Research Article© ①Effect of 17α-methyl testosterone, tamoxifen, and letrozole on
growth performance and sex reversal of rainbow trout
(Oncorhynchus mykiss)

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Abstract

Sex reversal and producing a monosex population is one of the most preferred growth promotion techniques of rainbow trout culture. Thus, the effects of 17α -methyltestosterone (2 mg/kg), tamoxifen (2, 20, and 100 mg/kg), letrozole (2, 20, and 100 mg/kg), and a combination of tamoxifen (100 mg/kg) and letrozole (100 mg/kg) on growth, masculinization and serum steroid content of rainbow trout were investigated in this research. Ethanol-dissolved chemicals were sprayed on commercial trout diet, and ethanol was evaporated overnight. The fish were fed the treated diet for two months and afterward, they were fed a commercial diet for six months. Results showed that 17α -methyltestosterone reversed the sex of rainbow trout effectively. The proportion of males, intersex, and females in this group were 76.67%, 10%, and 13.33%, respectively. In contrast with 17α -methyltestosterone, using tamoxifen and letrozole showed no effect on sex reversal of rainbow trout. Growth performance was adversely affected by all chemical-treated diets. However, compensatory growth occurred during first month after ending treatment period.

Keywords: Masculinization, Rainbow trout, Sex reversal

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Introduction

Sex determination is strongly related to genes in higher vertebrates: consequently, it is challenging to reverse the sex in these vertebrates. Conversely, in fish sex differentiation process can be easily affected by internal and external conditions (Piferrer et al., 1994). Thus, for sex reversal in fish. oral administration of steroid hormones is used by aquaculturists.

Fish sex is separated into genotype, determined by the genes responsible for gonads formation, and phenotype, the appearance of ovary or testis. The genotype of sex, related to genes, is determined at the time of fertilization, whereas, phenotype differentiation, the appearance of male or female sex organs, occurs in embryonic and larval development stages. This process is changed with administration of exogenous androgens and estrogens. Because monosex population of some cultured species, having a higher growth rate in comparison with mixed sexes, it is preferred by aquaculturists. For example, male tilapia, female common carp, and female rainbow trout are preferred for culture.

Monosex populations can be achieved either by directly treating fish with hormones to produce preferred sex or by indirect method with two steps; in the first step, embryos/ larvae/ juveniles are treated with androgens or estrogens to produce neo male $(XX \stackrel{\sim}{\triangleleft})$, neo female $(XY^{\bigcirc}_{+}, ZZ^{\bigcirc}_{+})$, or super male (YY^{\land}_{-}) populations. In the second step, these sex-reversed fish are used as breeders for producing all female or all-male populations (Hoga *et al.*, 2018). Similar to humans, sex determination system in rainbow trout is XX/XY type; thus, to obtain all-female population, XX males are produced by administration of androgens in the first step, and these sexreversed males are used for mating with normal females in the second step. Among androgens, 17α methyltestosterone (MT) is numerously used for producing neo male population of teleost fishes (Asadi Eidivand et al., 2022).

In addition, effects of chemicals such as tamoxifen (TM) and letrozole (LZ), with anti-estrogenic activities, on sex differentiation process and sex reversal have been studied in many species of vertebrates (Singh and Srivastava, 2015; Alijani et al., 2022). It is well demonstrated that TM and LZ suppress estradiol production in mammals (Bhatnagar et al., 2001; Simpson et al., 2002) and fish (Kwon et al., 2000; Sun et al., 2007; Singh et al., 2012). Based on acting mechanisms and pathways, there are two groups of anti-estrogenic chemicals. Chemicals, including TM, directly affect actions of estrogens because they inhibit the binding of estrogens estrogen receptors. to Additionally, some studies indicated that masculinization effect of TM is related to suppression of ovary-type P450arom mRNA expression (Kitano et al., 2007; Hulak et al., 2010). In contrast to the first group, chemicals like LZ act indirectly by inhibiting the excretion of estrogens. In this process, LZ inhibits aromatase activities, the enzyme that plays a crucial role in turnover of estrogens (Sun et al.,

2011). Consequently, ovarian development is suppressed, and sex differentiation leads to development of testis (Singh and Srivastava, 2015).

However, the efficacy of TM and LZ is yet to be evaluated in rainbow trout. Thus, effects of tamoxifen (a receptor blocker), letrozole (an aromatase inhibitor), and 17α -methyltestosterone on growth, masculinization, and serum steroid content of rainbow trout were investigated in this study.

Material and methods

Fish and rearing system

All-female larvae of rainbow trout before yolk sac uptake were transferred to Urmia University by nylon bags containing water and oxygen (1:3 ratios) from Rashekan rainbow trout hatchery. After two days of adaptation, the larvae were randomly distributed among 27 tanks, 125 individuals per 25-liter tank (three replicates per treatment). Water flow rate of 1.3-2.9 l/m and an aeration system were established for each tank in flow-through system. Water a temperature, pH, and dissolved oxygen values were 13.1-15.3°C, 7.85-8.04, and 7.96-8.22 mg/L, respectively. The fish were kept in these conditions for 2.5 months. Afterward, all fish were weighed, counted, and transferred to 200-liter tanks. Water flow rate of 5.7-6.4 l/m and an aeration system were established for each tank in a flowthrough system. Water temperature, pH, and dissolved oxygen values were 14.8-15.6°C, 7.94-8.08, and 7.11-7.84 mg/L, respectively (Table 1). Under these conditions, the fish were raised for 5.5 months.

Treatmen t	Temperature (°C)		рН		DO (mg/L)		water flow rate (l/m)	
	1-75 daf	76-240 daf	1-75 daf	76-240 daf	1-75 daf	76-240 daf	1-75 daf	76-240 daf
С	14.3±0.7	15.1±0.2	7.96±0.0	7.99±0.0	8.07±0.1	7.53±0.2	2.13±0.6	6.07±0.2
	5	6	8	5	1	5	3	3
М	14.1±0.7	15.3±0.2	7.94±0.0	7.98 ± 0.0	8.11±0.1	7.69 ± 0.3	2.06 ± 0.6	5.95 ± 0.2
	7	9	8	5	3	1	5	5
T2	14.2 ± 0.7	15.1±0.2	7.93±0.0	8.04 ± 0.0	8.05 ± 0.1	7.59 ± 0.2	2.10±0.6	6.01±0.3
	2	4	9	4	0	4	2	1
T20	14.3±0.7	15.2±0.2	7.96±0.0	8.00±0.0	8.08 ± 0.1	7.42±0.2	2.03±0.5	6.21±0.2
	0	2	8	4	1	0	9	0
T100	14.3±0.7	15.0 ± 0.2	7.97±0.0	7.98 ± 0.0	8.01±0.1	7.61±0.2	2.25±0.6	6.17±0.2
	3	8	7	5	4	7	3	6
L2	13.9±0.6	15.3±0.3	7.91±0.0	7.97±0.0	8.05 ± 0.1	7.54 ± 0.2	2.08 ± 0.6	6.13±0.2
	9	0	9	6	0	1	6	8
L20	14.1±0.7	15.1±0.2	7.96±0.0	8.04 ± 0.0	8.07 ± 0.1	7.52 ± 0.1	2.33±0.6	6.05 ± 0.2
	4	5	9	4	1	9	0	3
L100	14.3±0.7	15.2 ± 0.2	7.92±0.0	8.01 ± 0.0	8.13±0.0	7.46 ± 0.2	2.16±0.6	6.01±0.2
	4	7	8	5	9	5	2	2
T/L100	13.9±0.7	15.1±0.2	7.90±0.0	7.98±0.0	7.97±0.1	7.71±0.3	2.19±0.6	5.99 ± 0.2
	1	0	7	4	5	2	5	4

Table 1: Physico-chemical	factors of the rearing	water during ex	perimental period
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Data represents mean \pm standard deviation; treatments include C: control, M: 17 α -methyltestosterone, T2: 2 mg tamoxifen/kg feed, T20: 20 mg tamoxifen/kg feed, T100: 100 mg tamoxifen/kg feed, L2: 2 mg letrozole/kg feed, L20: 20 mg letrozole/kg feed, L100: 100 mg letrozole/kg feed, and T/L100: 100 mg tamoxifen and 100 mg letrozole/kg feed; daf: days after initiation of feeding.

Experimental feed

The alcohol evaporation method was used to prepare the chemical-treated diets (Navarro-Martín et al., 2009). 17amethyltestosterone (Aburaihan Pharmaceutical Co., Tehran, Iran), tamoxifen Hormone (Iran Pharmaceutical Co., Tehran, Iran), and letrozole (Soha Pharmaceutical Co., Tehran, Iran) after dissolving in 95% ethanol, were sprayed on the commercial trout diet (Faradaneh Co., Tehran, Iran). 95% ethanol was added to the diet of the control group without the chemicals. Alcohol of the diets was evaporated at room temperature overnight then stored at 4°C in refrigerator. There were nine treatments (with triplicates) containing 0 (control), 2 mg MT/kg feed (M), 2, 20, and 100 mg TM/kg feed (T2, T20, and T100, respectively), 2, 20, and 100 mg LZ/kg feed (L2, L20, and L100, respectively), and combination of 100 mg TM/kg feed and 100 mg LZ/kg feed (T/L100). The fish were fed with treated diets for two months after initiation of the exogenous feeding.

Sampling and measurements

A portable multimeter (WTW, Multi 3630 IDS, Weilheim, Germany) was record rearing used to water temperature, pH, and dissolved oxygen. At the end of the experiment, body weight of all fish from each tank were measured. Weight gain was calculated using the formula "Weight gain=Final weight-Initial weight". The formula **"SGR** $(\% day^{-1}) = 100 \times (\ln Wt - \ln W0)/t"$ was applied for calculating specific growth rate (SGR), where Wt and W0 are final and initial weight, respectively, and t is growth time in days (Gisbert and Williot, 1997). Growth retarding rate was calculated using the formula "GRR=(Wc-Wt)/Wc", where Wc and Wt represent mean body weights of control and experimental groups. respectively. For calculating GRR values, means of replicates were used (Shen et al., 2015). GSI values were measured using the formula "GSI=(gonad weight/body weight)×100".

For hormonal histological and analyses, 30 fish from each treatment were sampled randomly at the end of the experimental period. The fish were anesthetized with 500 mg/L carnation, and the blood of six specimens of each group was pooled. The samples allowed clotting in serum separator tubes for two hours at room temperature. The samples were centrifuged at 1000 rpm for 15 min at 4°C to obtain serum and stored at -80°C in freezer. Commercial kits (Monobind Inc., Lake Forest, USA) and microplate reader (BioTek, Synergy HT, USA) were used to estimate the serum 17β - estradiol and testosterone.

The fish were dissected after collecting blood. Gonads were weighed and fixed in Bouin solution. Standard histological techniques and microscopic analyses were used for sex determination of experimental fishes (Shen *et al.*, 2015).

Statistical analysis

SPSS 22 was used for all statistical analyses of data. Levine's Test was used for Homogeneity of variances testing,

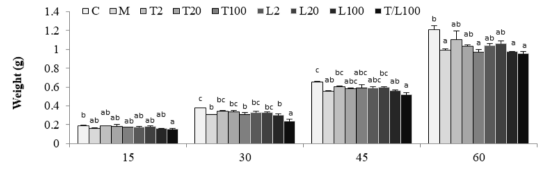
and Shapiro-Wilk test was used for normality of distribution testing. Oneway ANOVA was used to analyze the mean values, then Duncan's *post-hoc* Test was used to separate significantly different groups. All analyses were performed at α =0.05 (Irani and Noori, 2020).

Results

Growth and survival

Values of body weight in the control group were significantly more than

T/L100 group 15 days after initiation of the experiment (Fig. 1). In contrast, there were no significant difference among the control, M, T100, L100, and T/L100 groups 60 days after initiation of the experiment (at the end of the hormonal treatment period). Similar results were observed 15 days after ending the hormonal treatment period. There was no significant difference among all groups in the rest of the experimental period (Fig. 2).



Time (DAF)

Figure 1: Mean values of rainbow trout body weight during hormonal treatment period. Error bars show standard deviation. Different superscripts represent significant differences among treatments, one-way ANOVA, α<0.05, C: control, M: 17α-methyltestosterone, T2: 2 mg tamoxifen/kg feed, T20: 20 mg tamoxifen/kg feed, T100: 100 mg tamoxifen/kg feed, L2: 2 mg letrozole/kg feed, L20: 20 mg letrozole/kg feed, L100: 100 mg letrozole/kg feed, and T/L100: 100 mg tamoxifen and 100 mg letrozole/kg feed; DAF: days after initiation of feeding.

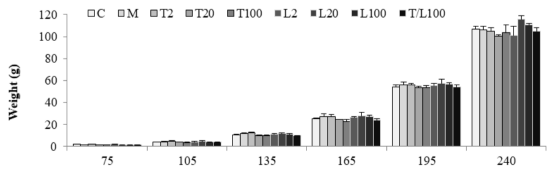
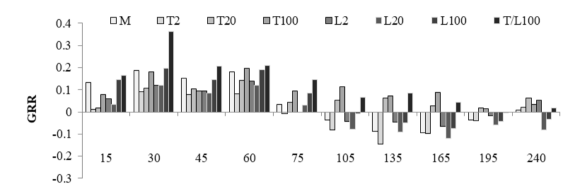




Figure 2: Mean values of rainbow trout body weight after hormonal treatment period. Error bars show standard deviation. C: control, M: 17α-methyltestosterone, T2: 2 mg tamoxifen/kg feed, T20: 20 mg tamoxifen/kg feed, T100: 100 mg tamoxifen/kg feed, L2: 2 mg letrozole/kg feed, L20: 20 mg letrozole/kg feed, L100: 100 mg letrozole/kg feed, and T/L100: 100 mg tamoxifen and 100 mg letrozole/kg feed; DAF: days after initiation of feeding.

Growth suppression, especially in M, T100, L100, and T/L100 groups, started 15 days after initiation of the hormonal treatment. Growth retarding rate increased during the first month, while it was almost constant during the second month (Fig. 3). Compensatory growth occurred 45 days after ending the hormonal treatments, as there were no significant difference in the growth performances among the groups from 105 DAF onwards.

SGR values were low on 15 DAF and increased afterward (Fig. 4). The values decreased gradually from 60 DAF onwards. There were significant differences between the control and T/L100 groups on 15 and 30 DAF.



Time (DAF)

Figure 3: Growth retarding rate of rainbow trout body weight during experimental period, M: 17αmethyltestosterone, T2: 2 mg tamoxifen/kg feed, T20: 20 mg tamoxifen/kg feed, T100: 100 mg tamoxifen/kg feed, L2: 2 mg letrozole/kg feed, L20: 20 mg letrozole/kg feed, L100: 100 mg letrozole/kg feed, and T/L100: 100 mg tamoxifen and 100 mg letrozole/kg feed; DAF: days after initiation of feeding.

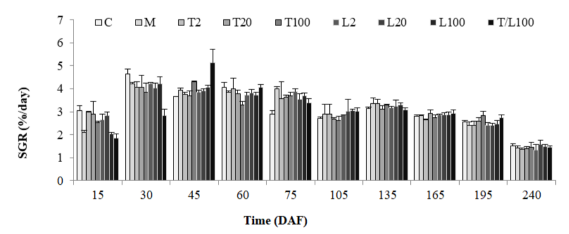


Figure 4: Mean SGR values of rainbow trout during the experimental period; error bars show standard deviation. C: control, M: 17α-methyltestosterone, T2: 2 mg tamoxifen/kg feed, T20: 20 mg tamoxifen/kg feed, T100: 100 mg tamoxifen/kg feed, L2: 2 mg letrozole/kg feed, L20: 20 mg letrozole/kg feed, L100: 100 mg letrozole/kg feed, and T/L100: 100 mg tamoxifen and 100 mg letrozole/kg feed; DAF: days after initiation of feeding.

With exception of 45 DAF, lowest FCR values were observed in control group

during the hormonal treatment period (Fig. 5). Values of M, T100, L100, and

T/L100 groups were higher than those of other groups during this period, whereas there were no significant difference among the experimental groups from 75 DAF onwards.

Sex reversal and GSI

Histological examination of gonads showed that sex reversal occurred only in the group treated with 17α methyltestosterone. Proportions of males, intersex, and females in this group were 76.67%, 10%, and 13.33%, respectively.

Mean GSI values in males of group M were significantly more than those in other groups. In contrast, the values in females of this group were significantly lower than those in other groups (Fig. 6). There was no significant difference among the rest of the groups.

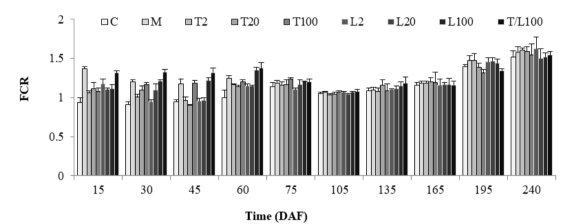


Figure 5: Mean FCR values of rainbow trout during the experimental period. Error bars show standard deviation. C: control, M: 17α-methyltestosterone, T2: 2 mg tamoxifen/kg feed, T20: 20 mg tamoxifen/kg feed, T100: 100 mg tamoxifen/kg feed, L2: 2 mg letrozole/kg feed, L20: 20 mg letrozole/kg feed, L100: 100 mg letrozole/kg feed, and T/L100: 100 mg tamoxifen and 100 mg letrozole/kg feed; DAF: days after initiation of feeding.

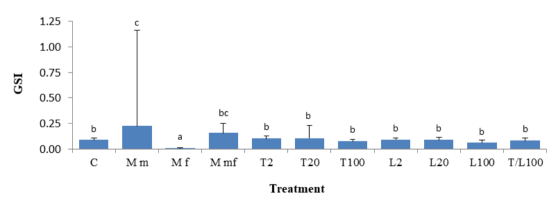


Figure 6: Mean GSI values of rainbow trout during the experimental period. Error bars show standard deviation. Different superscripts represent significant differences, one-way ANOVA, α<0.05; C: control, M: 17α-methyltestosterone, T2: 2 mg tamoxifen/kg feed, T20: 20 mg tamoxifen/kg feed, T100: 100 mg tamoxifen/kg feed, L2: 2 mg letrozole/kg feed, L20: 20 mg letrozole/kg feed, L100: 100 mg letrozole/kg feed, and T/L100: 100 mg tamoxifen and 100 mg letrozole/kg feed; DAF: days after initiation of feeding.

Steroids

Tamoxifen and letrozole caused no significant change in both testosterone and estradiol of experimental groups in comparison with the control group, whereas sex-reversed males in the 17α -methyltestosterone treated group showed significantly higher and lower

concentrations of testosterone and estradiol, respectively, compared to other groups. Testosterone concentration in intersex fish was also significantly higher than that in females of all groups (Fig. 7).

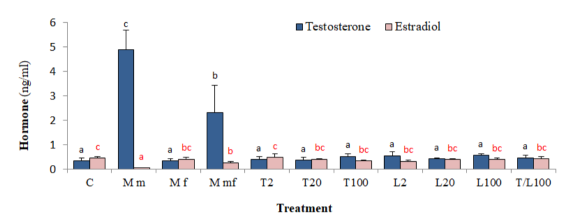


Figure 7: Testosterone and estradiol values of rainbow trout during the experimental period. Error bars show standard deviation. Different superscripts represent significant differences, one-way ANOVA, α<0.05; C: control, M: 17α-methyltestosterone, T2: 2 mg tamoxifen/kg feed, T20: 20 mg tamoxifen/kg feed, T100: 100 mg tamoxifen/kg feed, L2: 2 mg letrozole/kg feed, L20: 20 mg letrozole/kg feed, L100: 100 mg letrozole/kg feed, and T/L100: 100 mg tamoxifen and 100 mg letrozole/kg feed; DAF: days after initiation of feeding.

Discussion

Oral administration effects of 17amethyltestosterone (2mg/kg), tamoxifen, (2, 20, and 100 mg/kg), letrozole (2, 20, and 100 mg/kg), and a combination of tamoxifen (100 mg/kg) and letrozole (100 mg/kg) on growth, GSI, steroids, and sex reversal of rainbow trout were investigated in this study. The hormones have been used for sex reversal of several farmed species to produce a monosex population. There are several economic advantages to rear the most profitable gender, which possesses more growth characteristics (Taranger et al., 2010; Singh, 2013; 2018). 17α-Hoga al., et

methyltestosterone (MT) is numerously used for producing neomale population, and all-female population can be produced with these neomale breeders (Piferrer, 2001). In addition, chemicals like tamoxifen and letrozole, with antiestrogenic activities, have been used for sex reversal in mammals, birds, amphibians, reptiles, and fishes (Singh and Srivastava, 2015).

In the current study, growth performance was significantly affected by all administrated chemicals during twomonth treatment period, and suppression of growth occurred, especially at 17α methyltestosterone and high dosages of both tamoxifen and letrozole administrations. Compensatory growth occurred one month after the chemical treatment period. Similar findings were reported by Shen et al. (2015) for use of 17α -methyl testosterone in vellow fulvidraco. catfish Tachysurus In contrast, they achieved different results in use of letrozole, as low dose (20 mg/kg) of letrozole enhanced growth performance, while in higher doses (50 and 100 mg/kg) growth was not affected (Shen et al., 2015). Betancur et al. (2014) reported promotion of growth performance in low-dose of LZ (25 mg/kg) in treated red tilapia. This effect could be observed until one month after the treatment period. In their study there was no significant difference among growth values of the control, 17amethyltestosterone (60 mg/kg), and high dosage (100 mg/kg) of letrozole-treated groups.

These findings indicate significant differences in effects of hormones and other chemicals on growth performance of farmed fish, which could be due to species-specific characteristics. However, these adverse effects were not permanent, and usually compensatory growth occurred during the first month after ending chemical treatment period.

In this study, GSI values in tamoxifen and letrozole-treated groups were not affected, as there were no significant differences between the abovementioned groups and the control group. Similarly, sex differentiation of rainbow trout was not affected bv oral administration of tamoxifen and letrozole, whereas GSI values and sex differentiation of the 17α-methyl

testosterone-treated group were significantly influenced as GSI values of males were significantly more than those of the rest. While, in females of the group M, GSI values were significantly lower than those in other groups, which means that the ovary development was suppressed by 17α -methyl testosterone administration.

In contrast to our results, tamoxifen and letrozole showed significant effects on sex reversal of warm-water fishes, like common carp, catfish, and tilapia. A study of tamoxifen and letrozole influences on common carp (Cyprinus carpio) and Nile tilapia (Oreochromis niloticus) by Singh and Sirvastava (2015)indicated 100 that mg letrozole/kg feed made about 79% masculinization in common carp, and 88% masculinization in Nile tilapia, while the same dose of tamoxifen brought about 63% and 78% masculinization, respectively (Singh and Sirvastava 2015). Oral administration of letrozole (20, 50, and 100 mg/kg feed) produced about 75-83% males in yellow catfish (Tachysurus fulvidraco) that were significantly higher than male ratio in the control group, while 17α -methyl testosterone did not affect sex reversal of yellow catfish (Shen et al., 2015).

Thus, non-effectiveness of tamoxifen and letrozole on sex reversal in this study might be a result of rainbow trout physiology because this fish is considered a cold-water fish with different physiological requirements in comparison with warm-water fishes.

Monosex population of some cultured species like rainbow trout, having higher

growth rate in comparison with mixed sexes, is preferred by aquaculturists. Thus. 17α -methyltestosterone is commonly used for masculinization of this species. In the current study, this hormone effectively changed sex proportion of rainbow trout. In addition 17α -methyltestosterone, to use of chemicals with anti-estrogenic activities (tamoxifen and letrozole) for sex reversal, has been attempted in some ornamental and warm-water fishes. In contrast with 17α -methyltestosterone, tamoxifen and letrozole showed no effect on sex reversal of rainbow trout in this study. Growth performance was adversely affected by all chemicaltreated diets. However, compensatory growth retardation occurred during first month after ending the treatment period.

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References

- Alijani H., Amini Chermahini M., Zargham D. and Mohiseni M., 2022. The effect of Letrozole and Letrozole-Chitosan nanoparticles on masculinization of Rainbow trout larvae (Oncorhynchus mykiss). Journal **Fisheries** Iranian of Sciences. 21. 1383-1396. DOI:0.22092/ijfs.2022.128212
- Asadi Eidivand A., Mousavi S., Fatemi Tabatabei S., Zakeri M., Zanguee N., 2022. Determining the optimal time for artificial propagation

in Hilsa shad (*Tenualosa ilisha*) based on thyroid and steroid hormones levels. *Iranian Journal of Fisheries Sciences*, 21, 741-757. DOI: 10.22092/ijfs.2022.127015

- Betancur López, J.J., Quintero Velez, J.C., Ostos Alfonso, H., Barreiro-Sanchez, F. and Olivera-Angel, M., 2014. Effectiveness of the aromatase (P450 Arom) inhibitors Letrozole and Exemestane for masculinization of red tilapia (*Oreochromis* spp.). *Revista Colombiana de Ciencias Pecuarias*, 27(1), 47-53.
- Bhatnagar, A.S., Brodie, A.M.H., Long, B.J., Evans, D.B. and Miller, W.R., 2001. Intracellular aromatase its relevance and to the pharmacological of efficacy aromatase inhibitors. The Journal of Steroid Biochemistry and Molecular Biology, 76(1-5), 199-202. https://doi.org/10.1016/S0960-0760(01)00050-4.
- Gisbert, E. and Williot, P., 1997. Larval behavior and effect of the timing of initial feeding on growth and survival of Siberian sturgeon (*Acipenser baeri*) larvae under smallscale hatchery production. *Aquaculture*, 156(1-2), 63–76. https://doi.org/10.1016/S0044-8486(97)00086-0.
- Hoga, C.A., Almeida, F.L. and Reyes,
 F.G.R., 2018. A review on the use of hormones in fish farming: Analytical methods to determine their residues. *CYTA Journal of Food*, 16(1), 679-691.

https://doi.org/10.1080/19476337.20 18.1475423. Hulak, M., Psenicka, M., Gela, D., Rodina, M. and Linhart, O., 2010.
Morphological sex change upon treatment by endocrine modulators in meiogyngenetic tench (*Tinca tinca* L). Aquaculture Research, 41(2), 233-239.

https://doi.org/10.1111/j.1365-2109.2009.02325.x.

- Irani, A. and Noori, F., 2020. Comparative study on the biochemical factors and antioxidant enzymes of Rainbow trout eggs and larvae in a recirculating and flowthrough system. *Aquaculture*, 523, 735202.
- Kitano, T., Yoshinaga, N., Shiraishi, E., Koyanagi, T. and Abe, S.I., 2007. Tamoxifen induces masculinization of genetic females and regulates P450 aromatase and Müllerian inhibiting substance expression in mRNA Japanese flounder (Paralichthys olivaceus). Molecular Reproduction and 1171-1177. 74(9), Development, https://doi.org/10.1002/mrd.20603.
- Kwon, J.Y., Haghpanah, V., Kogson-Hurtado, L.M., McAndrew, B.J. Penman, **D.J.**, 2000. and Masculinization of genetic female Nile tilapia (O. niloticus) by dietary administration of an aromatase inhibitor during sexual differentiation. Journal of Experimental Zoology, 287, 46-53.
- Navarro-Martin, L., Blázquez, M. and Piferrer, F., 2009. Masculinization of the European sea bass (*Dicentrarchus labrax*) by treatment with an androgen or aromatase

inhibitor involves different gene expression and has distinct lasting effects on maturation. *General and Comparative Endocrinology*, 160(1), 3–11.

https://doi.org/10.1016/j.ygcen.2008. 10.012.

- Piferrer, F., Benfey, T.J. and Donaldson, E.M., 1994. Gonadal morphology of normal and sexreversed triploid and gynogenetic diploid coho salmon (*Oncorhynchus kisutch*). Journal of Fish Biology, 45(4), 541–553. https://doi.org/10.1111/j.1095-8649.1994.tb00923.x.
- Piferrer, F., 2001. Endocrine sex control strategies for the feminization of teleost fish. *Aquaculture*, 197(1-4), 229–281. https:/doi.org/10.1016/S0044-

8486(01)00589-0.

- Shen, Z.G., Fan, Q.X., Yang, W., Zhang, Y.L. and Wang, H.P., 2015. Effects of 17α-methyltestosterone and aromatase inhibitor letrozole on sex reversal, gonadal structure, and growth in yellow catfish *Pelteobagrus fulvidraco. The Biological Bulletin*, 228(2), 108–117. https://doi.org/10.1086/BBLv228n2p 108.
- Simpson, E.R., Clyne, C., Rubin, G., Boon, W.C., Robertson, K., Britt, K., Speed, C. and Jones M., 2002. Aromatase-a brief overview. Annals of Reviews in Physiology, 64, 93-127. https://doi.org/10.1146/annurev.phys iol.64.081601.142703.
- Singh, A.K. and Srivastava, P.P., 2015. A CYP19 based sex

determination and monosex production in aquaculture species *Oreochromis niloticus* L. and a Cyprinid *Cyprinus carpio* L. *Fisheries and Aquaculture Journal*, 6(1), 112. https://doi.org/10.4172/2150-3508.1000112.

- Singh, R., Singh, A.K. and Tripathi, M., 2012. Effect of a non steroidal tamoxifen on the gonad and sex differentiation in Nile tilapia *Oreochromis niloticus. Journal of Environmental Biology*, 33(4), 799-803.
- Singh, A.K., 2013. Introduction of modern endocrine techniques for the production of monosex population of fishes. *General and Comparative Endocrinology*, 181, 146–155. https://doi.org/10.1016/j.ygcen.2012. 08.027.
- Sun, L., Zha, J., Spear, P.A. and Wang, Z., 2007. Tamoxifen effects on the early life stages and reproduction of Japanese medaka

(*Oryzias latipes*). Environmental Toxicology and Pharmacology, 24(**1**), 23-29. https://doi.org/10.1016/j.etap.2007.0 1.003.

- Sun, L., Shao, X., Chi, J., Hu, X., Jin, Y. and Fu, Z., 2011. Transcriptional responses in the brain, liver and gonad of Japanese ricefish (*Oryzias latipes*) exposed to two antiestrogens. *Comparative Biochemistry and Physiology, Part C: Toxicology and Pharmacology*, 153(4), 392–401. https://doi.org/10.1016/j.cbpc.2011.0 1.003.
- Taranger, G.L., Carrillo, M., Schulz,
 R.W., Fontaine, P., Zanuy, S.,
 Felip, A., Weltzien, F.A., Dufour,
 S., Karlsen, Ø., Norberg, B.,
 Andersson, E. and Hansen, T.,
 2010. Control of puberty in farmed
 fish. *General and Comparative Endocrinology*, 165(3), 483-515.
 https://doi.org/10.1016/j.ygcen.2009.
 05.004.