The importance of thermoperiod for proper gametogenesis and successful egg and sperm production in meagre (Argyrosomus regius) breeders in aquaculture

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Abstract

We examined the effect of constant water temperature throughout the year on gametogenesis, spawning success and egg/sperm/embryo quality in meagre (Argyrosomus regius). Two broodstocks were exposed to simulated natural photoperiod, and either attenuated seasonal water temperature (SeasT, 16.4 to 19.6°C) or relatively constant water temperature (CoT, 19.4 ± 0.6°C). In the spawning period (May), 4 couples per group were induced to spawn with gonadotropin releasing hormone agonist (GnRHa). Gonadal stage of development, sperm quality parameters and plasma levels of sex steroids were evaluated prior to the GnRHa treatment. Spawning success and egg/sperm quality were examined over the following 4 weeks. Constant temperature did not prevent gametogenesis, but exposure to attenuated seasonal water temperature with the inclusion of winter low temperature was beneficial to both sexes. The mean (±SD) diameter of the largest vitellogenic oocytes prior to GnRHa administration was significantly higher in the SeasT compared to the CoT group (598 ± 27 vs 520 ± 17 µm). Testosterone plasma levels in the females were significantly higher in the SeasT group, but all other hormones were similar in both sexes. SeasT females spawned more consistently with higher relative fecundity, and 24-h embryo survival of the produced eggs. A more pronounced negative effect of constant water temperature was observed in males, since CoT males exhibited a spermiation index of 0 prior to GnRHa treatment, the latter clearly having a beneficial effect over the following 4 weeks. The study demonstrated that meagre do undergo gametogenesis to a significant extent even under constant water temperatures during the year. However, a seasonal thermal regime -even an attenuated one- was necessary for the proper development of the gametes, allowing for the successful spawning induction using the established GnRHa induction protocol.

Keywords: Argyrosomus regius; GnRHa; spawning; temperature; gametogenesis; sperm quality.

Introduction

Meagre (Argyrosomus regius) belongs to the Sciaenidae family and it is a species of interest for the diversification of Mediterranean aquaculture production (Quéméner et al., 2002; Stipa & Angelini, 2005). The rapid growth, excellent flesh taste and low-fat content of this fish led to its increasing production in the last decade (Poli et al., 2003; Cárdenas, 2010; Chatzifotis et al., 2010; Monfort, 2010; Grigorakis et al., 2011; Duncan et al., 2013). However, females do not reproduce readily in captivity, exhibiting dysfunctions with oocyte maturation, ovulation and spawning (Duncan et al., 2012; Mylonas et al., 2013a; Soares et al., 2015). As a result, significant research efforts have been invested in developing appropriate methods for controlling reproduction, based on the use of gonadotropin releasing hormone agonists (GnRHa) administered in the form of multiple injections or sustained release implants, resulting in the successful production of good quality eggs (Duncan et al., 2012; Mylonas et al., 2013b; Mylonas et al., 2013a; Fernández-Palacios et al., 2014; Mylonas et al., 2015; Soares et al., 2015; Duncan et al., 2018; Ramos-Júdez et al., 2019).

The annual reproductive cycle in fishes in the temperate or higher latitudes is controlled by environmental cues, mainly photoperiod and temperature, with photoperiod being the principal environmental regulator of the process of gametogenesis, and temperature acting as a secondary cue, being more important during final maturation and spawning (Bromage et al., 2001; Pankhurst & Porter, 2003; Falcon et al., 2010; Migaud et al., 2010; Wang et al., 2010; Zohar et al., 2010; Gordo & Carreras,
Although some fishes seem to have a specific thermal requirement in parallel with the photoperiod regime, others do not (Brown et al., 2006). This information is of a practical importance for the aquaculture industry, which usually simulates both photic and thermal conditions to achieve seasonal reproductive development and production of eggs (Bromage et al., 2001; Migaud et al., 2005; Povoa et al., 2011), as well as off-season spawning (Carrillo et al., 1989; Bromage & Roberts, 1995; Cerdà et al., 1995). Due to biosecurity requirements, aquaculture broodstocks are commonly maintained in recirculating aquaculture systems (RAS) using borehole seawater that is sterile, but has a relatively constant temperature throughout the year. As a result, significant costs for infrastructure and energy are incurred to heat and cool the water in order to provide the thermal cycling usually encountered in nature. If thermal cycling is found to be not so relevant for the gonadal development of a given species, then maintaining broodstocks under simulated photoperiods only, but under relatively constant borehole water temperatures would save in infrastructure and energy.

For example, in gilthead seabream (Sparus aurata) and common dentex (Dentex dentex), it was demonstrated that constant temperatures did not prevent full reproductive maturation or spawning of fish maintained in natural photoperiods (Pavlidis et al., 2001; Karamanlidis, 2017). A similar situation was also reported in salmonids when borehole water was used (Bromage et al., 2001). In an effort to reduce maintenance costs and also prolong the spawning season in meagre, previous studies have shown that females complete vitellogenesis and can spawn reliably after GnRHa administration, if exposed to a modified annual thermal cycle, with lower spring-summer-fall maxima than the natural temperature profile in the Mediterranean (Mylonas et al., 2013a; Mylonas et al., 2015). Similarly, males completed spermatogenesis and spermatogenesis under these conditions (Fakriadis et al., 2020). Then, spawning could be maintained for up to 17 weeks in response to weekly GnRHa injections to the females and a once-every-3-weeks GnRHa implantation to the males, under a water temperature of 19-20°C (Mylonas et al., 2016).

In the present study, we examined the hypothesis that meagre, as other fishes, may also undergo vitellogenesis and spermatogenesis in captivity when exposed solely to a natural photoperiod, while exposed to the relatively constant thermal regime provided by borehole seawater. Therefore, we evaluated a) the effect of a constant thermal regime compared to an attenuated seasonal thermal regime established earlier, in the process of gametogenesis, sperm production and quality, and plasma sex steroid levels during the expected spawning period; and b) the response to GnRHs administration in terms of spawning kinetics, and egg/sperm production and quality.

Materials and Methods

Broodstock maintenance

The experiment was held in the AQUALABS facilities of the Institute of Marine Biology, Biotechnology, and Aquaculture (IMBBC) of the Hellenic Centre for Marine Research (HCMR), Crete, Greece. From May 2018 until May 2019, two mixed-sex broodstocks (N = 14-16, mean body weight ± SD of 8.7 ± 0.5 kg) were maintained in 15-m³ rectangular tanks in recirculating aquaculture systems (RAS) under simulated natural photoperiod and supplied with borehole seawater, exposed to either a simulated “attenuated” seasonal temperature profile (SeasT group; 16.4 - 19.6°C) previously established for meagre maturation (Mylonas et al., 2015; Mylonas et al., 2016), or a constant temperature profile (CoT group; 19.4 ± 0.6°C) resulting from the use of a typical borehole seawater source used in commercial hatcheries in the Mediterranean (Fig. 1). Fish were fed five days per week to apparent satiation with commercial feed (Vitalis, Skretting S.A., Norway). Monitoring of temperature was done on a daily basis, while monitoring of pH, dissolved oxygen (DO, %), NH₃-N (mg l⁻¹) and NO₂-N (mg l⁻¹) was done once a week (CoT - pH = 7.54 ± 0.06; DO = 91 ± 4%; NH₃-N = 0.32 ± 0.22 mg l⁻¹; NO₂-N = 0.031 ± 0.026 mg l⁻¹ and SeasT- pH = 7.57 ± 0.06; DO = 92 ± 3%; NH₃-N = 0.32 ± 0.22 mg l⁻¹; NO₂-N = 0.031 ± 0.026 mg l⁻¹).

The experimental protocol was approved by the National Veterinary Service (PN 255356 - AΔΑ: 6Α17ΛΚ-ΠΛΩ). All procedures were conducted in accordance to the “Guidelines for the treatment of animals in behavioral research and teaching” (Anonymous, 1998), the Ethical justification for the use and treatment of fishes in research: an update (Metcalfe & Craig, 2011) and the “Directive 2010/63/EU of the European parliament and the council of 22 September 2010 on the protection of animals used for scientific purposes” (EU, 2010).

Broodstock reproductive evaluation and spawning induction

On the 6th of May 2019 both broodstocks were evaluated for reproductive stage. Fish were first tranquilized in their tanks after a 2-day starvation period with a dose of 0.01 ml l⁻¹ of clove oil, and then were transferred one-by-one for complete sedation to an anesthetic bath of a dose of 0.03 ml l⁻¹ clove oil (Mylonas et al., 2005). Ovarian biopsies were collected using an endometrial catheter (Pipele de Cornier, Laboratoire CCD, France) after applying gentle aspiration. A part of the biopsy was examined immediately as a wet mount under a compound
light microscope (x40 magnification) to evaluate the reproductive stage of the fish and to estimate the mean diameter of the most advanced vitellogenic oocytes (n = 10, at x100 magnification), and microphotographs were taken. The other part of the biopsy was stored for further histological processing (see below). Sperm production was evaluated after applying gentle abdominal pressure (stripping), using a subjective index (Spermiation Index, SI) developed earlier and used in meagre (Fakriadis et al., 2020) as follows: S0 = no milt released, S1 = only a drop of milt released after multiple stripping attempts, S2 = milt released after the first stripping attempt and, S3 = copious amount of sperm released with very little pressure. Milt samples (50–100 μl) were collected for sperm quality evaluation (when the spermiation index was ≥S2) after the genital pore was rinsed with clean water and blot dried, taking care to avoid contamination of samples with feces or urine. Milt was collected using a positive displacement pipette. The collected milt sample was stored in a 500-μl centrifuge tube, placed on ice and then transferred to a 4°C refrigerator until evaluation, immediately after the completion of the sampling.

Four females per group were selected at the expected spawning period (day 0) based on their ovarian maturity stage evaluation (mean oocyte diameter of the most advanced vitellogenic oocytes >550 μm), and were injected with desGly10, nAla6, Pro3-GnRH-NEthylamide (GnRHa, H-4070, Bachem, Switzerland) at an effective dose of 14.3 ± 0.2 μg GnRHa kg⁻¹ BW for the CoT females and 14.1 ± 0.2 μg GnRHa kg⁻¹ BW for the SeasT females (Mylonas et al., 2016). After treatment, females were placed individually in eight separate 5-m³ rectangular flow-through tanks supplied with aerated borehole seawater at 19.7 ± 0.4°C under simulated natural photoperiodic conditions. The four selected males from each thermal group (SeasT or CoT) were treated with a GnRHa implant at an effective dose of 48.0 ± 2.7 μg GnRHa kg⁻¹ BW (Mylonas et al., 2013b) constructed with [Ethylene-Vinyl Acetate]-copolymer and were also placed in the 5-m³ tanks with the females from their respective temperature profile group. This way four couples per temperature group were formed and induced to spawn. The spawning tanks were fitted with passive egg collectors supplied with water from the surface outflow. All GnRHa-treated fish were sampled weekly for the following four weeks. Females were injected with the same dose of a GnRHa injection at each of the following three weeks (days 7, 14, 21). Males were treated with the same GnRHa implant of the same dose once again at the beginning of the third week (day 14). At each sampling, biopsies and sperm were collected when it was possible, mean oocyte diameters of the largest vitellogenic oocytes were calculated and the spermiation index was determined. On day 28 the experiment was completed and all fish were transferred back to their original tanks.

**Sperm quality evaluation**

Sperm quality was assessed evaluating the following parameters: (a) sperm density (number of spermatozoa ml⁻¹ of milt), (b) survival of spermatozoa stored at 4°C (spermatozoa survival, days), (c) duration of forward motility of ≥ 5% of the spermatozoa in the field of view (mo-
tility duration, min). Estimation of density was performed in duplicates using a Neubauer haemocytometer under a compound light microscope (200x magnification, Nikon, Eclipse 50i, Japan), after diluting sperm 2121-fold with saline. Spermatozoa survival was estimated as follows: after collection, milt was stored at 4°C and examined every other day for spermatozoa motility until forward motility was less than 5%. Motility duration evaluation was conducted in duplicates after mixing 1 µl of milt and a drop of seawater (50 µl) on a microscope slide (400x magnification) and examining spermatozoa motility until forward motility was less than 5%.

Sperm quality was also assessed using computer-assisted sperm analysis (CASAS, ISAS, Spain). Milt samples were activated with seawater containing 2% bovine serum albumin (1:334) to obtain 200-300 cells in the field and placed in a counting chamber with a fixed depth (Sperm-Track 10). Evaluation with CASA was done immediately after milt collection, using a compound light microscope (Proiser UB 200i) under x200 magnification, on which a digital camera was mounted recording at 100 frames per second. Milt samples were evaluated in triplicates every 15 sec after activation until less than 5% of motile cells were present in the field of view. The analyzed parameters were curvilinear velocity (VCL), straight-line velocity (VSL), average path velocity (VAP) (µm sec⁻¹), motile cells, progressive cells (> 80% straightness - STR), rapid cells and STR (%). The software settings were adjusted to 1 to 90 µm for the head area. Spermatozoa were considered immotile, when showing a VCL < 10 µm sec⁻¹, whereas they were classified as rapid when VCL was higher than 100 µm sec⁻¹.

**Egg collection and embryo/larval survival evaluation**

Egg collectors were checked every day for the presence of eggs. After spawning, all eggs in the egg collector were taken with a dip-net and placed into a 10-l bucket. After whisking the bucket content, a sub-sample of 10-ml was used to estimate the fecundity (x 1000 eggs spawn⁻¹) and fertilization success (%) using a stereoscope with an egg-counter plate. To evaluate embryonic development, the method of Panini et al. (2001) was used by placing individual fertilized eggs in a 96-well microtiter (mct) plate (one egg per well). Daily monitoring of the mct-plates was carried out using a stereoscope, taking records of embryo viability 24 h after egg collection, hatching success, and larval survival 7 days after egg collection (yolk sac absorption). Embryo viability at 24 h after egg collection was estimated as the ratio of viable eggs (having live embryos) to the number of live eggs initially placed in the mct plate. Hatching success was calculated as the ratio of the hatched larvae to the number of viable embryos 24 h after egg collection. The survival 7 days after egg collection was estimated as the ratio of live larvae to the number of hatched larvae. Determination of survival of a specific developmental stage was done having as a denominator the live individuals from the previous stage to avoid masking effects and to have a more independent evaluation (Mylonas et al., 1992; Mylonas et al., 2004).

**Plasma sex steroid analysis**

Using slightly modified enzyme-linked immunosorbent assays (ELISAs) (Cuisset et al., 1994; Nash et al., 2000; Rodríguez et al., 2000), the plasma concentrations of testosterone (T), Estradiol (E2), 11-Ketotestosterone (11-KT) and 17,20β-P (17,20β-dihydroxy-4-pregnen-3-one) were quantified. Briefly, before running the samples in the ELISAs, plasma extraction was performed twice, by adding diethyl ether (3 ml) to plasma (300 µl) and vortexing vigorously the solution for 3 minutes (Vibramax 110, Heidolph, Germany). Once separated, the organic phase was transferred to new tubes in which it was dried under a nitrogen stream (React-vap III, Pierce, USA). Eventually, samples were reconstituted in 600 µl of assay buffer.

**Histological analysis**

Ovarian biopsies, obtained during the evaluation of the reproductive stage, were dehydrated in ethanol of gradually increasing concentration (70-96%), and were embedded in methacrylate resin (Technovit 7100®), Her-aeus Kulzer, Germany). Using a microtome (Leica RM 2245, Germany), 3 µm sections were cut and stained with Methylene Blue (Sigma, Germany)/Azure II (Sigma, Germany)/Basic Fuchsin (Polysciences, USA) according to the procedure of Bennett et al. (1976). A compound optical microscope (Nikon, Eclipse 50i, Japan), with a digital camera (Jenoptik progress C12 plus, Germany) mounted on top, was used to examine and photograph the sections.

**Statistical analysis**

Differences in mean oocyte diameters between the two thermal profile groups (SeasT vs CoT) were tested using either a t-test for the initial sampling of all fish, or using a two-way analysis of variance (ANOVA) for the GnRHα-treated selected females during the four weeks of the experiment. Differences in mean plasma sex steroid concentrations, relative fecundity, fertilization success and egg/larval survival between the two groups were tested using a t-test. Differences in spermiation index were tested using the nonparametric Mann-Whitney’s U test at the initial sampling of all fish, or the nonparametric Friedman’s test for the GnRHα-treated males during the four weeks spawning experiment, followed by Dunn’s post-hoc test. Statistical significance was set at P ≤ 0.05. Normality was tested using Shapiro-Wilk test. Statistical analysis was performed using GraphPad Prism version 8.4.3 for Mac (GraphPad Software, La Jolla California USA, www.graphpad.com). Results are presented as mean ± standard error of the mean (SEM) unless mentioned otherwise.
Results

Reproductive stage after different thermal regimes

The constant thermal regime did not prevent gametogenesis, but it affected negatively oocyte development, since the mean diameter of the largest vitellogenic oocytes of females from the CoT group was significantly lower (520 ± 17 μm) compared to the females from the SeasT group (598 ± 27 μm) (t-test, P = 0.02) (Fig. 2A). Further to a larger diameter, the ovaries of SeasT females were at a more advanced stage of development than those of CoT females, since CoT ovaries had a higher occurrence of oocytes in earlier developmental stages, such as primary (PO), cortical alveoli (CA) and early vitellogenic (eVg) oocytes (Fig. 3). In males, the influence of constant temperature on spermatogenesis was more pronounced, since no milt samples could be collected (Spermiation Index = S0) after stripping males from the CoT group (Fig. 2B). On the contrary, all males from the SeasT group where spermiating, though not all at the same extent, having a significantly higher mean spermiation index compared to the CoT group (Mann-Whitney’s U test, P = 0.002).

The plasma levels of T were also significantly higher in SeasT females (0.25 ± 0.01 ng ml⁻¹) compared to CoT females (0.17 ± 0.02 ng ml⁻¹) (t-test, P = 0.02), but no differences were observed in plasma levels of E₂ or 17,20β-P between the females from the two groups (Fig. 4A). No significant differences between the males from the two thermal groups were observed in the mean plasma levels of T, 11-KT or 17,20β-P (Fig. 4B).

Spawning induction after GnRHa injections

In the SeasT females, spawning commenced 2 days after the first GnRHa injection and a second spawn was obtained also the following day, with one female (No 4) spawning with a one-day delay compared to the rest (Fig. 5). On the contrary, not all females from the CoT group spawned after the first GnRHa injection, and spawning was very erratic. Fecundity was also many folds higher in the SeasT females at the first spawning induction.

In subsequent weekly ovarian evaluations, no significant differences in mean diameters of the largest vitellogenic oocytes were observed between the SeasT and CoT females, but a trend of lower values was noted in the CoT females (Fig. 6A). Spawning after the subsequent GnRHa injections was not as consistent in the SeasT females as it was after the first GnRHa injection, while in the CoT females it seemed to occur in more synchrony than before (Fig. 5), though individual fecundity was still markedly less than in the SeasT females. Overall relative fecundity of the SeasT females was significantly (t-test, P = 0.02) and 3-fold higher compared to the CoT group (Fig. 7A), while overall fertilization success was not significantly different (t-test, P = 0.31) between the two thermal groups (data not shown). Embryo survival 24-h after egg collection from SeasT females (77 ± %) was significantly higher (t-test, P = 0.003) compared to the CoT group (35 ± 21%), while no differences between the two thermal regimes were observed for hatching success and larval survival (Fig. 7B).

Sperm production and quality after GnRHa implantation

After the first GnRHa implantation, the males of the CoT group that had a Spermiation Index of S0 started progressively releasing sperm upon stripping, with 50% of males on day 7, 25% on day 14 and 75% on day 21 having a spermiation index of ≥ S2 (Fig. 6B). Still, males from the SeasT group were overall in higher spermatogenesis.
index compared to their CoT counterparts during all the sampling days (Friedman’s test, *P* = 0.002). Since day 0, all SeasT males released milt, and the percentage of males with an index equal to S3 steadily increased from 25% on day 7 to 75% on day 21. Statistical analysis on sperm quality parameters during the experiment was performed only for SeasT males (one-way ANOVA, *P* ≤ 0.05), due to the shortage of milt samples from CoT
males (Table 1), therefore no statistical comparison could be made between the two thermal regimes. For the SeasT sperm samples, no significant differences were observed in any of the evaluated parameters during the experiment. Sperm density ranged between 10 - 20 x 10⁹ szoa ml⁻¹, progressive motility between 43 - 69%, motility duration between 2.2 - 5.0 min, and survival under cold storage between 2.0 - 4.5 days (Table 1). The percentage of progressive and rapid cells evaluated by CASA was higher than 40% for the whole experiment. The VCL ranged between 145-210 μm sec⁻¹, VSL between 109-146 μm sec⁻¹, VAP between 115-170 μm sec⁻¹ and STR between 86 - 95% (Table 1).

Discussion

As meagre is one of the emerging species for the aquaculture development in the Mediterranean, the existing knowledge on environmental factors influencing its reproductive success needs to be improved. In order to reduce the energy cost of maintaining broodstocks in aquaculture facilities, previous studies exposing meagre to seasonal photoperiod, but relatively constant temperatures typical of borehole seawater (18-20°C) during the spring, summer and fall did not have any negative effect on gametogenesis in males or females in earlier studies (Mylonas et al., 2015). Based on these results, a very successful spawning induction protocol was proposed (Mylonas et al., 2016), which involved exposing meagre broodstock to low temperatures in the winter time, followed by a gradual increase to reach those typical of
spring (18-20°C) and then inducing spawning, with up to 17 weekly injections of GnRHa in females and Gn-RHa implantation every 3 weeks in males. This protocol resulted in the production of eggs of high fecundity and quality, as well as the necessary amount and quality of milt to ensure high fertilization success (Mylonas et al., 2015; Mylonas et al., 2016; Fakriadis et al., 2020). Extending these previous reports, the present study demonstrated that exposure throughout the year to relatively constant water temperatures typical of borehole water in the Mediterranean, did not prevent gametogenesis in either males or females. However, the results underlined the necessity of at least a winter thermal profile for the proper progression and completion of the gametogenic process. Therefore, maintaining broodstocks on seasonal photoperiod but constant borehole water temperature throughout the year cannot be utilized as a cost-effective method for aquaculture production for meagre.

The ability, however, of meagre - as well as other Mediterranean species (Pavlidis et al., 2001; Papadaki et al., 2018) - to undergo gametogenesis to a great extent responding only to photoperiod cues provides interesting information on the relative importance of temperature in controlling reproductive development in fishes in the temperate zone. At the onset of the spawning period, meagre females from the CoT group were in full vitellogenesis,

Fig. 6: Mean (± SEM) diameter of the largest vitellogenic oocytes from ovarian biopsies (A) and percentage (%) males at different spermiation index stages (B) of meagre (Argyrosomus regius) breeders exposed to a Constant (CoT, n = 4) or attenuated Seasonal (SeasT, n = 4) thermal regime, and selected for spawning induction. Arrows on the x-axis indicate the time of GnRHa administration (injection in females, implantation in males). Numbers inside the bars indicate the N value of the means. No significant differences were observed during the spawning induction experiment in oocyte diameters, but significant differences in spermiation index among thermal regime/sample time combinations were observed (Friedman’s test, Dunn’s post hoc, P≤0.05), indicated by different lowercase letters above the bars.

Fig. 7: A. Mean (± SEM) total relative fecundity (eggs kg⁻¹ female) of meagre (Argyrosomus regius) exposed to Constant (CoT) or attenuated Seasonal (SeasT) thermal regimes (t-test, P = 0.02). B. Mean (± SEM) percentage of embryo viability 24-h after egg collection (t-test, P = 0.003), hatching success and larval survival 7 days after egg collection. When present, lowercase letters above the means indicate significant differences between thermal regimes. Numbers inside the bars indicate the N value of the means.
Table 1. Mean (± SEM) values of sperm quality parameters of meagre (*Argyrosomus regius*) exposed to a Constant (CoT) or attenuated Seasonal (SeasT) thermal regime and sampled at the start of the spawning period and the onset of the GnRHa spawning induction experiment (day 0) and at weekly intervals thereafter. Fish were implanted with GnRHa at days 0 and 14, and were allowed to spawn with females induced with weekly injections of GnRHa. Sperm could be collected only from males with Spermiation Index of >S2 (see Materials and Methods).

| Day | n | Density¹ (10⁶ sperm ml⁻¹) | Motility¹ (%) | Duration¹ (min) | Survival¹ (days) | Progressive² (%) | Rapid² (%) | VCL² (µm s⁻¹) | VSL² (µm s⁻¹) | VAP² (µm s⁻¹) | STR² (%) |
|-----|---|--------------------------|--------------|----------------|-----------------|-----------------|------------|-------------|-------------|-------------|----------|
|     |   |                          |              |                |                 |                 |            |             |             |             |          |
| 0   | 0 | n/a                      | n/a          | n/a            | n/a             | n/a             | n/a        | n/a         | n/a         | n/a         | n/a      |
| 7   | 2 | 19.6 ± 3.9               | 62 ± 6.6     | 2.8 ± 0.5      | 2.5 ± 1.5       | 53 ± 1.0        | 58 ± 6.1   | 171 ± 24.7  | 132 ± 9.8   | 142 ± 16.6  | 94 ± 4.0 |
| 14  | 1 | 14.2 ± 10.0              | 94           | 5              | 4               | 69.8           | 92.2       | 210.3       | 145.6       | 170.5       | 85.8     |
| 21  | 2 | 10.2 ± 1.0               | 52 ± 41.8    | 3.4 ± 2.1      | 3.0 ± 1.0       | 43 ± 35.7       | 47 ± 44.5  | 145 ± 46.0  | 110 ± 46.2  | 117 ± 52.4  | 95 ± 3.2 |

1 The sperm parameters analyzed were density (Density, number of spermatozoa ml⁻¹ of milt), forward motility (Motility, %), duration of forward motility (Duration, min), and survival of spermatozoa stored at 4º C (Survival, days).

2 The CASA parameters analyzed were the percentage of spermatozoa with progressive movement (Progressive, %) and rapid movement (Rapid, %), curvilinear velocity (VCL), straight-line velocity (VSL), average path velocity (VAP)(µm sec⁻¹), and straightness of spermatozoa movement (STR, %).

3 Statistical analysis was performed only for data from the SeasT group, since very few males from the CoT group produced collectable milt (Spermiation Index ≤S1). No significant differences were observed among sampling times (one-way ANOVA, P ≤ 0.05).
but their ovaries had high occurrence of early developmental stage oocytes and the mean diameter of the largest vitellogenic oocytes had not reached the required minimum (550 µm) for the females to be successfully induced to spawn (Duncan et al., 2012; Mylonas et al., 2013a; Mylonas et al., 2016; Duncan et al., 2018). Similarly, males from the CoT group did not produce releasable milt at this time, but they did spawn and fertilized eggs 2 days after induction with GnRHa implants, suggesting that spermatogenesis was also completed, but not to the same extent as males exposed to the SeasT thermal regime. So, it appeared that the lack of winter temperature delayed or prevented the full progression of gametogenesis in meagre. Negative effects of inappropriate thermal conditions on gametogenesis and spawning success have been observed also in other teleosts, such as the common wolffish (Anarhichas lupus) (Tveiten & Johnsen, 1999), the Eurasian perch (Perca fluviatilis) (Migaud et al., 2002), the yellow perch (Perca flavescens) (Shewmon et al., 2007), the pikeperch (Sander lucioperca) (Zakęś, 2007) the European sea bass (Dicentrarchus labrax) (Carrillo et al., 1995) and the striped bass (Morone saxatilis) (Clark et al., 2005), which need to be exposed to a seasonal thermal regime simulating the winter season, in order to fully achieve vitellogenesis. Apparently, as shown in this and earlier studies of meagre (Mylonas et al., 2013b; Mylonas et al., 2013a; Mylonas et al., 2015; Mylonas et al., 2016), removing the summer from the thermal regime does not affect gametogenesis at all, but removing the winter has pronounced negative effects.

The fact that gametogenesis in meagre under a constant thermal regime was completed to a great extent in the present study, was also reflected in the lack of dramatic differences in plasma sex steroid hormones from the SeasT regime at the expected time of spawning, when GnRHa induction was implemented. The plasma levels measured were generally low in both sexes and thermal regimes, as observed previously in captivity (Mylonas et al., 2013a). Group-synchronous species, such as meagre, possess simultaneously oocytes at different developmental stages during the spawning season, and as a result plasma sex steroid levels do not exhibit dramatic differences among them and over time, as they are all required to support the process of vitellogenesis at the same time that oocyte maturation and ovulation take place. The slightly reduced T levels observed in the CoT females may be explained by a possible disruption of the sex steroid pathway caused by the warmer water during the gametogenesis period, as shown in studies with Atlantic salmon (Salmo salar) (Pankhurst et al., 2011), pikeperch (Heremlink et al., 2011) and Atlantic halibut (Hippoglossus hippoglossus) (van Nes & Andersen, 2006). Although it has been reported that both expression and activity of the gonadal aromatase (Cyp19a1a) gene may be inhibited by high-temperature exposure (Anderson et al., 2012; Elisio et al., 2012), in the present study plasma E2 did not differ between females of the two thermal regimes, again in support of the lack of an absolute requirement of a seasonal thermal regime in the reproductive cycle of meagre.

Similarly, in males the negative influence -but not prevention- of the constant thermal regime on spermatogenesis was not reflected by the similar levels of the measured sex steroid hormones. In other fishes, when a warmer than optimal temperature was applied for a prolonged period, a reduction or inhibition of spermatogenesis or spermatogenesis was observed (Vikingstad et al., 2016; Fenkes et al., 2017; Hani et al., 2019). The hormonal levels of males from both thermal groups were found to be similar to those reported in other studies of meagre in captivity (Mylonas et al., 2013a; Fakriadiis et al., 2020). This was surprising, since the failure to express milt from the CoT males prior to the GnRHa therapy was expected to be associated with lower plasma levels of 11-KT and/or 17,20β-P, as it has been amply demonstrated that these hormones are directly responsible for spermatogenesis and the testicular hydration associated with the increase in releasable milt during the spawning season (Schulz et al., 2010).

Perhaps expectedly, the negative consequences of exposure to the constant thermal regime during the winter season were not limited to the maturation stage of both sexes at the onset of the spawning period. In fact, the negative effects extended also during the spawning season in late spring (May-June) - when both treatments were exposed to the same temperature- compromising the final reproductive output. Previous studies on meagre demonstrated that using weekly GnRHa injections can improve vitellogenesis, favor the maturation of new batches of vitellogenic oocytes, and allow multiple spawning with eggs of high quality (Fernández-Palacios et al., 2014; Mylonas et al., 2016). Although after the first GnRHa injection the mean diameter of the largest vitellogenic oocytes increased in the CoT females and did not differ significantly from the SeasT females, there was a clear tendency for smaller oocytes throughout the spawning induction experiment. Similar results were found in the spiny chromis (Acanthochromis polyacanthus), where higher temperature decreased the ability of the fish to undergo proper vitellogenesis, and in the Atlantic salmon, where higher temperature led to a large imbalance in the size of the oocytes (Donelson et al., 2010; Vikingstad et al., 2016). Spawning kinetics and egg production of the SeasT females were more consistent to earlier studies (Mylonas et al., 2015; Mylonas et al., 2016; Duncan et al., 2018), although they differed from the expected two spawns per female on Days 2 and 3 after each GnRHa injection. On the contrary, CoT females spawned erratically, producing significantly and markedly less eggs, albeit of similar fertilization success to the SeasT females who experienced the low temperatures during the winter season. Additionally, although the mean fertilization success of fertilized spawns was not statistically different between the two thermal groups, in more than 50% of the spawns obtained from the CoT females, the eggs were not fertilized at all. Exposing fish during gametogenesis to a higher than optimal temperature had comparable effects on the relative fecundity of Atlantic salmon and river lamprey (Lampetra fluviatilis) exposed to 22 and 10°C, respectively (Pankhurst et al., 2011; Cejko et al., 2016). Therefore, although the repeated GnRHa injections were
successful in inducing further vitellogenesis to some extent in some CoT females, the initial reproductive condition of the females due to their exposure to a constant thermal regime over the whole year had a dramatic negative influence on their reproductive performance.

Embryo development and survival to hatching was significantly and drastically reduced in the eggs obtained from the constant thermal regime, while embryos from the SeasT group had a high survival, comparable to a previous study (Mylonas et al., 2016). Embryo mortality was also decreased significantly in common wolfish eggs obtained from females exposed to higher than optimal water temperature (Tveiten & Johnsen, 1999). Reduced survival at this stage could be related with altered intake of phospholipids and free fatty acids during vitellogenesis, as was observed in females of Arctic char (Salvelinus alpinus) and brown trout (Salmo trutta) exposed to higher temperatures (Jobling et al., 1995; Lahnsteiner & Leitner, 2013). Therefore, it is obvious that the overall egg production and quality from meagre females exposed to constant temperature was inadequate for profitable production in a commercial hatchery.

Regarding the males, the administration of two Gn-RHa implants improved steadily the spermiation index of individuals from the SeasT group, and milt could be collected and analyzed from almost all males at all sampling times. A similar, but much less pronounced effect was observed in GnRHa treated males from the CoT group. This demonstrates that the fish had the capacity to respond to the GnRHa stimulation -and the expected increase in plasma Luteinizing Hormone (LH) and sex steroid production (Mylonas et al., 2017)- but their initial stage of reproductive development was such that spermiation and releasable sperm production was reduced. A significantly lower sperm production was probably one of the reasons that many spawns from CoT females were not fertilized at all, as mentioned above.

Unfortunately, it was not possible to make any sperm quality comparisons between the two thermal groups due to the limited availability of releasable milt from CoT males. The milt obtained from the SeasT groups showed comparable or higher values with other studies with meagre under similar environmental conditions. For example, the percentage of motile spermatozoa from SeasT males was between 62 and 81%, which is similar to values between 80% and 90% (Mylonas et al., 2013a), 73% (Santos et al., 2018) or between 53 and 74% (Schiavorene et al., 2012). The VCL in the present study was higher than the of 140.90 ± 7.75 μm/s reported when testing sperm extenders for cryopreservation (Santos et al., 2018) or in vitro fertilization (Ramos-Júdez et al., 2019). In the latter study VAP values ~90 μm sec⁻¹ were considered satisfactory for high fertilization success, and in the present study even higher values were recorded. As has been reported for other species, such as the gillhead seabream (Sparus aurata) (Beirão et al., 2011), the percentage of motile cells and VCL are of critical importance for fertilization success. The sperm characteristics during the study did not change, suggesting that the GnRHa enhanced milt production without affecting sperm quality, either positively or negatively, which is commonly the effect of GnRHa implants in a number of teleosts (Mylonas et al., 2017).

In conclusion, although exposure throughout the year to relatively constant water temperatures did not prevent gametogenesis in either males or females, the present study demonstrated that a winter thermal profile is necessary for the proper progression and completion of the gametogenic process in meagre, in order to achieve the high reproductive performance required for successful industrial egg production. The results also point to the potential negative effects of global warming in the future reproduction of meagre in the wild.

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References

Anderson, K., Swanson, P., Pankhurst, N., King, H., Elizur, A., 2012. Effect of thermal challenge on plasma gonadotropin levels and ovarian steroidogenesis in female maiden and repeat spawning Tasmanian Atlantic salmon (Salmo salar). Aquaculture. 334-337, 205-212.

Anonymous, 1998. Guidelines for the treatment of animals in behavioural research and teaching. Animal Behaviour. 55, 251-257.

Beirão, J., Soares, F., Herráez, M.P., Dinis, M.T., Cabrita, E., 2011. Changes in Solea senegalensis sperm quality throughout the year. Animal Reproduction Science. 126, 122-129.

Bennett, H.S., Wyrick, A.D., Lee, S.W., McNeil, J.H., 1976. Science and art in preparing tissues embedded in plastic for light microscopy, with special reference to glycol methacrylate, glass knives and simple stains. Stain Technology. 51, 71-97.

Bromage, N., Porter, M., Randall, C., 2001. The environmental regulation of maturation in farmed finfish with special reference to the role of photoperiod and melatonin. Aquaculture. 197, 63-98.

Bromage, N.R., Roberts, R.J., 1995. Broodstock Management and Egg and Larval Quality. Blackwell Science, Oxford, 424 pp.

Brown, N.P., Shields, R.J., Bromage, N.R., 2006. The influence of water temperature on spawning patterns and egg quality in the Atlantic halibut (Hippoglossus hippoglossus L.). Aquaculture. 261, 993-1002.

Cárdenas, S., 2010. Crianza de la Corvina (Argyrosomus regius). Fundación Observatorio Español de Acuicultura, Madrid, Spain, 96 pp.

Carrillo, M., Bromage, N., Zanuy, S., Serrano, R., Prat, F., 1989. The effect of modifications in photoperiod on spawning time, ovarian development and egg quality in the sea bass (Dicentrarchus labrax L.). Aquaculture. 81, 351-365.
Carrillo, M., Zanuy, S., Prat, F., Cerda, J., Ramos, J. et al., 1995. Sea bass (Dicentrarchus labrax). p. 138-168 In: Broodstock Management and Egg and Larval Quality. Bromage, N.R., and Roberts, R.J. (Eds.). Blackwell Science, Oxford.

Cejko, B.I., Judycka, S., Juchno, D., Boron, A., Leska, A. et al., 2016. Hormonal treatment affects sperm motility in the spined loach (Cobitis taenia, Pisces, Cobitidae). Aquaculture Research, n/a-n/a.

Cerdà, J., Zanuy, S., Carrillo, M., Ramos, J., Serrano, R., 1995. and et al., 1995.

Chatzifotis, S., Panagiotidou, M., Papaioannou, N., Pavlidis, M., Nengas, I. et al., 2010. Effect of dietary lipid levels on growth, feed utilization, body composition and serum metabolites of meagre (Argyrosomus regius) juveniles. Aquaculture. 307, 65-70.

Clark, R.W., Henderson-Arzapalo, A., Sullivan, C.V., 2005. Disparate effects of constant and annually-cycling daylength and water temperature on reproductive maturation of striped bass (Morone saxatilis). Aquaculture. 249, 497-513.

Cuisset, B., Pradelles, P., Kime, D.E., Kühn, E.R., Babin, P. et al., 1994. Enzyme immunoassay for 11-ketotestosterone using acetylcholinesterase as label: application to the measurement of 11-ketotestosterone in plasma of Siberian sturgeon. Comparative Biochemistry and Physiology. 110C, 229-241.

Donelson, J.M., Munday, P.L., McCormick, M.I., Pankhurst, N.W., Pankhurst, P.M., 2010. Effects of elevated water temperature and food availability on the reproductive performance of a coral reef fish. Marine Ecology Progress Series. 401, 233-243.

Duncan, N.J., Estévez, A., Fernández-Palacios, H., Gairin, I., Hernández-Cruz, C.M. et al., 2013. Aquaculture production of meagre (Argyrosomus regius): hatchery techniques, ongrowing and market. p. 519-541 In: Advances in Aquaculture Hatchery Technology. Allan, G., and Burnell, G. (Eds.). Woodhead Publishing Limited, Cambridge, UK.

Duncan, N.J., Mylonas, C.C., Milton Sulton, E., Karamanlidis, D., Français Nogueira, M.C. et al., 2018. Paired spawning with male rotation of meagre Argyrosomus regius using GnRHa injections, as a method for producing multiple families for breeding selection programs. Aquaculture. 495, 506-512.

Duncan, N.J., Estévez, A., Porta, J., Carazo, I., Norambuena, F. et al., 2012. Reproductive development, GnRHa-induced spawning and egg quality of wild meagre (Argyrosomus regius) acclimatised to captivity. Fish Physiology and Biochemistry. 38, 1273-1286.

Durant, J.M., Hjermann, D.O., Ottersen, G., Stenseth, N.S., 2007. Climate and the match or mismatch between predator requirements and resource availability. Climate Research. 33, 271-283.

Elisio, M., Chalde, T., Miranda, L.A., 2012. Effects of short periods of warm water fluctuations on reproductive endocrine axis of the pejerrey (Odontesthes bonariensis) spawning. Comparative Biochemistry and Physiology Part A: Molecular & Integrative Physiology, 163(1), 47-55.

EU, 2010. Directive 2010/63/EU of the European parliament and the council of 22 September 2010 on the protection of animals used for scientific purposes. Official Journal of the European Union. L 276, 33-79.

Fakriadi, I., Zanatta, E.M., Fleck, R., Sena Mateo, D.L., Papadaki, M. et al., 2020. Endocrine regulation of long-term enhancement of spermatogenesis in meagre (Argyrosomus regius) with GnRHa controlled-delivery systems. Gen Comp Endocrinol. 297, 113549.

Falcon, J., Migaud, H., MuOoz-Cueto, J.A., Carrillo, M., 2010. Current knowledge on the melatonin system in teleost fish. General and Comparative Endocrinology. 165, 469-482.

Fenkes, M., Fitzpatrick, J.L., Ozolina, K., Shiels, H.A., Nudds, R.L., 2017. Sperm in hot water: direct and indirect thermal challenges interact to impact on brown trout sperm quality. Journal of Experimental Biology. 220 (14), 2513-2520.

Fernández-Palacios, H., Schuchardt, D., Roo, J., Izquierdo, M., Hernández-Cruz, C. et al., 2014. Dose dependent effect of a single GnRHa injection on the spawning of meagre (Argyrosomus regius) broodstock reared in captivity. Spanish Journal of Agricultural Research. 12, 1038-1048.

Gordo, A., Carreras, G., 2014. Determination of Temporal Spawning Patterns and Hatching Time in Response to Temperature of Atlantic Bluefin Tuna (Thunnus thynnus) in the Western Mediterranean. PLoS ONE. 9, e90691.

Grigorakis, K., Fountoulaki, E., Vasilaki, M., Mikos, I., Thanailides, C., 2011. Lipid quality and filleting yield of reared meagre (Argyrosomus regius). International Journal of Food Science and Technology. 46, 711-716.

Hani, Y.M.I., Turies, C., Palluel, O., Delahaut, L., Bado-Nilles, A. et al., 2019. Effects of a chronic exposure to different water temperatures and/or to an environmental cadmium concentration on the reproduction of the threespine stickleback (Gasterosteus aculeatus). Ecotoxicology and environmental safety 174, 48-57.

Hermelink, B., Wuerz, S., Trubirolha, A., Rennert, B., Klos, W. et al., 2011. Influence of temperature on puberty and maturation of pikeperch, Sander lucioperca. General and Comparative Endocrinology. 172 (2), 282-292.

Jobling, M., Johnsen, H., Pettersen, G.W., Henderson, R.J., 1995. Effect of temperature on reproductive development in Arctic char, Salvelinus alpinus (L.). Journal of Thermal Biology. 20.

Karamanlidis, D., 2017. Evolution of sex ratio and egg production in a population of gilthead seabream (Sparus aurata) over the course of five reproductive seasons, Biology Department. University of Crete, Crete, Greece, pp. 46.

Lahnsteiner, F., Leitner, S., 2013. Effect of temperature on gametogenesis and gamete quality in brown trout, Salmo trutta. Journal of Experimental Zoology. 319, 138-148.

McClell, J.D., Craig, J.F., 2011. Ethical justification for the use and treatment of fishes in research: an update. Journal of Fish Biology. 78, 393-394.

Migaud, H., Davie, A., Taylor, J.F., 2010. Current knowledge on the photoneuroendocrine regulation of reproduction in temperate fish species. Journal of Fish Biology, 76 (1), 27-68.

Migaud, H., Wang, N., Gardeur, J.N., Fontaine, P., 2005. Influence of photoperiod on reproductive performances in Eurasian perch Perca fluviatilis. Aquaculture. 252: 385-393.

Migaud, H., Fontaine, P., Sulisty, I., Kestemont, P., Gardeur,
J.N., 2002. Induction of out-of-season spawning in Eurasian perch Perca fluviatilis: effects of rates of cooling and cooling durations on female gametogenesis and spawning. *Aquaculture*. 205, 253-267.

Monfort, M.C., 2010. Present market situation and prospects of meagre (Argyrosomus regius), as an emerging species in Mediterranean aquaculture, *Studies and Reviews. General Fisheries Commission for the Mediterranean No*. 89: Food and Agriculture Organization of the United Nations, Roma, pp. 28.

Mylonas, C.C., Mitrizakis, N., Papadaki, M., Sigelaki, I., Mylonas, C.C., Cardinaletti, G., Sigelaki, I., Polzonetti-Mag, Mylonas, C.C., Hinshaw, J.M., Sullivan, C.V., 1992. Gn-RH-induced ovulation of brown trout (Salmo trutta) and its effects on egg quality. *Aquaculture*. 106, 379-392.

Mylonas, C.C., Duncan, N.J., Asturiano, J.F., 2017. Hormonal manipulations for the enhancement of sperm production in cultured fish and evaluation of sperm quality. *Aquaculture*. 472, 21-44.

Mylonas, C.C., Papadaki, M., Pavlidis, M., Divanach, P., 2004. Evaluation of egg production and quality in the Mediterranean red porgy (Pagrus pagrus) during two consecutive spawning seasons. *Aquaculture*. 232, 637-649.

Mylonas, C.C., Cardinaletti, G., Sigelaki, I., Polzonetti-Magni, A., 2005. Comparative efficacy of clove oil and 2-phenoxethanol as anesthetics in the aquaculture of European sea bass (Dicentrarchus labrax) and gilthead sea bream (Sparus aurata) at different temperatures. *Aquaculture*. 246, 467-481.

Mylonas, C.C., Mitrizakis, N., Papadaki, M., Sigelaki, I., 2013. Reproduction of hatchery-produced meagre Argyrosomus regius in captivity I. Description of the annual reproductive cycle. *Aquaculture*. 414-415, 309-317.

Mylonas, C.C., Mitrizakis, N., Castaldo, C.A., Cerviño, C.P., Papadaki, M., et al., 2013b. Reproduction of hatchery-produced meagre Argyrosomus regius in captivity II. Hormonal induction of spawning and monitoring of spawning kinetics, egg production and egg quality. *Aquaculture*. 414-415, 318-327.

Mylonas, C.C., Fatira, E., Karkut, P., Sigelaki, I., Papadaki, M., et al., 2015. Reproduction of hatchery-produced meagre Argyrosomus regius in captivity III. Comparison between Gn-RHa implants and injections on spawning kinetics and egg/larval performance parameters. *Aquaculture*. 448, 44-53.

Mylonas, C.C., Salone, S., Biglino, T., de Mello, P.H., Fakridis, I., et al., 2016. Enhancement of oogenesis/spermatogenesis in meagre Argyrosomus regius using a combination of temperature control and GnRHα treatments. *Aquaculture*. 464, 323-330.

Nash, J.P., Davail-Cuisset, B., Bhattacharyya, S., Suter, H.C., Le Menn, F., et al., 2000. An enzyme linked immunosorbent assay (ELISA) for testosterone, estradiol, and 17,20β-dihydroxy-4-pregnen-3-one using acetycholinesterase as tracer: application to measurement of diel patterns in rainbow trout (Oncorhynus mykiss). *Fish Physiology and Biochemistry*. 22, 355-363.

Panini, E., Mylonas, C.C., Zanuy, S., Carrillo, M., Ramos, J., et al., 2001. Incubation of embryos and larvae of marine fish using microtiter plates. *Aquaculture International*. 9, 189-196.

Pankhurst, N.W., Porter, M.J.R., 2003. Cold and dark or warm and light: variations on the theme of environmental control of reproduction. *Fish Physiology and Biochemistry*. 28, 385-389.

Pankhurst, N.W., King, H.R., Anderson, K., Elizur, A., Pankhurst, P.M., et al., 2011. Thermal impairment of reproduction is differentially expressed in maiden and repeat spawning Atlantic salmon. *Aquaculture*. 316, 77-87.

Papadaki, M., Mazzella, D., Santinelli, V., Fakridis, I., Sigelaki, I., et al., 2018. Hermaphroditism and reproductive function of hatchery-produced sharpsnout seabream (Diplodus puntazzo) under attenuated annual thermal cycles. *Aquaculture*. 482, 231-240.

Pavlidis, M., Keravec, L., Greenwood, L., Mourot, B., Scott, A.P., 2001. Reproductive performance of common dentex, Dentex dentex, broodstock held under different photoperiod and constant temperature conditions. *Fish Physiology and Biochemistry*. 25, 171-180.

Poli, B.M., Parisi, G., Zampacavallo, G., Iurzan, F., Mecatti, M., et al., 2003. Preliminary results on quality and quality changes in reared meagre (Argyrosomus regius); body and fillet traits and freshness changes in refrigerated commercial-size fish. *Aquaculture International*. 11, 301-311.

Povoa, I., Davie, A., Treasurer, J., Miguad, H., 2011. Broodstock spawning and larviculture of whiting (Merlangius merlangus L.) reared in captivity. *Aquaculture Research*. 42, 386-398.

Quéméné, L., Suquet, M., Mero, D., Gaignon, J.-L., 2002. Selection method of new candidates for finfish aquaculture: the case of the French Atlantic, the Channel and the North Sea coasts. *Aquatic Living Resources*. 15, 293-302.

Ramos-Júdez, S., González, W., Dutto, G., Mylonas, C.C., Faue, vel, C., et al., 2019. Gamete quality and management for in vitro fertilisation in meagre (Argyrosomus regius). *Aquaculture*. 509, 227-235.

Rodríguez, L., Begtashi, I., Zanuy, S., Carrillo, M., 2000. Development and validation of an enzyme immunoassay for testosterone: effects of photoperiod on plasma testosterone levels and gonadal development in male sea bass (Dicentrarchus labrax, L.) at puberty. *Fish Physiology and Biochemistry*. 23, 141-150.

Santos, M., Soares, F., Moreira, M., Beirão, J., 2018. Evaluation of different extenders for the cold storage of meagre (Argyrosomus regius) semen. *Aquaculture Research*. 49, 2723-2731.

Schiaivone, R., Zilli, L., Storelli, C., Vilella, S., 2012. Changes in hormonal profile, gonads and sperm quality of Argyrosomus regius (Pisces, Scianidae) during the first sexual differentiation and maturation. *Theriogenology*. 77, 888-898.

Schulz, R.W., de França, L.R., Lareyre, J.-J., LeGac, F., Chiari, ni-Garcia, H., et al., 2010. Spermatogenesis in fish. *General and Comparative Endocrinology*. 165, 390-411.

Shewmon, L.N., Godwin, J.R., Murashige, R.S., Daniels, H.V., 2007. Environmental Manipulation of Growth and Sexual Maturation in Yellow Perch, Perca flavescens. *Journal of the World Aquaculture Society*. 38, 383-394.

Soares, F., Ribeiro, L., Gamboa, M., Duarte, S., Mendes, A.C., et al., 2015. Comparative analysis on natural spawning of F1 meagre, Argyrosomus regius, with wild broodstock spawns in Portugal. *Fish Physiology and Biochemistry*. 41, 1509-1514.

Stipa, P., Angelini, M., 2005. Cultured Aquatic Species Infor-
Argyrosomus regius, FAO Fisheries Division, FAO, Rome.

Tveiten, H., Johnsen, H.K., 1999. Temperature experienced during vitellogenesis influences ovarian maturation and the timing of ovulation in common wolffish. *Journal of Fish Biology*. 55, 809-819.

van Nes, S., Andersen, O., 2006. Temperature effects on sex determination and ontogenetic gene expression of the aromatases cyp19a and cyp19b, and the estrogen receptors esr1 and esr2 in Atlantic halibut (*Hippoglossus hippoglossus*). *Molecular Reproduction and Development*, 73 (12), 1481-1490.

Vikingstad, E., Andersson, E., Hansen, T.J., Norberg, B., May-er, I. *et al.*, 2016. Effects of temperature on the final stages of sexual maturation in Atlantic salmon (*Salmo salar* L.). *Fish physiology and biochemistry*, 42 (3), 895-907.

Wang, N., Teletchea, F., Kestemont, P., Milla, S., Fontaine, P., 2010. Photothermal control of the reproductive cycle in temperate fishes. *Reviews in Aquaculture*. 2, 209-222.

Zakęś, Z., 2007. Out-of-season spawning of cultured pikeperch (*Sander lucioperca* (L.)). *Aquaculture Research*. 38, 1419-1427.

Zohar, Y., Muñoz-Cueto, J.A., Elizur, A., Kah, O., 2010. Neuroendocrinology of reproduction in teleost fish. *General and Comparative Endocrinology*. 165, 438-455.