Osmotic induction marking with Alizarin Red S on juveniles of pejerrey,
*Odontesthes bonariensis* (Atherinopsidae)

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Juveniles of pejerrey, *Odontesthes bonariensis*, were exposed to 0.1% Alizarin Red S (ARS) alone or with a previous immersion in 2.2% saline solution (Osmotic Induction, OI) to enhance the ARS marking method. Fish were marked in the field and immediately released in 1 m³ cages in “La Salada de Monasterio” lagoon, Chascomús, Buenos Aires, Argentina. After 73 days, clear marks were observed in the otoliths, caudal fin rays and scales with both treatments, being the intensity of the signal in the scales of OI+ARS treated fish higher. On the other hand, no marks were observed in the control group on the same structures. Approximately one year post-treatment (385 days), only marks in caudal fin rays were found clearly in OI+ARS treated fish. After this period, no significant differences in total length or weight between marked or control fish were observed and the mortality ranged between 30-40 % in all cages. These results provide strong evidence for the potential applicability of this cost-effective marking technique in differentiation of wild and hatchery-produced pejerrey. The success in the caudal fin rays marking is also important because it is easy to do and does not require the sacrifice of fish.

*Key words:* Fin rays, Fish tag, Otolith, Scales.

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**Introduction**

Pejerrey *Odontesthes bonariensis* (Atherinopsidae) is a brackish water species native to lakes and lagoons of Buenos Aires Province, Argentina and Rio Grande do Sul, Brazil (Dyer, 2006). Among other continental species from Argentina, the pejerrey is considered the most popular by local anglers because of its culinary value (López *et al*., 2001). It is also considered as a promising candidate for aquaculture (Somoza *et al*., 2008) as intensive culture methods had been successfully achieved in Argentina (Miranda *et al*., 2006; Velasco *et al*., 2008). As part of a long time tradition in Argentina, pejerrey larvae and juveniles are being released every year in shallow lakes and rivers of different argentine...
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provinces (Velasco *et al*., 2008). Nevertheless no evaluation of the success of this practice had been ever performed.

Several methods have been developed with the common goal of stock identification and/or discrimination from wild populations including the use of tags, mark-recapture, morphology of scales or otoliths, and thermal and chemical marking (Campana & Neilson 1985; Begg & Waldman, 1999; Crook *et al*., 2007). Among these methods, chemical immersion of fish in a solution that contains a fluorescent dye has been used to mark internal calcified structures, such as otoliths (Brooks *et al*., 1994; Skov *et al*., 2001; van der Walt & Forager 2003). Similar procedures have also resulted in the obtention of external marks, which allowed for a non-invasive detection (Mohler, 2003; Bashey, 2004; Negus & Tureson, 2004). The most commonly used were oxytetracycline (OTC), hydrochloride, calcein, alizarin complexone (AC) and alizarin red S (ARS). However, calcein, ARS and AC performed better than other chemicals because: marks can be detected and read clearly, the effect over survival rates was proven to be less negative, and all can be used in salt water (Tsukamoto *et al*., 1989; Van der Walt & Faragher, 2003; Taylor *et al*., 2005; Baer & Rösch, 2008; Liu *et al*., 2009). Even though previously performed experiments using calcein have obtained clear and long-lasting marks, this chemical compound is expensive and was found to be toxic when fishes were exposed to it for long periods of time (Brooks *et al*., 1994; Bumgardner & King, 1996; Frenkel *et al*., 2002). On the other hand, the use of ARS has been proven to be successful in marking fish at high scale (mass-marking) and it provides an alternative at only a fraction of the cost of the calcein and AC markers (Blom *et al*., 1994; Nagiec *et al*., 1995; Lagardere *et al*., 2000). Results from different studies that used ARS have shown that the compound produces highly readable marks for long periods of time (up to 842 days) in laboratory reared individuals (Crook *et al*., 2007).

The development of the ‘osmotic induction’ (OI) technique for fish marking with calcein has made possible to mark thousands of fish in less than 10 minutes (Mohler, 2003). Also, ARS has been utilized as a less expensive alternative to calcein for the OI, and results had shown the obtention of clear visible external marks after 9 months in the golden perch *Macquaria ambigua* (Crook *et al*., 2007).

In this context, the main objective of this study was to develop a quick and efficient protocol for administering ARS in pejerrey, in order to produce visible fluorescent marks and optimize this technique using a previous osmotic induction. The results obtained will be useful for the identification of captive raised juveniles stocks to be then released in water bodies.

**Material and Methods**

**Experimental fish and immersion marking.** On July 8th of 2009, 90 juveniles (total length 11±0.2 cm; weight 7±0.45 g) were selected from a stock of captive pejerrey from IIB-INTECH aquaculture facilities reared following Colautti *et al*., (2010) in the “La Salada de Monasterio” lagoon (35.8331S., 57.8871W), Chascomús, Buenos Aires, Argentina. The marking procedures were performed *in situ*, on a boat, using plastic containers for immersion and rinsing purposes. Fish were divided in three groups of 30 individuals and immersed in: A) ARS solution 0.1% for 10 min; B) Osmotic shock solution of 2.2% of salinity for 10 minutes and a posterior immersion in 0.1% ARS solution for 10 min; and C) Water from the lagoon (0.2% of salinity) for 10 min (control group) at approximately 15°C. All the solutions were prepared with water from the lagoon and after treatments all fish were rinsed with the same water and released in cages of 1 m³ that were already set up in the lagoon. During approximately one year, fish from each cage were sampled in order to verify the presence of fluorescent marks as well. Optimal ARS dosages and exposure times were established during preliminary assays, where increasing ARS concentrations led to an increase in mortality, and decreasing ARS concentrations produced no mark in any structure.

**Sampling and mark analyses.** All the structures were analyzed immediately after the obtention including the sagitta otolith (Fig. 1a), scales (Fig. 1b) from the lateral flank of the body (scales were extracted from right sub-ocular region), and caudal fin rays (Fig. 1c). The criterion used to choose the sagitta as the optimal pair of ear bones was its larger size compared to the lapilli and asterisci, which facilitated the extraction (Campana & Neilson, 1985). The otoliths were all freed from adherent tissues and rinsed with water. All samples were observed directly without resin and polishing (Liu *et al*., 2009). After removed, pictures of the structures were taken in a darkened room using a fluorescence microscope (Nikon Eclipse E600- G-2E/C TRITC Filter) attached to a digital camera (Nikon Digital Sight DS-U2). All photographs were obtained using the same magnification and exposure time. To assess the intensity of the ARS marks with and without osmotic induction a blind test was performed where the reader did not know the origin of the samples. At least three fish for each treatment and time period were analyzed. At the end of the experiment, growth and survival among the control and treatment groups were compared using ANOVA test.

**Results**

During the initial minutes of the marking procedures, pejerrey juveniles seemed to have lost equilibrium and were slowly floating towards the water surface. A few minutes later, all individuals showed signs of recovery, such as normal swimming behavior near the surface and no mortality was observed.

After 73 days post-treatment, clear marks were observed for both marking procedures, in otoliths (Fig. 2a-b), scales (Fig. 2c-d) and caudal fin rays (Fig. 2e-f). The intensity of the signal in scales was higher in fish marked with OI+ARS (Fig. 2c-d) and caudal fin rays (Fig. 2e-f). The intensity of the ARS marks with and without osmotic induction was observed.

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After approximately one year (385 days) only marks in caudal fin rays were observed in fish treated with OI+ARS (Fig. 2g). All control fish structures and, otoliths and scales for 385 days post-immersion showed no marks under fluorescence microscope and viewed like figure 2h.

At the end of the analyzed period no significant differences in total length or weight between marked or control fish were observed (P>0.05; data not shown), and the mortality ranged between 30-40% in all cages.

Discussion

The pejerrey swimming behavior observed during the marking process (lost equilibrium and floated immediately to the surface) was also reported for the golden perch subjected to the same treatment (Crook et al., 2007). It is important to note that no differences in mortality and growth during the analyzed period were observed between treatment and control group.

The immersion of pejerrey juveniles in ARS or OI+ARS solutions resulted in a 100% marking success after 73 days post-treatment. Despite variations in the quality and lasting of the marks, fluorescent signals were detected in both treatments and all the structures analyzed. These findings are in agreement with several studies that showed ARS effectiveness for immersion marking, as well as its potential in biological research and evaluation of stock enhancement programs (Bashey, 2004; Baer & Rösch, 2008; Crook et al., 2007; Liu et al., 2009).

It is important to note that no autofluorescence was observed in the structures analyzed, and marked and unmarked fish were easily distinguishable. Similar observations were reported by Bashey (2004) in juvenile guppies Poecilia reticulata and by Crook et al (2007) in golden perch marking with ARS. None of the treatments have affected pejerrey growth and survival (Colautti et al., 2010), and these results are in agreement with data reported by Liu et al (2009) for the Japanese flounder.

A higher signal intensity was found in the scales of OI+ARS treated fish after 73 days post-treatment, and only marks in caudal fin rays were found after more than 1 year post-immersion in OI+ARS. These results showed that the osmotic induction before immersion in ARS produces higher intensity marks than the direct immersion in ARS as it was previously reported in golden perch by Crook et al (2007) and using calcein in Atlantic salmon (Mohler, 2003). The success of this technique lies on the concept of the osmotic potential, as fish are exposed to a hyperosmotic environment when compared to the internal individual tissues and fluids. Hypothetically, this osmotic difference results in water loss from body fluids and then, when exposed to the marking solution, a fast uptake of the compound as some sort of replacement (Conte, 1969). Afterwards, the ARS binds to the calcium in fin rays, scales and otoliths.
Fig. 2. After 73 days post-treatment, clear marks were observed for both marking procedures, in otoliths (Figs. 2a, b), scales (Figs. 2c, d) and caudal fin rays (Figs. 2e, f). The intensity of the signal in scales was higher in fish marked with OI+ARS (Fig. 2d). After 385 days, only marks in caudal fin rays were observed in fish treated with OI+ARS (Fig 2g). All control fish structures and, otoliths and scales for 385 days post-immersion showed no marks under fluorescence microscope (Fig. 2h). a, c, e: ARS treatment; b, d, f, g: OI+ARS treatment. Scale bar = 200 µm.
What is important to stress is that, as shown in this study, otolith marks could be easily observed without any preparation of the sample such as the use of glycerin, or polishing the earbone surface (Vigliola, 1997; Taylor et al., 2005). Therefore, the amount of time required in order to verify the existence of the mark has been greatly reduced. However, when it comes to the verification of the marks, otoliths still require the sacrifice of the individuals. But as the OI+ARS treatment resulted in clear marks in the scales and caudal fin rays, marked fish can be easily detected without killing them.

Marking of fish by immersion requires a trade-off among cost, compound concentration, immersion time, salinity, mortality, growing condition and retention time to produce the best mark (Taylor et al., 2005). Also, it is important to assess which is the best life stage to perform the marking immersion, as chemical dyes form complexes with calcium and these were deposited in the bone as the fish grows (Eckmann, 2003).

To the best of our understanding, this study has shown for the first time that *Odontesthes bonariensis* juveniles can successfully retain ARS mark up to one year after the marking immersion. Future research must also be conducted working with earlier pejerrey stages, because only in the last year approximately 25 million of pejerrey embryos, larvae and juveniles were released in Argentinian water bodies by government hatcheries without any evaluation of the stock enhancement effectiveness. Even though the methods reported will require further research and testing, osmotic induction marking with ARS has considerable potential as a low cost, effective and practical technique for mass marking juvenile pejerrey.

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**Literature Cited**


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