediterranean commercial cultivation of this species from the length of finger are made. The leaping mullet is a species that is often hunted in lagoons. In recent years, the lack of good breeding and uncontrolled hunting has led to a decrease in the population of this species.

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1974; Sukuraman et al., 2003; Magnotti et al., 2016), there are no study for Liza saliens. Therefore, this study is the first record. The present study aimed to evaluate sperm viability in fresh semen and in semen diluted with different media during short-term storage at 0-4°C.

Materials and Methods

Broodstock
The experiments were carried out during the natural spawning period (i.e. July 2018) on sexually mature Leaping mullet (380.2±21.1 g and 31.3±7.3 cm, for weight and length, respectively) originating from Homa Lagoon (İzmir/Turkey). New harvested dead fish (n: 9) were used in the study.

Collection of Gametes
For semen collection, the abdominal and genital areas of each individual were cleaned with deionized water and dried with paper towel. After gentle abdominal pressure, semen was collected with 2 mL syringes and immediately stored in an ice-cold container at 4±2°C protected from light. The extracted semen was collected with a sterile syringe (without needle) and transferred in containers stored at a constant temperature of 4°C to Ege University, samples with urine or blood were discarded.

Evaluation of Fresh Semen

Sperm motility
Sperm motility (mot) was activated according to the procedure previously described by Valdebenito et al., (2017) with some modifications. Briefly, 1μl of semen was diluted in 3μl of 90% sea water (pH 7.0) on a slide and evaluated by subjective ex-amination using a phase-contrast microscope (Olympus C×31) at 40× magnification. Motility time was measured with a stopwatch from the moment of activation until the total stoppage of all spermatozoa. Motility was classified into five categories, determining the percentage of actively moving spermatozoa: 1 = 0 to <5%; 2 = 5–25%; 3 = 25–50%; 4 = 50–75%; 5 = 75–100% (Cosson et al., 2008). Each sample was analysed in triplicate. Only samples with high motility (≥80%) were used in this study.

Sperm concentration
Sperm concentration was determined in a Neubauer haemocytometer according to the criterion of Figueroa et al. (2013).

pH and osmolarity
Each sample was analysed in triplicate. subsequently, and due to the low volume of semen produced by each male (< 0.5 ml), a pool was prepared. Merck pH indicator (5.5 to 9.0) paper was used for sperm pH.

In each trial (n = 9), to obtain the gamete quantities required for the experiment, semen samples were pooled (with a minimum of six males), in order to reduce the variability in sperm quality inherent among males (DeGraaf and Berlinsky, 2004).

Experimental Design
In each trial (n = 9), to obtain the gamete quantities required for the experiment, semen samples were pooled, in order to reduce the variability in sperm quality inherent among males (DeGraaf and Berlinsky, 2004). The semen pool was divided into five groups: 1) Control: undiluted semen; 2) treatment 1 (DS1): diluted semen in Hepes in a ratio of 1:3 semen: diluent (v/v); 3) treatment 2 (DS2): diluted semen in DMSO in a ratio of 1:3 semen: diluent (v/v); 4) treatment 3 (DS3): diluted semen in BSA in a ratio of 1:3 semen: diluent (v/v); 5) treatment 4 (DS4): diluted semen in Tris in a ratio of 1:3 semen: diluent (v/v); at 4°C and in the dark with constant stirring during 11 days. The stored samples were analysed at hourly.

Statistical Analysis
Results from spermatological characteristics are given as mean±SD. The variance homogeneity of the data was performed using Levene’s test. Differences among groups, data were compared by one-way ANOVA, followed by Tukey multiple range test a when significant differences were found at a 0.05 level. All measurements were carried out in triplicate.

Results

Fresh Semen Characteristics
Milt of M. cephalus is highly viscous, white in colour and the quantities of milt varied among individuals. Length-weight relationship of Liza saliens is W = 0.04491L2.6201 (R² = 0.88) (Figure 1).

Figure 1. Length-weight relationship of Liza saliens’s

The parameters in fresh semen were: duration of sperm motility = 80.9±1.5 s; sperm concentration = 14.0±8.9×10⁹ spermatozoa ml⁻¹ and pH = 6.7±0.2. The average volume of sperm was determined as 4.46 ± 0.72 ml.kg⁻¹. The highest and the lowest sperm volume was 5.88 ml.kg⁻¹ and 3.17 ml.kg⁻¹, respectively. The length-weight relationship of L. saliens’s with sperm volume is given below (Figure 2).
Evaluation of Sperm Function

Sperm motility

There were no significant differences in sperm motility between the DSI, DS2 and DS3 groups at the time of the first activation (P>0.05). At the second days, motility decreased significantly in the fourth group and the control group (mot =10±2.82%). From day 4 to 8 sperm MOT decreased significantly in diluted semen (DS2: 55.8±2.8% to 10.1±2.0% and DS3: 70.3±1.5% to 38.7±8.5%, respectively, Figure 3). End of the study highest motility on DS3 group. Sperm of undiluted samples was immotile at 2 day of storage.

Figure 3 Effect of short-term storage on sperm function of undiluted and diluted semen (1:1 v/v and 1:3 v/v) of Leaping mullet (L. saliens) for 8 days at 4°C.

Discussion

The results of this study indicate there may be a method to effectively short term storage and maintain the viability of Liza saliens sperm. For this purpose, the average weight of 753.6±47.21 g, the average length of 40.60±1.14 cm in the natural breeding period from the rootstocks identified as a natural breeding period, without any hormonal intervention was taken by the milking method. In a different study, it was determined that the sperm volume is higher in male fish which has hormone application, while there is no statistically significant difference in sperm motility, concentration and survival (Basavaraja and Hegde, 2005). It has been reported in previous studies that storage at low temperatures has a negative impact on sperm motility, viability and function ((Aguilar-Juárez et al., 2014; Aramli, 2013; Shalutina et al., 2013; Trigo et al., 2015). In our study, a significant decrease in motility was observed after storage and the results were in parallel with previous studies.

As in other species, it is important to know the pH of the sperm in the fish to be applied to the sperm (Akçay et al., 1995; Suquet et al., 1993). In this study, many researchers in the measurement of semen pH (Suquet et al., 1993; Billard and Cosson, 1992; Cevik et al., 2000; Hatipoglu et al., 2007) as in the pH range of 5.5-9.0 indicator papers were used. The pH of the sperm obtained in the study was found to be 6.7.

In previous studies, it was reported that by Astuirano et al. (2001) and Fauvel et al. (1999), sperm viability of cultured sea bass was found around 0.6–2 and 0.7–2.9 min under culture conditions. Additionally, this period was reported 0.4–2 and 3.6 min for Oncorhynchus mykiss (Yanagimachi et al., 1992; Billard, 1983). In Sparid, it was estimated 1.5–2 min for white sea bream, Diplodus sargus (Lahnsteiner and Patzner, 1998), 2.6–3.9 min for P. pagrus (Mylonas et al., 2003) and 1–3 min for sharpsnout sea bream, Diplodus puntazzo (Taddei et al., 2001). Besides, currently it was 6.9±0.8 min for sea bream and also in other cultured marine fish, Psetta maxima, 3 different periods for viability of sperm were reported as 3–5 min (Dreanno et al., 1999), 1–17 min (Suquet et al., 1994) and 2–3 min (Chauvaud et al., 1995). It is well reported that viability of sea bream (Sparus aurata) sperm was determined as 6.9±0.8 min (Engin et al., 2018). There is no study about this species for survival times of sperm. In our study, it is determined that the survival times of the leaping mullet sperm; 6 min in fresh sperm, 6.5 min in Hepes, 5 min 20 min in BSA, 3 min 20 min in Tris and 8.5 min in DMSO.

Dilution of milt in an extender is an important step for improving the quality of sperm after collection in aquatic animals (Ciereszko et al., 2000; Bobe and Labbe, 2008). Further, dilution ratio determines the success of short term preservation of milt at low temperature and production of fingerlings during fertilisation. In the present study, short term preservation of sperm in Liza saliens’s 1:3 dilution was found to be the best compared to other dilution ratios. Motility grade of spermatozoa decreased at higher dilution ratios probably indicating that all the intrinsic substances present in the seminal plasma, for example protein, glucose and ions loses their role at higher dilution (Jimenez et al., 2004; Mommens et al., 2008).

For cryopreservation, sperm membrane may be damaged after freezing and thawing procedure. All these processes negatively affect sperm fertilization capacity. After cryopreservation, sperm cell membrane, mitochondria, chromatin structure and acrosome may be damaged. Most of these damages were between -10 and -40°C (Cabrita et al., 2007). As a result of the short-term storage protocol, sperm samples will be transported to the laboratory environment in order to determine the effects of the factors mentioned in reproduction biology. In our study, at the end of the experiment, no significant difference was found for the duration of survival in the control and DS4 groups (P>0.05). The highest survival rate was found in DS3 group.

Conclusions

In previous studies, it was reported that short term storage for Mugil cephalus and Liza parsia (Chao et al., 1974; Balamurugan et al., 2016; Sukuraman et al., 2003). But there is no study about Liza saliens therefore it is a first record for short term storage. However, further studies are required to improve the sperm motility for longer storage period at low temperature by modifying the extender composition. The data generated from sperm quality studies from the fresh milt will be useful for effective management of cryopreservation of gametes for captive seed production in hatcheries.
References


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