Effect of European Seabass (*Dicentrarchus Labrax*) Fry Source (Wild, NIOF and GAFRD K21) on its Growth Performance and Physiology

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The present study aimed to evaluate the impacts of different sources on the European seabass (*D. labrax*) fry growth and physiology. One thousand and five hundreds fry from 3 sources were transferred to the lab., the 1st fry group was collected from the wild habitat in El- Meadya region, El- Behira Gov. from the wild fry capture fisheries. The 2nd fry group was 94 (dph) day post hatching produced in the Marine NIOF Hatchery, El- Anfoushy from induced spawning and the 3rd group fry was 94 dph produced in the Marine GAFRD Hatchery from induced spawning. Fry samples were randomly collected to statistically determine the length, weight growth and physiology performances. The comparative effects of European seabass (*D. labrax*) fry sources (wild and two hatcheries) and their impacts on length and weight growth parameters, total length, standard length, body width, total weight, gutted weight, hepatosomatic and vesrosomatic indexes. The wild collected fry achieved the best significant (P < 0.05) results in all this growth parameters while the NIOF marine hatchery produced fry achieved the best significant (P < 0.05) results in all these physiology parameters. In conclusion, this study explained why fish farmers prefer wild collected European seabass (*D. labrax*) fry than hatchery produced fry. Egypt marine hatcheries need more efforts and technology transfer to help marine aquaculture and fisheries development, that need ranching and restocking with hatchery produced fry.

Keywords: *Dicentrarchus labrax*, fry, wild, hatchery, growth, hepatosomatic and vesrosomatic indexes.

Introduction

Egypt is the tenth world aquaculture producer and also the first Mediterranean Sea, Arab and African aquaculture producer [1]. Egypt marine aquaculture still depending on fry collected from natural resources (95.437 million fry, only 0.797 million fry of them recorded as Seabreams and Seabass fry). Although Egypt had 6 hatcheries producing 8,906 million fry, only 3,500 million were Seabass fry in 2015 [2]. Not enough or misbalanced or low quality live feeds and microdiets affecting negatively marine hatcheries larval productivity and quality [3, 4]. In 2013, Egypt mass mortalities in Kafr El-Sheik and El- Behira governorates tilapia fish farms which diagnosed as *A. hydrophilla* outbreak and also, in Maryut Valley, Alexandria governorate in European seabass, gilthead seabream and meagre farms with different reasons as high temperatures and low oxygen stress in seabass and meagre, 100 tons equals 7 million L. E. losses [5] and because of *Vibrio* sp. outbreak in gilthead sebream after seabass and seabream wild collected fry were transported to these farms [5]. Therefore, the aim of the study were to evaluate the impacts of different resources on the European seabass (*D. labrax*) fry growth and physiology.

Global fish production from capture and aquaculture supplied about 167.2 million tons in 2014, with 73.8 million tons from aquaculture (47.1 from inland and 26.7 million tons from marine aquaculture), providing an apparent per capita supply of 20.1 kg [1]. China is the world leading aquaculture producer and had produced 58.16% of the world aquaculture production. FAO [1] reported that in 2014, the composition of world aquaculture production was; freshwater fishes (49.9 million tons), molluscs (16.1 million tons) and crustaceans (6.9 million tons). The number of species recorded in FAO aquaculture production.
statistics increased to 580 species and/or species groups farmed around the world, including those once farmed in the past, had been registered with production data by FAO. These species items include 362 finfishes (including hybrids), 104 molluscs, 62 crustaceans, 6 frogs and reptiles, 9 aquatic invertebrates, and 37 aquatic plants [1]. Turkey is the major producer of European seabass (Dicentrarchus labrax), Greece dominates the production of gilt-head seabream (Sparus aurata), while Egypt is the main producer of mullets (Mugilidae spp) in the Mediterranean [6].

Materials and Methods

Experimental fish, duration and location

The present study was conducted using one thousand and five hundreds fry from 3 sources and transferred to the Fish Reproduction and Spawning Lab., the 1st group were 500 hundreds fry collected from the wild habitat in El-Maadyia region, El- Behira Governorate from local fry trader in the seabass fry collecting season from the wild fry capture fisheries., the 2nd group were 500 hundreds of 94 dph fry produced in the Marine General Authority for Fish Resources Development (GAFRD) Hatchery, Km 21 Abou Talat, Alexandria, Egypt from induced spawning of 1600 gm female and 1000 gm male broodstock managed and three generations selected in the hatchery and the 3rd group were 500 hundreds of 94 (dph) day post hatching fry produced in the Marine NIOF Hatchery, El- Anfoushy, Alexandria from induced spawning of 1250 gm female and 600 gm male broodstock collected from Maryut valley fish farm twelve ripe fish of European seabass broodstock. Fry samples collected during the spawning season of 2014/2015 from the wild and both hatcheries.

Induced spawning using hormonal manipulation of the hatchery produced fry

In both hatcheries, Induced spawning were done using luteinizing releasing hormone analogue (LHRHa) and human chronic*-gonadotropin hormone (HCG). After broodstock acclimatization, resting and feeding then checked for ripeness. Ripe females injected in the abdominal cavity using two doses, the priming dose were 10 µg LHRHa kg⁻¹ and the second dose were 200 IU HCG kg⁻¹ releasing dose. Ripe males were injected using the releasing dose only using 200 IU HCG kg⁻¹. The post larvae rearing protocol was almost the same in both hatcheries as shown by Salem [12, 3] and Hebalah [13].

Growth criterias

Growth of post larvae were measured carefully as total length were to nearest 0.1 mm and total weight to nearest 0.001 mg to calculate total weight gain, average daily gain, specific growth rate in weight according to Castell [14], Condition factor were according to Yildiz et al. [15] and average daily gain or increase in length were according to Garber [16] and specific growth rate in length were according to Nour [17].

Weight growth parameters

Total weight gain (WG g/fish)

\[ \text{Total Weight Gain (WG)} = Wt - W0 \]

Where: W0: the initial fish weight at the start of the experiment.
Wt: the final fish weight at the end of the experiment.

Weight average daily gain (WADG mg/day)

\[ \text{ADG (mg)} = \frac{Wt - W0}{n} \]

Where: W0: the initial fish weight at the start of the experiment.
Wt: the final fish weight at the end of the experiment.
n: the duration period of the experiment in days.

Weight specific growth rate in weight (WSGR%)

\[ \text{SGR} = \frac{(\ln Wt - \ln W0)}{n} \times 100 \]


Weight gain % (WG%)

\[ \text{Weight Gain %} = \frac{\text{WG}}{W0} \times 100 \]

Length growth parameters

Total length gain (LG cm/fish)

\[ \text{Total length gain (LG)} = Lt - L0 \]

Where: L0: the initial fish length at the start of the experiment.
Lt: the final fish length at the end of the experiment.

Length average daily gain (LADG mm/day)

\[ \text{LADG (mm)} = \frac{Lt - L0}{n} \]

Where: L0: the initial fish length at the start of the experiment.
Lt: the final fish length at the end of the experiment.
n: the duration period of the experiment in days.

Length specific growth rate % (LSGR %)

\[ \text{LSGR %} = \frac{(\ln FL - \ln IL)}{n} \times 100 \]

IL: the initial fish length at the start of the experiment.
FL: the final fish length at the end of the experiment.
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**Length gain % (LG %)**

\[ \text{Length Gain} \% = \frac{\text{LG}}{\text{L} \times 100} \]

**Physiology parameters**

**Condition factor (K)**

\[ K = \left( \frac{\text{Wt (gm)}}{\text{L (cm)}} \right) \times 100 \]

Where: Wt: the total body weight.
L: the total body length.

**Hepatosomatic index (HSI)**

\[ \text{HSI} = \frac{100 \times (\text{liver weight (gm)} \times \text{body weight(gm)}^{-1})}{1} \]

**Viscerosomatic index (VSI)**

\[ \text{VSI} = \frac{100 \times (\text{carcass weight (gm)} \times \text{body weight(gm)}^{-1})}{1} \]

**Measurement of water quality**

Water quality measurements were done using AquaReed®. Water quality portable electric device were done in water quality and the electrod device was gently putted in the bottom of the experimental tanks away from the airstone and water quality mesurments are Dissolved Oxygen % (DO%), pH, Conductivity and Total Dissolved Solids (TDS) while Salinity as part per thousand (ppt) were measured using the referectometer.

**Counting the bacterial groups**

Microbiological measurements were done in randomly collected rotifer enrichment environment water samples in the water source and algal source for greenwater for Colony Forming unit (CFU) of *Vibrio* and *Bacillus* were done in the Microbiology Lab., Marine Environment Division, NIOF., Serial dilutions from $10^{-2}$ through $10^{-6}$ were made using filtered sterilized sea water. A portion (0.1 ml) from each appropriately diluted sample was used to inoculate plates prepared with seawater agar for total bacterial counting. *Bacillus* plates were incubated at 30°C for 72 h and *Vibrio* plates were incubated at 38°C for 72 h. Plates of two selective media were inoculated with 1 ml of appropriately dilution sample for counting the different bacterial groups: *Vibrio* and *Bacillus* bacterial counts.

**Chemicals and media**

All chemicals used for biochemical tests and extraction of antimicrobial activity were of pure grade and purchased from Sigma chemicals, USA. Ingredients of media were all of analytical grade and obtained from recognized chemical suppliers (mainly Oxoid Co.). Media used throughout the work are described below. The composition is given in gl^{-1}. The pH value of the media was adjusted to 7.5 prior to sterilization. Autoclaving was occurred at 121°C for 15 min.

Media used for isolation and enumeration of the different bacterial groups Sea water agar

A selective medium by [18] used for determining Bacillus count bacteria. Peptone, 5, ferric phosphate, 0.1, agar, 15, sea water 1L.

**Thiosulfate citrate bile salt sucrose agar (TCBS)**

A selective medium by [19] used for isolating Vibrio spp.: Yeast extract, 5, peptone, 10, sodium thiosulfate, 10, sod-citrate, 10, Ox bile, 8, sucrose, 20, sodium chloride, 10, ferric chloride, 1, Bromothymol blue, 0.04, thymol blue, 0.04 and agar, 14.

**Statistical analysis**

Statistical analysis were performed using analysis of variance (ANOVA), differences among means were considered significant at $p < 0.05$ multiple range of post hoc comparisons were performed using Duncan’s multiple range test and critical ranges to resolve the differences among the means of replication according to Duncan [20] using basic statistics, STATISTICA® software for Windows [21].

**Results and Discussion**

The effects of European seabass (*D. labrax*) fry sources (wild and two hatcheries) on length and weight growth parameters, total length, standard length, body width, total weight and gutted weight in gm (Table 1.). The wild collected fry achieved the best significant ($p < 0.05$) results in all this growth parameters. Morphometric relationship between length and weight (LWR) is of great importance in fish biology assessments. According to Lagler [22], Length-weight relationship leads itself to comparison of individuals within and between different populations. LWR can be used to assess the well-being of individuals and to determine the possible differences between separate unit stocks of the same species [23, 24].

The European seabass (*D. labrax*) fry sources (wild and two hatcheries) impacts on length growth performance parameters, total length, length gain, average daily gain in mm, specific growth rate in %/day and length gain % growth parameters (Table 2.). The wild collected fry achieved the best significant ($p < 0.05$) results in
all this length growth performance parameters. Salem and Ahmed [12], Salem et al. [3], Salem [5], recorded that results of larvae length gains showed that better significantly results achieved by GMW+MP and GMW+S than GMW in most larvae body length and length gains parameters of 7dph, 14dph, 21dph, 25dph, 35dph and 40dph, showed potential application of them in these critical stages and these may be due to many factors and mechanisms such as positive effects of these treatments as supported by Arığ [25] who showed enhanced growth of gilthead seabream larvae using Bacillus sp. bacteria enriched rotifers and artemia.

The comparative effects of European seabass (D. labrax) fry sources (wild and two hatcheries) on weight growth performance parameters, total weight, weight gain in gm, average daily gain in mg, specific growth rate in percentage/day and weight gain % growth parameters (Table 2.). The wild collected fry achieved the best significant (P < 0.05) results in all these weight growth performance parameters. Salem [12, 3] recorded that larvae body weight better results achieved by GMW+MP and GMW+S than GMW, El-Dakar [26] studied various inclusion levels of a feed additive that contains probiotic bacteria, digestive enzymes and prebiotic spices to evaluate the effect on siganid survival, growth, feed conversion, nutrient utilization, body composition and economics and showed that rabbitfish offered the control diet exhibited lower growth and feed utilization than all experimental treatments. There was no effect of probiotic inclusion level on survival but growth was better at all inclusion levels than in the control. No significant differences in growth were observed among fish groups fed various levels of the probiotic.

### TABLE 1. Length, weight, condition factors, hepatosomatic and viscososomatic indexes of wild collected and hatcheries produced European seabass (D. labrax) fry.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Wild</th>
<th>GAFRD K21</th>
<th>NIOF</th>
</tr>
</thead>
<tbody>
<tr>
<td>TL (cm)</td>
<td>7.320a</td>
<td>3.760b</td>
<td>3.260b</td>
</tr>
<tr>
<td>± 0.657</td>
<td>± 0.152</td>
<td>± 0.207</td>
<td></td>
</tr>
<tr>
<td>STL (cm)</td>
<td>6.240b</td>
<td>3.220b</td>
<td>2.680b</td>
</tr>
<tr>
<td>± 0.639</td>
<td>± 0.110</td>
<td>± 0.148</td>
<td></td>
</tr>
<tr>
<td>BW (cm)</td>
<td>1.380b</td>
<td>0.560b</td>
<td>0.580b</td>
</tr>
<tr>
<td>± 0.148</td>
<td>± 0.055</td>
<td>± 0.045</td>
<td></td>
</tr>
<tr>
<td>TW (gm)</td>
<td>4.048b</td>
<td>0.483b</td>
<td>0.399b</td>
</tr>
<tr>
<td>± 1.090</td>
<td>± 0.030</td>
<td>± 0.067</td>
<td></td>
</tr>
<tr>
<td>GW (gm)</td>
<td>3.386b</td>
<td>0.355b</td>
<td>0.246b</td>
</tr>
<tr>
<td>± 0.919</td>
<td>± 0.030</td>
<td>± 0.046</td>
<td></td>
</tr>
<tr>
<td>KTL</td>
<td>1.013b</td>
<td>0.910b</td>
<td>1.155b</td>
</tr>
<tr>
<td>± 0.034</td>
<td>± 0.063</td>
<td>± 0.188</td>
<td></td>
</tr>
<tr>
<td>KSTL</td>
<td>1.650b</td>
<td>1.449b</td>
<td>2.068b</td>
</tr>
<tr>
<td>± 0.227</td>
<td>± 0.112</td>
<td>± 0.258</td>
<td></td>
</tr>
<tr>
<td>HSI</td>
<td>1.168b</td>
<td>1.610b</td>
<td>1.932b</td>
</tr>
<tr>
<td>± 0.215</td>
<td>± 0.202</td>
<td>± 0.182</td>
<td></td>
</tr>
<tr>
<td>VSI</td>
<td>7.308b</td>
<td>7.533b</td>
<td>18.673b</td>
</tr>
<tr>
<td>± 1.462</td>
<td>± 0.226</td>
<td>± 0.822</td>
<td></td>
</tr>
</tbody>
</table>

Values in the same row with different letters are significantly different at P = 0.05.

TL: Total length, STL: Standard length, BW: Body width, TW: Total weight, GW: Gutted weight, Condition factor (K) = ([Wt (g)] / [L (cm)]^3 x 100) Where Wt: the total body weight (gm) and L: the total body length (cm), Hepatosomatic index (HSI) = 100 × (liver weight (g) × body weight(gy)^-1), Viscerosomatic index (VSI) = 100 ×(carcass weight (g) × body weight(gy)^-1).

TABLE 2. Growth performances of wild collected and hatcheries produced European seabass (D. labrax) fry.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Wild</th>
<th>GAFRD K21</th>
<th>NIOF</th>
</tr>
</thead>
<tbody>
<tr>
<td>TL (mm)</td>
<td>73.200a</td>
<td>37.600b</td>
<td>32.600c</td>
</tr>
<tr>
<td>± 6.573</td>
<td>± 1.517</td>
<td>± 2.074</td>
<td></td>
</tr>
<tr>
<td>69.400b</td>
<td>33.800b</td>
<td>29.200b</td>
<td></td>
</tr>
<tr>
<td>TLG (mm)</td>
<td>± 6.573</td>
<td>± 1.517</td>
<td>± 2.074</td>
</tr>
<tr>
<td>0.780b</td>
<td>0.380b</td>
<td>0.328b</td>
<td></td>
</tr>
<tr>
<td>TLADG (mm)</td>
<td>± 0.074</td>
<td>± 0.017</td>
<td>± 0.023</td>
</tr>
<tr>
<td>2.067b</td>
<td>1.717c</td>
<td>1.646c</td>
<td></td>
</tr>
<tr>
<td>TLSGR (%/d)</td>
<td>± 0.046</td>
<td>± 0.022</td>
<td>± 0.034</td>
</tr>
<tr>
<td>6940.00b</td>
<td>3380.00b</td>
<td>2920.00b</td>
<td></td>
</tr>
<tr>
<td>TLG% (%)</td>
<td>± 657.27</td>
<td>± 151.66</td>
<td>± 207.36</td>
</tr>
<tr>
<td>TW (gm)</td>
<td>4.048a</td>
<td>0.483b</td>
<td>0.399b</td>
</tr>
<tr>
<td>± 1.090</td>
<td>± 0.030</td>
<td>± 0.067</td>
<td></td>
</tr>
<tr>
<td>4.045b</td>
<td>0.480b</td>
<td>0.396b</td>
<td></td>
</tr>
<tr>
<td>WG (gm)</td>
<td>± 1.090</td>
<td>± 0.030</td>
<td>± 0.067</td>
</tr>
<tr>
<td>45.445b</td>
<td>5.389c</td>
<td>4.454c</td>
<td></td>
</tr>
<tr>
<td>WADG (mg)</td>
<td>± 12.244</td>
<td>± 0.335</td>
<td>± 0.757</td>
</tr>
<tr>
<td>0.668b</td>
<td>-0.359b</td>
<td>-0.458b</td>
<td></td>
</tr>
<tr>
<td>WSGR (%/d)</td>
<td>± 0.130</td>
<td>± 0.032</td>
<td>± 0.091</td>
</tr>
<tr>
<td>404.460b</td>
<td>47.960b</td>
<td>39.640b</td>
<td></td>
</tr>
<tr>
<td>WG% (%)</td>
<td>± 108.975</td>
<td>± 2.984</td>
<td>± 6.734</td>
</tr>
</tbody>
</table>

Values in the same row with different letters are significantly different at P = 0.05.
TL: Total length, TW: Total weight, Length gain (LG) = Lf – Li, where Li and Lf are initial and final lengths (mm), Length average daily gain (LADG) = Lf – Li/t, where Li and Lf are initial and final lengths (mm) and t is the time of experiment (days), Length specific growth rate % (LSGR) = (Lin Lf – Lin Li) 100/t, where Li and Lf are initial and final lengths (mm) and t is the time of experiment (days), Length gain % (LG %) = LG/ Li x 100, where Li is initial length and LG is Length gain (mm), Total weight gain (WG) = Wf – Wi, where Wi and Wf are initial and final weights (g), Weight average daily gain (WADG) = Wf - Wi/t, where Wi and Wf are initial and final weights (g) and t is the time of experiment (days), Weight Specific growth rate (WSGR) = 100 (lnWf – lnWi)/t, where Wi and Wf are initial and final weights (g) and t is the time of experiment (days), Weight gain % (WG %) = WG/ Wi x 100, where Wi is initial weight and WG is weight gain (g).

The physiology effects of European seabass (D. labrax) fry sources (wild and two hatcheries) on condition factor of total and standard length growth, hepatosomatic and vesrosomatic indexes. The NIOF marine hatchery produced fry achieved the best significant (P < 0.05) results in all this physiology parameters. Hebalah [27] found that Length-weight relationship gives an indication of the degree of the well-being of farm fish than that inhabit in El-Maadyia region. The condition factor is used in order to compare the condition, fatness or wellbeing of fish. It is based on the hypothesis that heavier fish of a particular length are in a better physiological condition [28]. Condition factor is also a useful index for monitoring of feeding intensity, age, and growth rates in fish [29]. It is strongly influenced by both biotic and abiotic environmental conditions and can be used as an index to assess the status of the aquatic ecosystem in which fish live [30]. Hebalah [27] showed that the results revealed that the mean value of condition factor for seabass from fish farm was higher than that captured from El-Maadyia region. Also, it may be attributed to fish farm do not suffer the deficiency of food. Also, it was obvious significant differences in the mean values of condition factor for different seasons of Seabass collected from El-Maadyia region, while fish farm showed no seasonal significant differences in the mean values of condition factor. Hebalah [27] reported that fishery research is necessary to understand the many
complex factors that contribute to the health and decline of our resources. This research is needed to provide management with guidance in making decisions to ensure sustainable fisheries. These research protocols must describe past and ongoing monitoring of the fishery. Little biological information on European seabass in Egyptian Mediterranean water has been gathered in the past years. Hebalah [27] reported that hepato-somatic index (HSI) has been used in fishery biology as a useful tool for assessing the fish condition.

Facey [31] stated that H.S.I. as a biomarker is often correlated with exposure to various contaminants (e.g., polychlorinated biphenyls, polycyclic aromatic hydrocarbons, and some heavy metals). Exposure to contaminants can lead to an increase in liver size as a result of increase in hepatocytes size (hypertrophy) or number (hyperplasia) [32]. Heath [33] and Facey [34] studies evaluating the relative liver size of fishes from contaminated and reference sites often utilize the HSI. Goede [35] and Ahmed [36] have used HSI as a biomarker of contaminant exposure. Hebalah [27] showed that the highest values of Hepato-somatic index for seabass from fish farm than that captured from El-Maadyia region can be attributed to feed on artificial food in aquaculture comparing with natural food organism for fish from El-Maadyia region. Rajaguru [37] reported that the lowest values of hepatosomatic indices were recorded only during the peak spawning period. This is in agreement with Hebalah [27] finding in El-Maadyia region, where the lowest values of hepatosomatic indices were found in spawning season (spring to summer).

The European seabass (D. labrax) fry sources (wild and two hatcheries) water quality parameters, Temperature in °C, pH: Acidity and alkalinity parameter, Dissolved oxygen%, conductivity in Ms/cm, TDS: Total dissolved solids in part per million and salinity in ppt: part per thousand (Table 3.). The wild collected fry achieved the best significant (P < 0.05) results in all this water quality parameters. Hebalah [27] showed that the water quality of European seabass (D. labrax) collecting seasons and locations recorded that El-Maadyia region in the summer season had the best significant (P < 0.05) DO% results compared with other location and other seasons. The fish farm in the summer season had the highest significant (P < 0.05) pH results compared with other location and other seasons. El-Maadyia region in the spring and winter seasons had the highest significant (P < 0.05) TDS results compared with other location and other seasons. El-Maadyia region in the spring, summer and autumn seasons had the highest significant (P < 0.05) salinity results compared with other location and other seasons. El-Maadyia region in the winter season had the best significant (P < 0.05) EC results compared with other location and other seasons. El-Maadyia region in the summer season and the fish farm in the winter season had the best significant (P < 0.05) turbidity results compared with other location and other seasons. Hebalah [13] recorded that European seabass (D. labrax) newly hatched larvae tanks water quality showed mostly no significant (P < 0.05) differences between treatments results within the optimum water quality ranges for European seabass larval rearing. Also Hebalah [13] recorded that European seabass (D. labrax) post larvae tanks water quality showed mostly no significant (P < 0.05) differences between treatments results within the optimum water quality ranges for European seabass post larval rearing.

These results in agreement with Hebalah [3] and Salem [5, 12] results that indicated European Seabass eggs, yolk sac larvae and first fed larvae to weaning tanks water quality results performances were within suitable limits for Seabass, also in agreement with Zaki [38], Nour [17] and EFSA [39] whom reported that the optimum European seabass eggs incubator water temperature is in the range of 12-15 °C, salinity 32-40 ppt, oxygen concentration 7.0-8.0 mg l⁻¹, oxygen saturation 80-100% and pH 7.9-8.1. Yeong [40] indicated that water quality monitoring showed no statistically significant differences between different diets, and the observed values indicate that none of the experimental diets affected the quality of water and culture water parameters were within the suitable range. Sonu [41] reported that the effects of beneficial bacteria in an aquaculture system can be explained by various mechanisms such as improvement of water quality, antagonism towards pathogens including competition for adhesion sites, enzymatic contribution to digestion in the host, and stimulation of the host immune response [42]. Grati [43] reported that although sea bass is known as a euryhaline species, and alterations in metabolic rate with varying salinity seem to be small [44], post larvae usually aggregate in brackish waters of shallow tidal lagoons and estuaries [44, 45] and then migrate offshore as they grow [46]. Environmental variables, such as water temperature, salinity, and oxygen

TABLE 3. Water and bacterial quality of wild collected and hatcheries produced European seabass (D. labrax) fry.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Wild</th>
<th>GAFRD K21</th>
<th>NIOF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temp</td>
<td>24.80 ± 0.14</td>
<td>24.35 ± 0.07</td>
<td>24.65 ± 0.21</td>
</tr>
<tr>
<td>pH</td>
<td>8.03 ± 0.08</td>
<td>8.20 ± 0.01</td>
<td>8.24 ± 0.04</td>
</tr>
<tr>
<td>DO%</td>
<td>109.60 ± 0.99</td>
<td>100.35 ± 0.78</td>
<td>94.75 ± 1.20</td>
</tr>
<tr>
<td>Ec</td>
<td>54.86 ± 13.44</td>
<td>94.85 ± 119.57</td>
<td>188.10 ± 0.28</td>
</tr>
<tr>
<td>TDS</td>
<td>31.59 ± 3.54</td>
<td>120.50 ± 0.28</td>
<td>120.75 ± 0.78</td>
</tr>
<tr>
<td>Salinity</td>
<td>6.50 ± 0.71</td>
<td>34.85 ± 0.00</td>
<td>36.80 ± 1.40</td>
</tr>
<tr>
<td>VBCF (CFU)</td>
<td>31.33 ± 3.21</td>
<td>20.33 ± 1.53</td>
<td>3.00 ± 1.00</td>
</tr>
<tr>
<td>BBCF (CFU)</td>
<td>1.00 ± 1.00</td>
<td>2.33 ± 0.58</td>
<td>3.00 ± 1.00</td>
</tr>
<tr>
<td>BBCW (CFU)</td>
<td>0.00 ± 0.00</td>
<td>4.00 ± 1.00</td>
<td>3.67 ± 1.53</td>
</tr>
</tbody>
</table>

Values in the same row within the subtales with different letters are significantly different at $P = 0.05$.

European seabass (D. labrax) fry sources (wild and two hatcheries) effects on water and fry Vibrio sp. and Bacillus sp. bacterial counts (Table 3.). Vibrio sp. hadn’t detected in the water source sample of the three fry sources. The wild collected fry achieved the best significant (P < 0.05) water and fry results in Bacillus bacterial counts. Salem [12, 3, 5] recorded that rotifers and artemia enrichment and yolk sac larvae and first fed larvae to weaning tanks bacterial counts of the potentially pathogenic bacteria such as Aeromonas sp., Staphilococcus sp. and Vibrio sp., or as potentially useful bacteria such as Bacillus sp and total bacterial count in the treatments showed that the potentially pathogenic bacteria reduced by GMW+MP followed by GMW+S than GMW and FMW. In 2013, Egypt mass mortalities in Kafr El-Sheik and El- Behira governorates tilapia fish farms which diagnosed as A. hydrophilla outbreak and also, in Maryut Valley, Alexandria governorate in European seabass, gilthead seabream and meagre farms but with different reasons of high temperatures and low oxygen stress in seabass and meagre, 100 tons equals 7 million L. E. loses [5] and because of Vibrio sp. outbreak in gilthead seabream after seabass and seabream wild collected fry were transported to this farms [5].

Conclusion

Egypt marine fish farmers prefer wild collected European seabass (D. labrax) fry than hatchery produced fry. Marine hatcheries need more efforts and technology transfer to help marine aquaculture and fisheries development, that need ranching and restocking with hatchery high quality produced fry to convince farmers that hatchery produced fry is better not only because wild fry would transfer pathogens causing outbreaks but also due to hatchery fry better growth and quality.

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References


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تأثير مصدر زريعة أسماك القاروص على أداء النمو والفسيولوجي

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تهدف هذه الدراسة إلى تقييم تأثير المصادر المختلفة لزريعة أسماك القاروص الأوروبي على أداء النمو والأداء الفسيولوجي. أجريت هذه الدراسة في معمل فسيولوجيا الأسماك شعبة المصايد بالمعهد القومي لعلوم البحار والمصايد بالإسكندرية. وأجريت هذه الدراسة باستخدام أفراد حسبن زريعة من 3 مصادر مختلفة وتم نقلها إلى المعمل. المجموعة الأولى: تم جمعها من مصدرها الطبيعي من منطقة بوغاز المعدية محافظة البحيرة. المجموعة الثانية: اليرقات كانت بعد الفقس وقد تم الحصول عليها من المفرخ البحري التابع للمعهد القومي لعلوم البحار والمصايد الأفريقي. والمجموعة الثالثة: اليرقات كانت بعد الفقس وتم الحصول عليها من المفرخ البحري. وقد تم الحصول عليها من المفرخ البحري للهيئة العامة لتنمية الثروة السمكية. المجموعة شاملاً زريعة أسماك القاروص عشوائياً ثم تم تقييم بيئة النمو والوزن والأداء الفسيولوجي لزريعة الأسماك. وقد أوضحت النتائج الدراسة المقارنة بين زريعة القاروص الأوروبي المجمعة من الطبيعة والمفرخات البحرية تأثيرها على أداء النمو الطولي والوزن، متمثل في الطول الكلي والطول القياسي وعمق الجسم والوزن الكلي والوزن منزوع الأحشاء. وقد حققت الزريعة المجمعة من المصادر الطبيعية فروق معنوية عالية في جميع معاملات النمو في حين حققت الزريعة المجمعة من المصدر البحري منفعة متميزة في جميع المعاملات الفسيولوجية. والخلاصة أن الدراسة أوضحت سبب تفضيل المربين لزريعة أسماك القاروص المجمعة من المصادر الطبيعية أكثر من الزريعة التي تنتجها المفرخات. ولذلك تحتاج المفرخات البحرية المصرية بكثير من التطور والجهد ونقل التكنولوجيا لتساعد في تنمية الأستزراع البحري وكذلك تنمية المصايد والتي تحتاج تعزيز مخزونها السمكي بأمدادات زريعة التي تنتجها المفرخات.

الكلمات الدالة: القاروص الأوروبي، الزريعة، الطبيعة، المفرخ، النمو، معامل حالة الكبد، معامل حالة الأحشاء.