

**Effect of Different Doses of Ovulin Hormone  
Suspended in Saline on the Induced Breeding  
Performance of African Catfishes *Clarias  
anguillaris* and *Clarias gariepinus* in Sokoto,  
Nigeria**

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**ABSTRACT**

A study on the effect of different doses of Ovulin hormone suspended in saline on the breeding performance of *Clarias anguillaris* and *Clarias gariepinus* was carried out. The experiment was conducted in a 2x5 factorial experiment in a Completely Randomized Design at the Hatchery Unit of the Department of Fisheries and Aquaculture, Usmanu Danfodiyo University, Sokoto. Species and hormone dilutions constituted the factors with specie having 2 levels (*C. anguillaris* and *C. gariepinus*) and Ovulin suspended in saline at 5 levels (0%, 25%, 50%, 75% and 100%). The result showed that species levels did not significantly ( $P>0.05$ ) affect the breeding performance in all the breeding performance parameters observed. However, fertilization rate, hatching rate and survival rate were significantly affected ( $P<0.05$ ) by different levels of Ovulin suspended in saline, but did not have significant influence ( $P>0.05$ ) on egg weight, spawning fecundity and relative fecundity. It could be concluded from this study that Ovulin suspended in saline can have significant influence on the breeding performance of African catfish.

*Keywords: African catfish, Ovulin, Hormone, Clarias anguillaris, Clarias gariepinus*

**1. INTRODUCTION**

The African catfish is widely considered as the leading cultured fish in Nigeria. Some of the credentials of African catfish are: high growth rate reaching market size of 1 kg in 5–6 months under intensive management conditions: highly adaptable and resistant to handling and stress; can be artificially propagated by induced spawning techniques for reliable mass supply of fingerlings; commands a very high commercial value where it is highly cherished as food in Nigerian homes and hotels [1] [2].

Fertilization, hatching and early survival of larvae are vital for successful aquaculture of the African catfishes [3]. Richter and Van der Hurk [4] reported that the problem of inadequate supply of fish seed can only be solved through induced breeding by the application of various inducement materials. Various types of fishes have been induced to spawn, using various hormonal materials [5] [6] [7] [8]. Some of these spawning agents are either difficult to quantity, ineffective or of short shelf life, and for that, many breeders are reluctant to use them in field conditions. However, the commercially available synthetic inducing hormones in ready-made form containing GnRH $\alpha$  and dopamine (Ovaprim, Ovopel, Ovulin, Ovatide,

35 Dagin and Aquaspawn) are becoming very popular and found to be efficient in successful  
36 spawning of fishes [9] [10] [11].

37

38 Prior to the work reported by Olumuji and Mustapha [12], the synthetic hormone for fish  
39 breeding has been used undiluted unlike natural hormone (pituitary extract). Normal saline  
40 which is commonly used form of saline solution is prepared by dissolution of 9g of NaCl in 1  
41 litre of water [13]. Therefore, not much, if any, has been reported on the use of Ovulin diluted  
42 with normal saline on induced breeding of *C. anguillaris* and *C. gariepinus*.

43

44 This study therefore, was carried out using Ovulin hormone diluted with normal saline for  
45 induced breeding of *C. anguillaris* and *C. gariepinus* in order to test the effectiveness and  
46 efficiency of the hormone in induced breeding of African catfish and to compare the effect of  
47 various doses of the normal saline diluted-hormone with undiluted ones on the fertilization  
48 rate of the eggs, hatching rate of the eggs and larval survival of both *Clarias anguillaris* and  
49 *Clarias gariepinus*.

50

## 51 **2. MATERIAL AND METHODS**

### 52 **2.1 Description of the Study Area**

53 The experiment was conducted at the Hatchery unit of the Department of Fisheries and  
54 Aquaculture located at the main campus of Usmanu Danfodiyo University, Sokoto (N13° 07'  
55 45.12" E5° 12' 18").

### 56 **2.2 Broodstock Collection and Management**

57 Eighty (80) broodstock, forty (40) each of *C. anguillaris* and *C. gariepinus* (30 females, 10  
58 males each) were collected from the Departmental Fish Farm of the department of Fisheries  
59 and Aquaculture, Usmanu Danfodiyo University Sokoto. The fish were conditioned at the  
60 hatchery complex of the farm and were fed commercially produced industrial feed (Coppens)  
61 at 3% body weight twice daily for two weeks.

### 62 **2.3 Experimental Design and Procedures**

63 The experiment was set up in a 2 factor (2 x 5) factorial experiment in a Completely  
64 Randomized Design (CRD) with two levels of *Clarias* species (*C. anguillaris* and *C.*  
65 *gariepinus*) and five inclusion levels of Ovulin hormone suspended in saline (at 0%, 25%,  
66 50%, 75% and 100%). All the treatments were replicated three times to give a total of 30  
67 spawning trials (i.e. 2 x 5 = 10, replicated 3 times = 30). Induced breeding was carried out  
68 and data collected was subjected to statistical analysis.

69

70 The treatment combinations were therefore; A<sub>0</sub>B<sub>0</sub>, A<sub>0</sub>B<sub>1</sub>, A<sub>0</sub>B<sub>2</sub>, A<sub>0</sub>B<sub>3</sub>, A<sub>0</sub>B<sub>4</sub>, A<sub>1</sub>B<sub>0</sub>, A<sub>1</sub>B<sub>1</sub>, A<sub>1</sub>B<sub>2</sub>,  
71 A<sub>1</sub>B<sub>3</sub>, A<sub>1</sub>B<sub>4</sub>. Where A<sub>0</sub> = *Clarias anguillaris*, A<sub>1</sub> = *Clarias gariepinus*, B<sub>0</sub> = 0%, B<sub>1</sub> = 25%, B<sub>2</sub> =  
72 50%, B<sub>3</sub> = 75% and B<sub>4</sub> = 100%. These were randomly distributed in triplicate tanks in a  
73 CRD.

74

## 75 **2.4 Induced Spawning and Hormone Treatment**

76 Artificial hormone (Ovulin) was used for inducing ovulation at a recommended dosage of  
77 0.5ml/kg body weight of female broodstock, while half dosage was administered to male  
78 broodstock [14]. Hormone administration was carried out via intramuscular injection with 0%,  
79 25%, 50%, 75% and 100% inclusion levels of normal saline and the injected fish were kept  
80 separated in well-labelled closed containers containing water. The containers with the  
81 injected fish were covered and heavy stones were put on the lid of the containers that  
82 prevented the fish from jumping out.

## 83 **2.5 Procurement of ripe eggs and milt**

84 After a latency period of about 8 hours, at a temperature of about 28<sup>0</sup>C, the eggs were  
85 collected from each female through stripping by gently pressing the abdomen of the fish. The  
86 eggs were collected into clean bowls labelled accordingly. The weights of the eggs were  
87 recorded. Milt was obtained by sacrificing the males. Each male was dissected carefully and  
88 their milt sac obtained. A small incision was made on the lobes of the testes with a sharp  
89 razor blade and the milt was squeezed into a dry Petri dish containing the collected eggs.

## 90 **2.6 Artificial fertilization**

91 Dry method of fertilization was used where the milt obtained from the male fishes was  
92 squeezed gently onto the stripped eggs obtained from the females accordingly and stirred  
93 gently and thoroughly using plastic spoon for about 2 minutes to allow contact and adequate  
94 fertilization. Normal saline was added before spreading the eggs on the spawning nets in the  
95 incubation units prepared earlier for that purpose [15].

## 96 **2.7 Incubation and hatching**

97 The mixture of the eggs and milt were distributed in a single layer on the spawning nets in  
98 the well aerated incubation bowls. Three gram of egg was collected from each sample and  
99 incubated in 60-litre plastic containers for the experiment, for easy assessment of fertilization  
100 and hatching rates [16] [17] [18].

## 101 **2.8 Data collection**

102 Data on induced breeding performance (ovulation response, fecundity, fertilization rate,  
103 hatching rate and larval survival rate) were recorded.

## 104 **2.9 Data analysis**

105 The data collected on the induced breeding parameters was subjected to statistical analysis  
106 using SPSS (Version 20). All data with discrete counts and percentages was transformed  
107 before analysis was carried out. The data were analyzed using analysis of variance  
108 (ANOVA) to test for significant differences ( $P < 0.05$ ) in fertilization rate, hatching rate and  
109 larval survival, and means were separated using Duncan's Multiple Range Test (DMRT)  
110 where significant difference exist [19].  
111

### 112 3. RESULTS AND DISCUSSION

#### 113 3.1 Water quality parameters

114 The water quality parameters recorded during the experiment are shown in table 2. The  
115 mean temperature recorded during latency period ranged from 27.47 to 28.17 for all the  
116 treatments and the mean temperature recorded during incubation ranged from 25.90 to  
117 28.00. pH of the water during the experiment was in the range of 7.10 to 7.73 for all the  
118 treatments while the mean dissolved oxygen value recorded was between 6.40 and 7.27.

119

120 According to [20], the physico-chemical parameters of water are important to the growth,  
121 productivity and survival of aquatic organisms especially fish as they play a vital role in the  
122 biology and physiology of the fish. This statement agrees with the findings of [14] who  
123 opined that the best temperature range for optimum production of *Clarias* species is 25 –  
124 31°C and the water quality parameters recorded during this experiment are within those  
125 recommended levels for catfish breeding. The mean temperature values recorded during the  
126 experiment was similar to what was observed by [21] for *Clarias gariepinus* that exhibited a  
127 latency period of about 8 hours at 28°C. The mean pH value recorded during the experiment  
128 also falls within the normal range of 6.5 to 8.0 for catfish according to [21]. And this is in  
129 agreement with many researchers that the best water for fish cultivation is that which has a  
130 pH range of between 7 to 8. Dissolved oxygen level during the experiment was also within  
131 the recommended level for catfish.

132

133 **Table 2: Mean water quality parameters recorded during the experiment with ovulin**

Treatments	Parameters			
	Latency temp (°C)	Incubation temp (°C)	pH (mg/l)	DO (mg/l)
A <sub>0</sub> B <sub>0</sub>	27.97	26.73	7.17	6.47
A <sub>0</sub> B <sub>1</sub>	27.73	25.90	7.23	6.50
A <sub>0</sub> B <sub>2</sub>	27.60	27.57	7.13	6.87
A <sub>0</sub> B <sub>3</sub>	27.70	27.40	7.53	6.63
A <sub>0</sub> B <sub>4</sub>	27.53	-	-	-
A <sub>1</sub> B <sub>0</sub>	27.93	27.00	7.27	6.93
A <sub>1</sub> B <sub>1</sub>	28.17	27.23	7.73	6.40
A <sub>1</sub> B <sub>2</sub>	28.17	28.00	7.10	6.77
A <sub>1</sub> B <sub>3</sub>	27.47	27.53	7.17	7.27
A <sub>1</sub> B <sub>4</sub>	27.63	-	-	-

134

#### 135 Mean weight, dosage, incubation period and latency period

136 The mean initial weight of the broodstocks used for the experiment ranged from 326.67g to  
137 480.00g (Table 3). Dosage administered for injection of the broodstocks ranged from 0.16 ml  
138 to 0.24 ml at the recommended dosage of 0.5 ml/kg body weight and the quantity of egg  
139 used for incubation was 3 grams for each treatment. The number of eggs obtained in 1 g of  
140 egg was between the range of 623 to 645. Latency period was between 7hrs and 58mins to  
141 8hrs and 12mins and incubation period ranged between 22hrs to 23hrs and 4mins for all the  
142 treatments.

143

144 The size range of the broodstocks used in the experiment was in agreement with [21] who  
145 opined that African catfish *clarias* can become mature and breed as from 200g body weight.  
146 And it agrees also with [22] who reported that the ideal broodfish weight should be between

147 300-800 grams, as larger fish are difficult to handle and often results in substantial egg  
 148 losses prior to stripping. The time taken to achieve ovulation (latency period) is dependent  
 149 upon water temperature as reported by [22], as such the higher the temperature the quicker  
 150 the eggs ovulate. In other words, the higher the temperature the shorter the latency period.  
 151 The mean latency period observed in this study fall within 8hrs at mean temperature of  
 152 between 27 – 28<sup>0</sup>C and was similar to what was reported by [22] for *Clarias gariepinus*. The  
 153 result also showed no significant variation of latency time between the treatments except for  
 154 treatment induced with 100% normal saline which could be the reason why ovulation did not  
 155 occur for that particular treatment in all the phases which is due to the lack of hormone effect  
 156 that foster ovulation in fish. This was similar to what was observed by [12] on induced  
 157 breeding of *Clarias gariepinus* using different doses of normal saline-diluted ovaprim.  
 158

159 de Graaf and Janssen [22] reported that the development process of fish from fertilized egg  
 160 to hatching is like all other biological processes, that is, it is dependent upon water  
 161 temperature, as such the higher the water temperature the faster the eggs hatch. The  
 162 incubation period observed in this experiment was in the range of 21 to 23hrs at a  
 163 temperature range of about 25.9 to 28.6<sup>0</sup>C which was similar to observations of Viveen *et al.*  
 164 (1985) and was also in comparison with the findings of [23] for *Clarias gariepinus* that  
 165 achieved incubation period of 15hrs at a temperature of 30<sup>0</sup>C  
 166

167 **Table 3: Weight, dosage, latency period and incubation period during induced**  
 168 **breeding with ovulin**

Treatments	Parameters				
	Initial weight (g)	Dosage (ml)	No. of Egg (1g)	Latency period	Incubation period
A <sub>0</sub> B <sub>0</sub>	453.33	0.23	623	8h 9m	23h 4m
A <sub>0</sub> B <sub>1</sub>	433.33	0.22	634	8h 2m	22h 57m
A <sub>0</sub> B <sub>2</sub>	426.67	0.21	645	8h	22h 10m
A <sub>0</sub> B <sub>3</sub>	480.00	0.24	629	8h 1m	22h
A <sub>0</sub> B <sub>4</sub>	346.67	0.17	-	-	-
A <sub>1</sub> B <sub>0</sub>	363.33	0.18	641	8h 12m	22h 32m
A <sub>1</sub> B <sub>1</sub>	380.00	0.19	643	7h 58m	22h 8m
A <sub>1</sub> B <sub>2</sub>	326.67	0.16	630	8h	22h 20m
A <sub>1</sub> B <sub>3</sub>	433.33	0.22	641	8h 3m	22h 25m
A <sub>1</sub> B <sub>4</sub>	413.33	0.21	-	-	-

169

170 **Breeding performance of *C. anguillaris* and *C. gariepinus* induced with ovulin**  
 171 **suspended in saline**

172 The result of breeding performance of *C. anguillaris* and *C. gariepinus* induced with different  
 173 levels of ovulin hormone suspended in saline is shown in Table 4. The result indicates that  
 174 species levels did not significantly ( $p>0.05$ ) affect the breeding performance in this  
 175 experiment with *C. gariepinus* producing relatively higher mean values compared to *C.*  
 176 *anguillaris* in egg weight, spawning fecundity and relative fecundity, as well as the breeding  
 177 performance parameters (fertilization rate, hatching rate and larval survival rate). However,  
 178 there was no significant difference ( $p>0.05$ ) between the means. The result further showed  
 179 that different doses of ovulin suspended in saline significantly affected the breeding  
 180 performance in this experiment. In terms of egg weight, spawning fecundity and relative  
 181 fecundity, positive control (0% saline), 25%, 50% and 75% dilution levels produced  
 182 significantly similar ( $p>0.05$ ) mean values while the negative control (100% saline) did not

183 produce any value since spawning did not occur. 0% normal saline dilution (positive control)  
184 produced relatively higher mean values that are significantly different ( $p < 0.05$ ) from the other  
185 dilution levels (25%, 50% 75% and 100%) in terms of breeding performance parameters  
186 (fertilization rate and hatching rate) except for survival rate where no significant difference  
187 ( $p > 0.05$ ) exist between the mean values. 25% and 50% dilution levels produced statistically  
188 similar ( $p > 0.05$ ) result in terms of fertilization rate while 50% and 75% dilution levels  
189 produced similar result statistically ( $p > 0.05$ ) in terms of hatching rate. There was no  
190 significant interaction between the factors in this experiment.

191  
192 The spawning fecundity observed in the study showed that different doses of ovulin  
193 suspended in saline at 25%, 50% and 75% inclusion levels can be effective in the induced  
194 breeding of *C. anguillaris* and *C. gariepinus*. The highest mean fecundity value 33,939 was  
195 observed in 75% dilution level. The value obtained was in agreement with [21], that larger  
196 female fishes contain more eggs than smaller fishes and therefore have higher fecundity  
197 values and this could also be due to the efficacy of the hormone used which indicates that  
198 even a small quantity of hormone can be diluted with saline and be effective in the induced  
199 breeding of African catfish.

200  
201 The highest mean fertilization rate was observed in positive control treatment (0% normal  
202 saline dilution) with mean values of 92.22 and this was significantly different ( $p < 0.05$ ) from  
203 the other dilution levels. This was similar to what was obtained by [12] who examined the  
204 effect of varying doses of normal saline-diluted ovaprim on the induced breeding of *C.*  
205 *gariepinus*. This work however, showed that suspending generic hormone in saline at 25%,  
206 50% and 75% dilution levels can be effective in the induced breeding of African catfish,  
207 which agrees with the findings of [23] that even small quantity of hormone below the  
208 manufacturer's recommended dose can successfully induce ovulation in African catfish.

209  
210 The highest mean hatching rate of 86.62 was observed in positive control treatment (0%  
211 dilution) and this was significantly different from the other dilution levels. This was relatively  
212 higher than what was obtained by [23] on the induced breeding of *C. gariepinus* using  
213 different doses of normal saline-diluted ovaprim and this could be attributed to the efficacy of  
214 the hormones used in this study. The mean hatching rate obtained was also higher than  
215 what was obtained by Moses *et al* for Kainji strains of *C. anguillaris* (58.58%) and *C.*  
216 *gariepinus* (52.44%) using ovaprim and [24] using ovatide and ovaprim on *C. gariepinus* with  
217 mean values of 59.70% and 66.37% respectively.

218  
219 The highest larval survival rate recorded in this experiment was 93.73 in 75% dilution level.  
220 The value obtained were comparatively higher than what was obtained by several authors  
221 working on *Clarias*; [25] who worked on induced breeding of *C. gariepinus* under varying  
222 broodstock ratios; [26] on the effect on breeding performance and egg quality of *C.*  
223 *batrachus* at various doses of ovatide during spawning induction. Likewise, the result  
224 obtained was higher than what was obtained by [12] on the induced breeding of *C.*  
225 *gariepinus* using different doses of normal saline-diluted ovaprim, and this can be related to  
226 the spawning medium (tank) used to run the experiment which was larger in this experiment  
227 with more space and constant aeration using aerators that provide dissolved oxygen into the  
228 medium which agrees with [27] and [20] that physico-chemical parameters of water such as  
229 high concentration of dissolved oxygen affects the hatchability and larval survival of fish.

230

231 **Table 4: Main effects specie and ovulin suspended in saline effects on induced**  
 232 **breeding performance of *C. anguillaris* and *C. gariepinus***

Factors	Parameters					
	EW(g)	SF	RF(g)	FR(%)	HR(%)	SR(%)
<b>Specie</b>						
<i>C. anguillaris</i>	34.92	22,245	49.13	60.67	56.87	73.09
<i>C. gariepinus</i>	35.77	22,876	61.90	65.56	58.84	74.27
<b>SEM</b>	3.71	2463.64	6.00	1.86	1.98	1.13
<b>Hormone dilution</b>						
0%	39.93 <sup>a</sup>	25,295 <sup>a</sup>	65.74 <sup>a</sup>	92.22 <sup>a</sup>	86.63 <sup>a</sup>	92.08 <sup>a</sup>
25%	45.42 <sup>a</sup>	29,034 <sup>a</sup>	71.82 <sup>a</sup>	82.78 <sup>b</sup>	75.09 <sup>b</sup>	90.61 <sup>a</sup>
50%	37.97 <sup>a</sup>	24,536 <sup>a</sup>	62.69 <sup>a</sup>	77.22 <sup>b</sup>	65.80 <sup>c</sup>	92.00 <sup>a</sup>
75%	53.42 <sup>a</sup>	33,939 <sup>a</sup>	77.33 <sup>a</sup>	63.33 <sup>c</sup>	61.76 <sup>c</sup>	93.73 <sup>a</sup>
100%	00.00	0.00	0.00	0.00	0.00	0.00
<b>SEM</b>	5.87	3895.35	9.49	2.94	3.13	1.70
<b>Interaction</b>	NS	NS	NS	NS	NS	NS

233 Means with the same superscripts on the same column are not significantly different  
 234 ( $p>0.05$ )

235 NS = Not significant

236 EW = Egg weight, SF = Spawning Fecundity, RF = Relative Fecundity, FR = Fertilization  
 237 Rate, HR = Hatching Rate, SR = Survival Rate

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#### 4. CONCLUSION

It has been observed from the result of this experiment that there is no statistically significant difference between *Clarias anguillaris* and *Clarias gariepinus* induced using Ovulin synthetic hormone in terms of egg weight, fecundity and all the breeding performance parameters such as fertilization rate, hatching rate and larval survival rate. It has also been observed that positive control (100% hormone) stand out as the best performing treatment in relation to breeding performance with respect to *C. anguillaris* and *C. gariepinus* compared to other treatments where hormone was diluted with normal saline.

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