Acclimation of juvenile *Mugil liza* Valenciennes, 1836 (Mugiliformes: Mugilidae) to different environmental salinities

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Survival and physiological parameters associated with metabolism and osmoregulation were evaluated in juveniles of the Lebranche mullet *Mugil liza* acclimated to different water salinities (5, 10, 20, 30, and 40‰) for 15 days. Room temperature (25°C) and photoperiod (12L:12D) were fixed. Fish were fed twice-a-day with commercial diet (28% crude protein) until satiation. After acclimation, whole body oxygen consumption was measured and fish were euthanized and sampled for blood, gills, and liver. Whole body oxygen consumption and plasma osmolality did not change in the range of salinities tested. The isosmotic point was estimated as 412.7 mOsmol kg⁻¹ (13.5‰). Gill Na⁺,K⁺-ATPase activity tended to be lower at 20 and 30‰, while liver glycogen content was significantly higher at 20‰ than at 5 and 40‰. These results indicate that juvenile *M. liza* is able to acclimate for a short-period of time (15 days) to a wide range of salinities (5-40‰). This condition is achieved through adjustments in gill Na⁺,K⁺-ATPase activity and carbohydrate metabolism to regulate plasma osmolality and aerobic/energy metabolism. Therefore, our findings support the idea of catching juveniles *M. liza* in sea water and rear them in estuarine and marine waters.

**Keywords:** Fish, Glycogen, Na⁺,K⁺-ATPase, Osmoregulation, Oxygen consumption.

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(Na⁺ and Cl⁻) is actively secreted at the gills. As a result, a water gain and salt loss is achieved, thus counteracting the obligatory osmotic loss of water and the diffusive gain of salts associated with the osmotic gradient existing between the fish blood and sea water. Using these homeostatic mechanisms, euryhaline teleost fish are able to keep their plasma osmolality more or less constant over a wide range of water salinities (Baldisserotto, 2009). Under steady state, marine teleosteos are hyposmotic to sea water, with plasma osmolality ranging from 370 to 480 mOsmol kg⁻¹. In turn, freshwater teleosts or euryhaline marine teleosts in fresh water have a lower plasma osmolality (260 to 330 mOsmol kg⁻¹), which is still much higher than that found in the ambient water (~1 mOsmol kg⁻¹) (for review: Jobling, 1995; Schmidt-Nielsen, 1996; Evans, 2008).

It is clear from the above that changes in energy consumption are expected to occur associated with fish osmotic regulation over a wide range of environmental salinities. According to Handeland et al. (1998), the osmoregulatory cost is proportional to the osmotic gradient. Therefore, lower energy expenditure would be expected to occur when fish blood plasma is isosmotic with the environmental medium. Indeed, an isosmotic environment has been reported to maximize growth in some euryhaline teleosts, since less energy would be spent in the osmoregulatory process (Woo & Kelly, 1995; Tsuzuki et al., 2007; Herrera et al., 2009; Nordlie, 2009). This phenomenon will be referred hereafter as the “Isosmotic Theory”. According to this theory, the energy spared on osmoregulation can be used for fish growth maximization. However, there is still no consensus about the importance of this energy sparing mechanism in improving fish growth. In fact, the effect of salinity on fish growth will depend on how this environmental parameter affects the standard metabolic rate, food intake, food conversion, and/or hormonal stimulation (Boeuf & Payan, 2001), which are shown to be species-specific (Jobling, 1994).

Accurate measurements of the amount of energy spared when fish are maintained at an isosmotic environment are difficult to perform because of the complex interactions existing among the osmoregulatory mechanisms and other physiological processes, which are in turn influenced by environmental factors, as well as the methodology employed. Boeuf & Payan (2001) reported that the energy spent by fish with osmoregulation can range from 10% to more than 50% of the total energy budget.

Plasma osmolality, gill Na⁺, K⁺-ATPase activity, whole body oxygen consumption and liver glycogen content are physiological parameters usually measured to evaluate the effect of salinity in fish (McCormick, 1995; Baldisserotto et al., 2007; Saoud et al., 2007). In fact, Na⁺, K⁺-ATPase is the main enzyme responsible for NaCl transport across the teleost gills. In freshwater fish, it is associated with salt absorption, while in seawater fish it is related to salt secretion. Therefore, gill Na⁺, K⁺-ATPase activity is usually increased in extreme (low and high) environmental salinities where the osmolality gradient between the fish plasma and ambient salinity is maximized. As a consequence, gill Na⁺, K⁺-ATPase activity is directly related to the maintenance of plasma osmolality homeostasis (Baldisserotto et al., 2007; Evans, 2008).

It is clear from the above that osmoregulation is an energy-demanding process. As previously mentioned, energy spent by fish with osmoregulation can be as high as 50% of the total energy budget (Boeuf & Payan, 2001). At this point, it is important to note that glucose is the main source of energy for fish osmoregulation through the oxidative metabolism (Pérez-Robles et al., 2012). In fish, as in other vertebrates, this carbohydrate is mainly stored as glycogen in the liver (Jobling, 1994; Schmidt-Nielsen, 1996). Therefore, measurements of whole body oxygen consumption and liver glycogen content can be good indirect indicators of the energy demand associated with osmoregulation. Theoretically, the optimal environmental salinity for rearing fish would be that showing a combination of lower gradient between ambient osmolality and fish plasma osmolality, where a resulting lower gill Na⁺, K⁺-ATPase activity would be observed. In this case, a lower whole body oxygen consumption and higher liver glycogen content would be expected to occur. These conditions would suggest that a likely higher amount of energy could be spared by fish, thus allowing a better growth.

The taxonomic status of Mugilidae from the Caribbean and South American Atlantic coast has been revised. Fish previously identified as M. platanus are now correctly identified as the Lebranche mullet M. liza (Menezes et al., 2010). This teleost fish is an important item for estuarine and coastal fisheries in Southern South America (Reis & D’Incao, 2000). Furthermore, it has been considered for aquaculture in estuarine and marine waters in Brazil. The Lebranche mullet shows a small dietary protein requirement (Carvalho et al., 2010) and occupies a low position in the food web (Oliveira & Soares, 1996). In the scope of the present study, it is important to mention that the mullet M. liza can face very significant changes in environmental osmotic conditions during its life cycle. In South America. It inhabits tropical and subtropical waters from Rio de Janeiro (southeastern Brazil) to Argentina, being especially found at the Patos Lagoon estuary and the adjacent coastal region (Rio Grande do Sul state, southern Brazil) (Vieira et al., 1998). The Lebranche mullet M. liza is considered as being an estuarine-dependent fish species (Vieira et al., 1998). As a catadromous fish, mature individuals displace from the estuarine area to the sea for reproduction and spawning. After spawning, its eggs and larvae are transported to the Patos Lagoon estuary, where they develop until reaching sexual maturity. Mature individuals move then to the sea for reproduction and spawning (Vieira et al., 1998). Therefore, juveniles M. liza face abrupt, frequent and marked changes in ambient osmolality during its development and growth in the Patos Lagoon estuary (Seeliger et al., 1998).
Little is known about the effect of environmental parameters on the physiology of juvenile Lebranche mullet. It was shown that growth of juvenile *M. liza* is improved at temperatures between 25 and 30°C (Okamoto et al., 2006). Regarding salinity, Sampaio et al. (2002) showed that tolerance of juveniles of the Lebranche mullet to ammonia and nitrite is improved in fish acclimated to intermediate and high salinities with respect to those maintained in fresh water. Fonseca Neto & Spach (1998/1999) reported no mortality after 96 h of abrupt transfer of juvenile *M. liza* from sea water (30‰) to lower salinities (15, 10 and 5‰). Very recently, we have shown that salinity influenced growth without significant changes in biochemical parameters (gill Na⁺/K⁺-ATPase and liver glycogen content) in juvenile Lebranche mullet *M. liza* after a 40-days rearing period in different salinities ranging from 0 to 24‰ (Lisboa et al., 2015).

The main objective was to evaluate survival and the response of physiological parameters associated with osmoregulation in juveniles of the Lebranche mullet *M. liza* acclimated sea water (30‰), abruptly transferred to a wide range of salinities (5-40‰) and maintained at these salinities for 15 days. Physiological endpoints analyzed included plasma osmolality, gill Na⁺,K⁺-ATPase activity, whole body oxygen consumption, and liver glycogen content. As observed for other euryhaline marine teleosts, it is expected that juveniles of the Lebranche mullet *M. liza* would show plasma osmolality varying from 260 to 480 mOsmol kg⁻¹ after full acclimation (15 days) to salinity in the range of ambient salinities tested (5-40‰). It is also expected that whole body oxygen consumption and gill Na⁺,K⁺-ATPase activity would be lower in intermediate salinities (10-30‰), while liver glycogen content would be higher under this condition.

**Material and Methods**

**Fish collection and acclimation.** Juvenile *M. liza* were captured at Cassino Beach (Rio Grande, RS, southern Brazil) and transferred to the Laboratory of Marine Fish Culture of the Universidade Federal do Rio Grande (FURG), Rio Grande, RS. Fish acclimation to laboratory conditions was performed in two 1,000-L circular fiber tanks containing sea water at 25°C and salinity corresponding to that of the collection site (30‰). Tanks were maintained under natural photoperiod (12 h L:12 h D). Fish were fed twice-a-day (from 10:00 to 11:00 h and from 17:00 to 18:00 h) with a commercial diet (28% crude protein) until satiation. Feces produced were siphoned out daily and at least 50% of the experimental medium was renewed. All procedures used for capture, maintenance and experimentation using juveniles of the Lebranche mullet followed the practices actually recommended by the Brazilian Council for Control of Animal Experimentation (CONCEA).

**Experimental design.** After acclimation to laboratory conditions for 20 days, 36 fish (wet body weight: 54.1 ± 0.9 g; total length: 17.7 ± 0.1 cm) were fastened for 24 h, individually weighed (wet body weight; electronic scale; precision: 0.01 g) and measured (total length), randomly divided into six groups (6 fish per group). The first group of fish were immediately analyzed for whole body oxygen consumption and had samples collected for plasma osmolality, gill Na⁺,K⁺-ATPase activity and liver glycogen content, as described below. Juvenile mullets from the other 5 groups were kept in sea water (30‰) or abruptly transferred to diluted sea water (5, 10, and 20‰) or concentrated sea water (40‰). Sea water at the different experimental salinities was obtained by diluting concentrated sea water (40‰ salinity) with tap water. In turn, concentrated sea water was obtained by evaporation at 50°C. Fish were maintained in 300-L circular fiber tanks at the experimental salinities for 15 days. Tanks were provided with a closed circulating system and 80% of the total water volume was renewed every day. Fish stocking density corresponded to ~1 g L⁻¹, as recommended for physiological studies.

The 15-days period of experiment was selected considering that juvenile *M. liza* shows ultrastructural changes in gill chloride cells, which are directly associated with acclimation to fresh and sea water, within few hours or days (4 days) after transfer to a wide range of environmental salinities (Fonseca Neto & Spach, 1998). Therefore, it would be expected that mullets employed in the present study would be fully acclimated 15 days after the transfer to the experimental salinities tested. In addition, the experimental time employed in the present study (15 days) was already used to acclimate *M. liza* to different salinities for previous physiological (Carvalho et al., 2010) and toxicological (Sampaio et al., 2002) studies.

Water temperature (hand termometer), dissolved oxygen content (oxy meter YSI Model), salinity (refractometer, Atago), pH (pH meter, Quimis), and total ammonia (American Public Health Association (APHA, 2005)) were measured daily. Water samples were also collected over the experimental period for osmolality measurements using a semi-micro osmometer (Knauer, Germany) based on the freezing depression point. Osmolality data were expressed in mOsm kg⁻¹.

**Whole body oxygen consumption.** At the end of the experimental period, whole body oxygen consumption was measured following the procedures described by Cunha et al. (2009). Fish were fastened for 24 h prior to oxygen consumption measurements. Water aeration was removed and the tank was completely sealed with a transparent plastic sheet to avoid any air diffusion. Dissolved oxygen content in the water was recorded every 15 min for up to 1 h. Measurements were performed in triplicate for each tank using an oximeter (YSI model Hexis 55). They were never performed when the water saturation level with oxygen
dropped to 70% of its maximum to avoid a possible effect of this water parameter on fish oxygen consumption. The rate of oxygen consumption (OC) was calculated using the following equation:

\[
OC = \frac{(O_f - O_i) x V}{T x B}
\]

where \(O_i\) and \(O_f\) correspond to the initial and final dissolved oxygen content in the water (mg O\(_2\) L\(^{-1}\)), respectively, \(V\) is the tank volume (L), \(B\) is the fish biomass (g), and \(T\) is the measurement duration (h). Results were expressed in mg O\(_2\) g wet body mass\(^{-1}\) h\(^{-1}\).

**Tissue collection and analysis.** After the oxygen consumption measurement, fish were anesthetized (50 ppm benzocain) and blood samples were collected by heart puncture using 1-mL heparinized disposable syringes and centrifuged. Plasma obtained was collected and stored in ultrafreezer (-80°C) for further osmolality measurement, as described below. The second gill arch at the left side of the gill chamber and the liver were dissected, immediately frozen in liquid nitrogen and stored in ultrafreezer (-80°C) for further measurement of gill Na\(^+\),K\(^-\)-ATPase activity and liver glycogen content, respectively, as described below.

Fish plasma osmolality was determined using the semimicro osmometer (Knauer, Germany) based on the freezing depression point. The fish isosmotic point was assessed by considering the intersection of the isosmotic line with the regression line estimated between plasma and water osmolality as described by Sampaio & Bianchini (2002). Results were expressed in mOsm kg\(^{-1}\).

Gill samples were homogenized with 300 µl of a buffer solution (pH 7.3) containing 150 mM sucrose, 10 mM ethylenediaminetetraacetic acid, 50 mM imidazole, and 11.5 mM sodium deoxycholate, and centrifuged at 10,000 g for 30 min (4°C). The supernatant was used for Na\(^+\),K\(^-\)-ATPase activity determination at 25°C, following procedures described by McCormick (1993). Sample absorbance was measured using a spectrophotometer (ELX 800 Universal Microplate Reader/Bio-Teck Instruments, Winooski, Vermont, USA). The protein concentration procedures described by Carr & Neff (1984). In this case, the homogenized sample was split into two aliquots of each sample using a commercial reagent kit based on the glucose-oxidase method (Doles, Goiânia, GO, Brazil). Sample absorbance was measured using the spectrophotometer (ELX 800 Universal Microplate Reader/Bio-Teck Instruments, Winooski, Vermont, USA). Glycogen concentration was calculated considering the difference in glucose concentration between the aliquots digested and nondigested with the enzyme. Results were expressed in mg g wet tissue\(^{-1}\).

**Data presentation and analysis.** Results were expressed as mean ± standard error (n = 6). For all parameters analyzed, significant differences among salinities were assessed by One-Way analysis of variance (ANOVA) followed by the Tukey’s test. For each salinity, comparison of mean values between water and plasma osmolality was performed using the non-paired Student t-test. Water and plasma osmolality data were subjected to linear regression analysis. Gill Na\(^+\),K\(^-\)-ATPase activity and liver glycogen content data were subjected to non-linear regression analysis (polynomial, quadratic). For all analyses, the level of significance adopted was 95% (\(α = 0.05\)).

**Results**

Water osmolality was significantly different among the experimental salinities (Table 1). Dissolved oxygen content, temperature and ammonia concentration (NH\(_3\)-N) in the water did not change throughout the experimental period and among treatments (\(P > 0.05\)). Therefore, a general mean value was calculated for dissolved oxygen content (6.34 ± 0.18 mg O\(_2\) L\(^{-1}\)), temperature (24.0 ± 0.20°C) and ammonia concentration (0.03 ± 0.00 NH\(_3\)-N). However, water pH increased significantly (\(P < 0.05\)) from 7.70 ± 0.06 at the lowest salinity (5‰) to 8.02 ± 0.06 at the highest salinity (40‰).

No fish mortality was observed over the whole experimental period. Mean whole body oxygen consumption of juvenile mullets before transference to the experimental salinity (time 0) was 0.34 ± 0.03 mg O\(_2\) g\(^{-1}\) h\(^{-1}\). No significant change was observed in whole body oxygen consumption in the range of salinities tested (Table 1).

**Table 1.** Water osmolality, plasma osmolality and whole body oxygen consumption of juveniles of the Lebranche mullet *Mugil liza* maintained in different salinities for 15 days. Data are expressed as mean ± standard error (n = 6). Different letters indicate significant different mean values among salinities for each parameter analyzed (\(P < 0.05\)).* indicates significant difference between water and plasma osmolality for each experimental salinity (\(P < 0.05\)).

<table>
<thead>
<tr>
<th>Salinity (%)</th>
<th>Water osmolality (mOsm kg(^{-1}))</th>
<th>Plasma osmolality (mOsm kg(^{-1}))</th>
<th>Oxygen consumption (mg O(_2) g(^{-1}) h(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>168 ± 3.1*</td>
<td>473.4 ± 41.4*</td>
<td>0.36 ± 0.08*</td>
</tr>
<tr>
<td>10</td>
<td>332 ± 4.7b</td>
<td>330.0 ± 41.0*</td>
<td>0.43 ± 0.03*</td>
</tr>
<tr>
<td>20</td>
<td>565 ± 2.5c</td>
<td>401.5 ± 11.8*</td>
<td>0.38 ± 0.07*</td>
</tr>
<tr>
<td>30</td>
<td>910 ± 5.2d</td>
<td>467.4 ± 35.8*</td>
<td>0.32 ± 0.05*</td>
</tr>
<tr>
<td>40</td>
<td>1080 ± 3.5e</td>
<td>414.4 ± 54.5*</td>
<td>0.36 ± 0.07*</td>
</tr>
</tbody>
</table>
Mean plasma osmolality of juvenile mullets before transference to the experimental salinities (time 0) corresponded to 422.0 ± 17.0 mOsmol kg\(^{-1}\). As observed for oxygen consumption, plasma osmotic concentration did not change among water salinities. However, a significant difference was observed between plasma and water osmolality in all salinities tested, except at 10‰. Plasma osmolality was higher than water osmolality in fish maintained at 5‰ and lower than water osmolality in fish kept at 20, 30 and 40‰ (Table 1). The slope value of the regression line between plasma and water osmolality was not significant (b = 0.023; \(P > 0.05\)). The isosmotic point of juveniles of the Lebranche mullet was estimated as 412.7 mOsmol kg\(^{-1}\), which corresponded to 13.5‰ (Fig. 1).

**Fig. 1.** Plasma osmolality in juveniles of the Lebranche mullet _Mugil liza_ as a function of ambient water osmolality. Values are expressed as mean ± standard error (n = 6). Data were fitted using a linear regression analysis (y = a + bx). The straight line corresponds to the isosmotic line while the arrow indicates the isosmotic point.

Mean gill Na\(^{+}\),K\(^{-}\)-ATPase activity of juvenile mullets before transference to the experimental salinities (time 0) corresponded to 0.81 ± 0.22 μmoles ADP mg protein\(^{-1}\) h\(^{-1}\). Enzyme activity was not significantly affected by salinity, but tended to be lower at intermediate salinities (20 and 30‰) and higher at extreme salinities (5 and 40‰). When taken together, mean values followed an inverse polynomial quadratic function (Fig. 2).

**Fig. 2.** Gill Na\(^{+}\),K\(^{-}\)-ATPase activity in juveniles of the Lebranche mullet _Mugil liza_ maintained at different salinities for 15 days. Values are expressed as mean ± standard error (n = 6). Data were fitted using a non-linear regression analysis (polynomial; quadratic).

Mean liver glycogen content of juvenile mullets before transference to the experimental salinities (time 0) was 0.22 ± 0.03 mg g\(^{-1}\). In contrast to the other parameters analyzed, liver glycogen content was significantly affected by salinity. It was significantly higher in fish maintained at 20‰ than in those maintained at 5 or 40‰, but similar to those at 10 and 30‰. When taken together, mean values of liver glycogen content followed a direct polynomial quadratic function (Fig. 3).

**Fig. 3.** Liver glycogen content in juveniles of the Lebranche mullet _Mugil liza_ maintained at different salinities for 15 days. Values are expressed as mean ± standard error (n = 6). Data were fitted using non-linear regression analysis (polynomial; quadratic).

**Discussion**

Juveniles of the Lebranche mullet _Mugil liza_ are naturally subjected to important changes in environmental salinity over its life cycle. An adequate ability to cope with these changes is imperative for _M. liza_ to complete its life cycle.
This ability was clearly observed in juvenile *M. liza* exposed to a wide range of salinities. This statement is based on the following facts: (1) juvenile mullets tested in the present study were able to survive after abrupt transfer from sea water (30‰) to a wide range of experimental salinities (5-40‰); (2) no significant change in plasma osmolality was observed after acclimation of juvenile mullets to the different environmental salinities, and (3) the slope of the regression line between plasma and water osmolality was not significant (b = 0.023), indicating a strong capacity of juvenile mullets to regulate plasma osmolality in a wide range of water salinities.

The well-developed osmoregulatory capability showed by juvenile *M. liza* tested in the present study (18 cm body length) combined with the fact that juvenile *M. liza* smaller than 2.8-3.3 cm are not able to tolerate abrupt transfer from sea water (30‰) to fresh water (Fonseca Neto & Spach, 1998/1999) suggests that juveniles of the Lebranche mullet have already fully developed their osmoregulatory ability. In fact, changes in this ability are shown to occur over the ontogenetic development of other fish species such as the fat snook *Centropomus parallelus* (Tsuzuki *et al.*, 2007) and the Brazilian flounder *Paralichthys orbignyanus* (Sampaio *et al.*, 2007). Unfortunately, there is no report describing the ontogenesis of osmoregulation in the Lebranche mullet *M. liza*. Full osmoregulatory capability of the stripped mullet *M. cephalus* is reached when individuals achieve 4.0-6.9 cm body length (Nordlie *et al.*, 1982), a range of body sizes lower than the mean body size of juveniles of the Lebranche mullet *M. liza* analyzed here (18 cm body length).

No significant effect of environmental salinity on plasma osmolality was observed in juveniles of the Lebranche mullet *M. liza*. However, a slightly positive but not significant correlation (linear regression; b = 0.023; P > 0.05) between environmental and plasma osmolality was observed. This fact could be associated with a possible dehydration suffered by fish when exposed to hyperosmotic environments (Sampaio & Bianchini, 2002; Resley *et al.*, 2006; Saoud *et al.*, 2007). In fact, plasma osmolality in juveniles of the Lebranche mullet *M. liza* maintained at 5‰ (mean value: 473.4 ± 41.4 mOsmol kg<sup>-1</sup>) or 30‰ (mean value: 467.4 ± 35.8 mOsmol kg<sup>-1</sup>) was higher than that reported for diadromous fish acclimated to fresh water (310.0 ± 6.5 mOsmol kg<sup>-1</sup>) or sea water (402.6 ± 20.4 mOsmol kg<sup>-1</sup>), including the stripped mullet *M. cephalus* (fresh water: 326.7 mOsmol kg<sup>-1</sup>; sea water: 356.7 mOsmol kg<sup>-1</sup>) (Nordlie *et al.*, 1982; Nordlie, 2009).

The isosmotic point for juveniles of the Lebranche mullet was estimated as 412.7 mOsmol kg<sup>-1</sup>, which corresponded to 13.5‰. In this case, fish is hyper-regulating above this environmental salinity and hypo-regulating above it. According to the idea that energy consumption would be reduced at the isosmotic condition, a lower gill Na<sup>+</sup>,K<sup>-</sup>-ATPase activity and oxygen consumption would be expected in water salinities close to the isosmotic point. Reduced energy consumption, indirectly indicated in the present study by higher liver glycogen content, would be also expected in water salinities close to that where fish is isosmotic respect to the environment. However, whole body oxygen consumption and gill Na<sup>+</sup>,K<sup>-</sup>-ATPase activity of juveniles of the Lebranche mullet did no change significantly among water salinities. Moreover, despite the fact that the gill Na<sup>+</sup>,K<sup>-</sup>-ATPase activity and liver glycogen content followed a U-shape curve according to the water salinity, the lower mean values of enzyme activity were observed at 20 and 30‰ and the higher mean value of liver glycogen content was found at 20‰. Also, it is important to note that juveniles of *M. liza* reared at 24‰ were shown to grow better than those maintained in fresh water for 40 days. However, fish reared at 12‰, which is very close to its iso-osmotic point (13.5‰), showed similar growth to those maintained at 0, 6 and 24‰ (Lisboa *et al.*, 2015). Taken together, these findings clearly do not support the “Isosmotic Theory” for juveniles of the Lebranche mullet *M. liza*.

Regarding gill Na<sup>+</sup>,K<sup>-</sup>-ATPase activity, our data are in contrast to those reported for juveniles of the coho salmon *Oncorhynchus kisutch* (Morgan & Iwama, 1998) and gilthead sea bream *Sparus aurata* (Laiz-Carrión *et al.*, 2005). In these fish species, gill enzyme activity also followed a U-shape pattern with the environmental salinity, but the lowest value was found in water salinity close to the isosmotic point. Nevertheless, our results are in agreement with those reported for the rabbitfish *Siganus rivulatus*, where enzyme activity also followed a U-shape pattern across a wide range of water salinities with the lower values being observed in water salinities not close to that corresponding to the isosmotic point (14.6‰) (Saoud *et al.*, 2007). Furthermore, it was shown that gill Na<sup>+</sup>,K<sup>-</sup>-ATPase activity in the milkefish *Chanoschanos* did not show a clear pattern of change across salinities, with the lower enzyme activity value being found at 35‰. Actually, Morgan & Iwama (1998) have suggested that the response of the gill Na<sup>+</sup>,K<sup>-</sup>-ATPase activity to environmental salinity may be species-dependent, which could explain the great variability observed among euryhaline fish species.

The U-shape curve displayed by the gill Na<sup>+</sup>,K<sup>-</sup>-ATPase activity in *M. liza* according to the water salinity combined with the observed lack of change in plasma osmolality is indicative that enzyme activity was stimulated at the extreme salinities (5 and 40‰). Under these conditions, the higher Na<sup>+</sup>,K<sup>-</sup>-ATPase activity would be a positive response to compensate the higher loss and gain of NaCl occurring at 5 and 40‰, respectively. In fact, increased changes in NaCl fluxes across the gills would be a consequence of the higher osmotic gradient existing between the ambient osmolality and fish plasma osmolality at 5 and 40‰. In this case, a higher gill Na<sup>+</sup>,K<sup>-</sup>-ATPase activity would increase the absorption and secretion of NaCl in fish at 5 and 40‰, respectively. Gill enzyme activity is described to be regulated by neurochemical and/or endocrine mediators released after stimulation by different triggers, including external salinity and blood osmolality (Evans, 2008).
Despite the gill Na⁺,K⁺-ATPase activity in *M. liza* followed a U-shape curve according to the water salinity, no significant difference was observed among the experimental salinities. This finding is in complete agreement with previous studies on the effect of salinity on the ultrastructural changes in gills of *M. liza* after acclimation to different salinities. Fonseca Neto & Spach (1998) reported that significant ultrastructural changes in chloride cells are only observed in *M. liza* subjected to salinities lower than 3‰. Furthermore, they showed that no significant difference was observed among mullets subjected to salinities ranging from 5 to 34‰, a similar range of salinities used in the present study.

In contrast to our findings, oxygen consumption in *M. cephalus* acclimated for 8 days in laboratory augmented with increasing water salinity (Nordlie & Leffler, 1975). In turn, the oxygen consumption changes observed in the present study with the juvenile *M. liza* acclimated for 15 days to different environmental salinities correspond to the Type 4 response described by Kinne (1967). Similar to our findings, Morgan & Iwama (1998) reported that water salinity did not influence oxygen consumption in juvenile coho salmon. Therefore, it seems that 15 days is a suitable period of time to acclimate juveniles of the Lebranche mullet *M. liza* to different environmental salinities without changes in the aerobic metabolism.

Osmoregulation is an energy demanding activity and is fuelled mainly by glucose (Pérez-Robles *et al.*, 2012). This carbohydrate is an important source that provides energy for animal metabolism and an increased plasma concentration of this metabolite after osmotic stress has been reported for several fish species (Herrera *et al.*, 2009). Depending on stress intensity and duration, the plasma glucose level can be sustained through glycogenolysis (Baldisserotto *et al.*, 2007), which is a known secondary response to stress. The liver glycogen concentration in juveniles of the Lebranche mullet was significantly lower at the extreme salinities (0 and 40‰) than at 20‰. This finding is in complete agreement with that observed for gill Na⁺,K⁺-ATPase activity, suggesting that the energy demand to regulate the plasma osmotic concentration in extreme conditions was likely provided through glucose oxidation. In this case, a higher glycogen mobilization from liver would occur, thus leading to the observed lower glycogen content in the liver.

Finally, many authors have discussed on the advantages of keeping fish under isosmotic conditions. This practice would imply that the energy saved in the absence of an important osmotic work could maximize fish growth (Boeuf & Payan, 2001; Saoud *et al.*, 2007; Tsuzuki *et al.*, 2007). However, this theory seems to be species-specific, since it has been proved for some fish species like *Sparus sarba* (Woo & Kelly, 1995), but not for others like *Paralichthys orbignyanus* (Sampaio & Bianchini, 2002). Very recently, we have shown that juveniles of the Lebranche mullet *M. liza* reared at 12‰, a salinity close to the isosmotic point, did not display a better growth than those cultivated in a wide range of salinities (0-24‰) (Lisboa *et al.*, 2015). Results from the present study are in agreement with this finding, thus suggesting that the idea of improving fish growth at isosmotic condition cannot be considered for juveniles of the Lebranche mullet *M. liza*. However, it is important to note that the present findings clearly indicate that juveniles of *M. liza* is able to keep its plasma osmotic homeostasis after 15 days of acclimation to a wide range of salinities through adequate adjustments in gill Na⁺,K⁺-ATPase and liver glycogen metabolism, without significant changes in the aerobic metabolism. This would suggest that juvenile mullets could be captured in sea water and abruptly transferred and reared in estuarine waters.

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References


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