The Effects of Some Physico-Chemical Parameters on the Breeding of Catfish (Clarias Gariepinus) using Ovaprim and Ovatide

Akombo, Pauline Mbakaan; Atile, John Iornyiman;* Obaje, John Agwajinya

Department of Biological Sciences, Benue State University, Makurdi, Nigeria

*Corresponding Author: Atile, John Iornyiman, Department of Biological Sciences, Benue State University, Makurdi, Nigeria

Abstract: A comparative assessment of the effects of some physico-chemical parameters on breeding of catfish (Clarias gariepinus) using ovaprim and ovatide was investigated from the month of July to August, 2016 in Makurdi, Benue State, Nigeria. Twelve (12) female brooders and six (6) males were obtained from Ocepson’s Farm and Aqua-heaven Fish Farm respectively in Makurdi. The doses of 0.8ml/kg, 0.5ml/kg and 0.3ml/kg of ovaprim and 0.3ml/kg, 0.2ml/kg and 0.1ml/kg of ovatide were used. Some physico-chemical parameters such as dissolved oxygen, temperature and pH were monitored during breeding of the fish. The maximum mean temperature of 26.43°C was observed in 0.3ml/kg dosage of ovaprim while in ovatide, the maximum mean temperature of 26.90°C was observed in a dosage of 0.1ml/kg. There was no significant difference (p>0.05) in the mean temperature amongst all the varying doses of the hormones used. The maximum mean pH of 7.01 was observed in 0.8ml/kg dosage of ovaprim whereas a maximum mean pH of 6.99 was observed in a dosage of 0.3ml/kg of ovatide. There was no significant difference (p>0.05) in the mean water pH amongst the varying doses of the hormones used. The minimum and maximum mean incubation period of 20.66±40 and 23.00±25.00 hours respectively were observed in ovaprim. The minimum and maximum mean incubation period of 20.83±38.00 and 22.83 hours respectively were observed in ovatide. There was a significant difference (p<0.05) in the mean incubation period of the eggs that were induced with varying doses of ovaprim and ovatide.

Keywords: ovaprim, ovatide, physico-chemical parameters, breeding, Clarias gariepinus

1. INTRODUCTION

Physico-chemical parameters of water are known to influence fish breeding. Some of these physico-chemical parameters are known to affect fish breeding either when they are in low, moderate or high states. Among the most prominent physico-chemical parameters that can influence fish breeding include Temperature, Dissolved oxygen and pH. According to Chukwuma and Henry (2012), water hardness does not affect the fertilization of Clarias gariepinus but was known to have affected the hatching rate of the fish.

A study conducted by Oyelese (2006) reported that fertilization and hatchability was favoured at higher temperature. According to Amaechi and Solomon (2015), optimum temperature range for fertilization and hatchability was 26-27°C. The study further stated that the dissolved oxygen level of 5.03mg/l and a pH range of 7.10-8.70 favoured the fertilization and hatchability of Clarias gariepinus.

Bhatnagar and Sangwan (2009) reported that dissolved oxygen in the range of 4.5-8.0mg/l was suitable for fish breeding. The importance of water dissolved oxygen was stressed by Bhatnagar and Grag (2000). The study reported that lower level in water oxygen can lead to poor feeding efficiency, starvation, slow growth rate as well as mortality of the fish.

The study conducted by Santhosth and Singh (2007) revealed that fish breeding was suitable at a pH range of 6.7-9.5. The study further revealed that a water pH level above and below the pH range posed stress to the fish strain used.

International Journal of Innovative Studies in Aquatic Biology and Fisheries (IJISABF)
Whenever a fish farmer or breeder embarks on fish breeding, the aim is to have a good success to produce large number of hatchlings that can grow up to fingerlings and eventually up to table size fish. However, this cannot be achieved unless the breeder knows the most prominent and suitable range of physico-chemical parameters that can be maintained throughout the breeding period why using a given doses of a particular hormone to induce the fish. Hence the need to check the effects of some physico-chemical parameters on breeding of catfish (Clarias gariepinus) using ovaprim and ovatide. This will provide baseline information on the most prominent physico-chemical parameters such as temperature, dissolved oxygen and pH as well as their suitable range that can be maintained during breeding of Clarias gariepinus using ovaprim and ovatide.

2. MATERIALS AND METHODS

2.1. Study Area

The research was conducted in Toc’s Mini Fish Hatchery (A private hatchery situated behind Tilley Gyado College North bank, Makurdi, Nigeria), located on latitude 7.7493N and longitude 8.5508E (Fig. 1).

2.2. Brood Stock Procurement and Acclimatization

Thirty (30) healthy brood stocks of the African Catfish Clarias gariepinus (20 females and 10 males) were purchased from two different fish farms in Makurdi to avoid inbreeding. The females were gotten from Ocepson’s Farm along University of Agriculture road, Makurdi, Benue State, Nigeria, while the males were gotten from Aqua-heaven Farm, North Bank, behind cattle market, Makurdi. All brood stocks were selected by examining their external morphological characteristics. Both males and females were acclimatized in separate earthen ponds of 6 x 6 x 2 meters for 3 weeks during which
they were fed with a formulated diet of 40% crude protein twice daily at 5% of total fish biomass. The fish samples were each weighed with a sensitive electronic weighing balance (Camry Emperors) of 1500g maximum capacity.

2.3. Experimental Design

Twelve (12) females and six (6) males were selected randomly at the ratio of 2:1 with three (3) treatments and three (3) replicates each for ovaprim and ovatide. Ovaprim consisted of Treatment A: 0.8 ml/kg, Treatment B: 0.5 ml/kg, and Treatment C: 0.3 ml/kg. Ovatide consisted of Treatment O: 0.3 ml/kg, Treatment P: 0.2 ml/kg, and Treatment Q: 0.1 ml/kg.

2.4. Selection of Brooders

Twelve (12) female and six (6) male brooders with weight between 1.0kg-1.5kg were selected. A female was considered ripped if the sex organ was reddish and the abdomen was well protruded and eggs ooze out freely when the abdomen was gently pressed antero-posteriorly while the male was considered ripped if the tip of the genital papilla was reddish in colour (Olubiyi et al., 2005).

2.5. Hormone Injection

Selected female brooders were injected using a 1 and 2ml graduated syringe intramuscularly at an angle of 30-45° at the dorsal fin with different doses of ovaprim and ovatide (Viveen et al., 1985). The males were not administered with hormones.

Broods were collected from earthen ponds just prior to injection. A single injection was given to the female from three doses of Ovaprim and Ovatide hormone. The broods were treated with inducing agent at evening so that fish ovulate in the morning. Injection at the interval of 30 minutes to prevent the fish from attaining latency period at once.

2.6. Stripping and Fertilization

The females were stripped by gently pressing the abdomen between 8-11 hours after injecting the fish. This was carried out by holding the fish at head and covering with a wet towel and at the tail. The ovulated eggs were stripped into a dry plastic bowl and 8.5g of eggs were collected from each sample into each labelled bowl for easy identification.

The males of the Catfish were sacrificed to obtain their milt (sperm) by dissecting them with a new razor blade into a dry plastic bowl and washed with 10ml of normal saline to enable gentle movement of sperms, reduces the stickiness of eggs and to prolong the fertilizing capacity of the milt. The eggs obtained from the stripping were weighed, followed by mixing the sperm with eggs and washing the sperm sac in to the eggs carefully for about 30-60 seconds by shaking the plastic plate and adding equivalent volume of clean water to rinse the fertilized eggs.

Thus fecundity was calculated using this formula as described by Brzuska (2003) as follows;

\[
\text{Stripping (\%)} = \frac{\text{Weight of stripped eggs}}{\text{Body weight}} \times 100
\]

2.7. Incubation

The fertilized eggs were evenly spread over the nylon mosquito net mesh size of 2 mm and placed in a 10 litres plastic trough containing about 7-8 litres of water.

2.8. Estimate of Percentage Fertilization, Hatchability and Survival Rates of African Catfish *Clarias gariepinus*

After 20-24 hours of fertilization, dead and unviable eggs which had turned whitish were collected after removals of the spawning netting by siphoning and were counted and percentage fertilization was estimated.

The fertilized eggs were counted in other to calculate the percentage fertilization as described by Adebayo and Popola, (2008) as follows;
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Fertilization (%) = \( \frac{\text{Number of fertilized eggs}}{\text{Total number of counted eggs}} \times 100 \)

Percentage hatchability was calculated after the period of 20-24 hours using a 500ml beaker, where the un-hatched eggs were estimated and this was used to calculate hatchability as described by Haniffa and Sridhar, (2002) as estimated below;

Hatchability (%) = \( \frac{\text{Number of hatchlings}}{\text{Total number of eggs counted}} \times 100 \)

2.9. Water Quality Parameters

pH, Temperature (°C) and Dissolved Oxygen (DO) of the water were monitored twice daily using pH meter, Thermometer (check temp °C) and dissolved oxygen meter (DO-5509) respectively.

2.10. Data Analysis

The data of the number of eggs stripped, fertilized, hatched and survival rate of fry were analyzed using Simple percentages and ANOVA, to compare the dosage of Ovaprim and Ovatide using SPSS (Statistical Package for Social Sciences) computer package was used to perform data analysis.

3. RESULTS

3.1. Water Quality

The physico-chemical conditions of water during the experiments are shown in the following table, Temperature, pH and dissolved oxygen of water during treatments ranged from 25.06-25.97°C, 7.00-7.01 and 5.78-5.83mgL\(^{-1}\) for ovaprim and 26.85-26.90°C, 6.94-6.99 and 5.71-5.86mgL\(^{-1}\) for ovatide.

3.2. Temperature

Temperature was measured twice daily and there was slight fluctuation during the study period.

The temperature ranges between 25.06-26.43°C in all ovaprim treatments. The maximum mean temperature was observed in 0.3ml doses of ovaprim hormone treated basins during the experimental period. (Fig.1)

In ovatide hormone, the temperature ranges between 26.85-26.90°C in all treatments while the maximum mean temperature was observed in 0.1ml doses of ovatide hormone treated basins during the experimental period (Fig.1).

Table 1. The Fish Weight, Mean Egg Weight, Mean Incubation and Latency Period during the Study.

<table>
<thead>
<tr>
<th>Hormone Doses (ml/kg)</th>
<th>Female weight (kg)</th>
<th>Male weight(kg)</th>
<th>Mean egg weight (g)</th>
<th>Mean latency period (hrs)</th>
<th>Mean incubation period (hrs)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OVAPRIM</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.8</td>
<td>1.0±0.029(^a)</td>
<td>1.2±0.101(^a)</td>
<td>185.20±0.20(^a)</td>
<td>8.16±10.00(^a)</td>
<td>20.66±40.00(^a)</td>
</tr>
<tr>
<td>0.5</td>
<td>1.0±0.031(^a)</td>
<td>1.5±0.203(^b)</td>
<td>166.25±0.25(^b)</td>
<td>11.66±20.00(^b)</td>
<td>23.00±25.00(^b)</td>
</tr>
<tr>
<td>0.3</td>
<td>1.0±0.029(^a)</td>
<td>1.2±0.122(^a)</td>
<td>178.30±0.26(^a)</td>
<td>10.00±12.00(^b)</td>
<td>22.66±27.00(^b)</td>
</tr>
<tr>
<td>P value</td>
<td>p&gt;0.05</td>
<td>P&lt;0.05</td>
<td>P&lt;0.05</td>
<td>P&lt;0.05</td>
<td>P&lt;0.05</td>
</tr>
<tr>
<td>OVATIDE</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3</td>
<td>1.2±0.019(^a)</td>
<td>1.3±0.100(^a)</td>
<td>162.70±0.24(^a)</td>
<td>8.33±12.50(^a)</td>
<td>20.83±30.00(^a)</td>
</tr>
<tr>
<td>0.2</td>
<td>1.0±0.031(^a)</td>
<td>1.5±0.201(^b)</td>
<td>175.60±0.23(^b)</td>
<td>11.50±15.00(^b)</td>
<td>22.83±32.00(^b)</td>
</tr>
<tr>
<td>0.1</td>
<td>1.1±0.029(^a)</td>
<td>1.2±0.022(^a)</td>
<td>178.20±0.26(^b)</td>
<td>10.00±11.00(^b)</td>
<td>22.50±38.00(^b)</td>
</tr>
<tr>
<td>P value</td>
<td>p&gt;0.05</td>
<td>P&lt;0.05</td>
<td>P&lt;0.05</td>
<td>P&lt;0.05</td>
<td>P&lt;0.05</td>
</tr>
</tbody>
</table>
3.3. pH

In ovaprim hormone, the pH value of each treatment basin was in the range between 7.00 - 7.01 in all treatment. There was no significant difference (p>0.05) in pH concentration among all treatments during the study periods (Fig. 2).

In ovatide hormone, the pH value of each treatment basin was in the range between 6.94-6.99 in all treatments. There was no significant difference (p>0.05) in pH among all treatments during the study periods (Fig. 2).

3.4. Dissolved Oxygen

In ovaprim hormone, the dissolved oxygen (DO) concentration was measured in the range between 5.78-5.83 mg L\(^{-1}\) in all treatments. The maximum dissolved oxygen (DO) concentration was 5.83 mgL\(^{-1}\) recorded in 0.2ml doses of ovaprim hormone treated basins during the experimental period (Fig. 3). There was no significant difference (p>0.05) in dissolved oxygen concentration among all treatments during the study period. The range of dissolved oxygen observed in this study slightly varied among all treatments.

In ovatide hormone, the dissolved oxygen (DO) concentration was measured in the range between 5.71-5.86 mg L\(^{-1}\) in all treatments. The maximum dissolved oxygen (DO) concentration was 5.86 mgL\(^{-1}\) recorded in 0.3ml doses of ovatide hormone treated basins during the experimental period (Fig. 3). There was no significant difference (p>0.05) in dissolved oxygen concentration among all treatments during the study periods.

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**Fig1. Mean Temperature in Ovaprim and Ovatide during the Study**

**Fig2. Mean pH in Ovaprim and Ovatide during the Study**
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### Table 2. Mean Water Quality Conditions and Latency Temperature of Ovaprim during the Study

<table>
<thead>
<tr>
<th>Hormone Dose (ml/kg)</th>
<th>DO (mg L⁻¹)</th>
<th>PH</th>
<th>TEMP (°C)</th>
<th>LATENCY TEMP</th>
</tr>
</thead>
<tbody>
<tr>
<td>OVA PRIM</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.8</td>
<td>5.7800±0.63*</td>
<td>7.0050±0.01*</td>
<td>25.0575±2.49*</td>
<td>27.7250±0.32*</td>
</tr>
<tr>
<td>0.5</td>
<td>5.8200±0.13*</td>
<td>7.0000±0.00*</td>
<td>25.9700±0.16*</td>
<td>27.5325±0.18bc</td>
</tr>
<tr>
<td>0.3</td>
<td>5.8325±0.13*</td>
<td>7.0000±0.00*</td>
<td>26.4275±0.10*</td>
<td>27.8900±0.24ac</td>
</tr>
</tbody>
</table>

* P value: p>0.05

### Table 3. Mean Water Quality Conditions and Latency Temperature of Ovatide during the Study

<table>
<thead>
<tr>
<th>Hormone Doses (ml/kg)</th>
<th>DO (mg L⁻¹)</th>
<th>PH</th>
<th>TEMP (°C)</th>
<th>LATENCY TEMP</th>
</tr>
</thead>
<tbody>
<tr>
<td>OVA TIDE</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3</td>
<td>5.8575±0.14*</td>
<td>6.9875±0.03*</td>
<td>26.8825±0.07*</td>
<td>27.4750±0.29*</td>
</tr>
<tr>
<td>0.2</td>
<td>5.7125±0.23*</td>
<td>6.9375±0.03*</td>
<td>26.8500±0.04*</td>
<td>26.9900±0.01*</td>
</tr>
<tr>
<td>0.1</td>
<td>5.7400±0.19*</td>
<td>6.9625±0.03*</td>
<td>26.9025±0.05*</td>
<td>28.0500±0.00*</td>
</tr>
</tbody>
</table>

* P value: p>0.05

### Fig 3. Mean Dissolved Oxygen in Ovaprim and Ovatide during the Study

**4. DISCUSSION**

During the study period, the mean temperature ranged between 25.06-26.43°C in incubated eggs that ovulated as a result of inducing the parent stock with varying doses of ovaprim. The maximum mean temperature was observed in 0.3ml/kg dose of ovaprim hormone (Fig. 1). In ovatide hormone, the mean temperature ranged between 26.85-26.90°C. The maximum mean temperature was observed in incubated eggs that were ovulated by using 0.1ml/kg of ovatide hormone (Table 2). The differences in the mean temperature in both hormones could be attributed to the quantity of heat absorbed by each egg in each hatchery tank, weather and environmental conditions. There was no significant difference (p>0.05) in the mean temperature among all the varying doses of the hormones used. The highest temperature which occurred at 0.3ml/kg in ovaprim hormone had the lowest fertilization and hatchability rates. These results disagreed with the previous study of Oyelese (2006) which stated that at higher temperatures, fertilization and hatchability gave the best results. However, this statement agrees with those trends in ovatide where at higher temperature, fertilization and hatchability rates were at their best. These results agrees with a study conducted by Amaechi and Solomon (2015) which reported that optimum temperature ranges for fertilization and hatchability was between 26-27 °C. Gomina (2011) reported that the best temperature for fertilization and hatchability was 25.9 °C and Alemayehu (2015) reported that the best temperature for fertilization and hatchability was between 25.10-26.12 °C.

The mean water pH of the incubated eggs that were stripped as a result of using varying dosages of ovaprim to stimulate their ovulation ranged between 7.00-7.01 (Fig. 2). The mean water pH of the incubated eggs that were stripped as a result of using varying dosages of ovatide to stimulate their
The differences in the mean water pH of both hormones could be attributed to the weather and environmental conditions, presence of blood stains on the eggs, fatty content of the eggs and variation in water level of the hatchery tanks used. There was no significant difference (p>0.05) in the mean water pH among the varying doses of the hormones used. These results agreed with the work of Santhosth and Singh (2007) which reported that a suitable pH range for fish breeding was between 6.7-9.5. This work also agrees with the work of Kutwal et al., (2015) which reported that pH of 6.0 favoured the breeding of Clarias gariepinus using Carp pituitary extract and ovulin-L. Isa et al., (2015) worked on the effect of monthly variation in water temperature on artificial breeding of common carp (Cyprinus carpio) on a pH range of 7.97-8.09. Muhammad et al., (2014) worked on the breeding of Sperata seenghala using different hormones such as ovaprim, Human Chorionic Gonadotropin, Leutennizing Releasing Hormone and ovatide on pH range of 7.0-7.5.

The dissolved oxygen of the water in which the eggs that were stripped as a result of using varying dosages of ovaprim were incubated ranged between 5.78-5.83mg/l. The maximum mean dissolved oxygen of 5.83mg/l was recorded in 0.3ml/kg of ovaprim (Fig. 3). On the other hand, the dissolved oxygen of the water in which the eggs that were stripped using varying dosages of ovatide were incubated ranged between 5.71-5.86mg/l. The maximum mean dissolved oxygen of 5.86mg/l in ovatide was recorded in a dose of 0.1ml/kg (Fig. 3). The differences in the mean dissolved oxygen of both hormones could be attributed to differences in the respiration of the incubated eggs, temperature of the hatchery tanks used, the influence of the prevailing atmospheric air pressure during the experiment, and continuous flowing of water in and out of the hatchery tanks. There was no significant difference (p>0.05) in the mean dissolved oxygen among all the varying doses of the hormones used. This study agrees with the work of Bhatnagar and Sangwan (2009) which reported that dissolved oxygen in the range of 4.5-8mg/l was suitable for fish breeding. This work is also similar to the findings of Tsadu et al., (2013) on induced breeding of Clarias anguilaris with Xenopus laevis (African clawed frog) crude pituitary glands at a dissolved oxygen level of 5.6-6.4mg/l. This study disagreed with the work of Ngueku (2015) on the efficacy of synthetic and non-synthetic hormones in the induced spawning of the African catfish at a dissolved oxygen level of 4.70mg/l; Nwadukwe (2013) on the breeding performance of Heterobranchus bidorsalis and Heterobranchus longifillos using three doses of ovaprim at a dissolved oxygen level of 1.64mg/l and that of Victor et al., (2016) which reported that a dissolved oxygen range of 2.20-5.60mg/l was observed during the study on the effect of water renewal on growth of Clarias gariepinus fingerlings.

The minimum and maximum mean incubation period of 20.66±40.00 and 23.00±25.00hours respectively were observed in ovaprim. On the other hand, a minimum and maximum mean incubation period of 20.83±38.00 and 22.83±32.00hours respectively were observed in ovatide (Table 1). There was a significant difference (p<0.05) in the mean incubation period (hours) of the eggs that were induced with varying doses of ovaprim and ovatide. The difference in the mean incubation period (hours) could be attributed to the effects of oxygen, temperature, pH and dissolved oxygen during incubation of the eggs and water levels in the hatchery tanks used. These results are in agreement with the work of Ngueke (2015) which reported that the incubation period of African catfish eggs was between 20.00-21.00hours. Similarly, a research conducted by Ayoola et al., (2012) reported a hatching period of 21-26 hours on the eggs of Clarias gariepinus. Sharma et al., (2010) reported an incubation period of 46.0 hours, 44.5 hours and 42.5 hours in Clarias batrachus injected with 0.8ml/kg, 1.0ml/kg and 0.6ml/kg (doses of ovatide) respectively. The differences in the mean incubation period (hours) obtained from this study with that of others compared could be attributed to environmental conditions and species of the fish used.

5. Conclusion

Fertilization and hatchability rates of the eggs of Clarias gariepinus were low at higher temperature in eggs that were induced with ovaprim, and on the other hand were at their best in those induced with ovatide. A pH range of 6.94-7.01 observed in this study favoured the fertilization and hatchability rates of the eggs that were induced with both ovaprim and ovatide. The dissolved oxygen range of 5.78-5.83mg/l observed in this study favoured fertilization and hatchability in the eggs that were induced with both varying doses of ovaprim and ovatide. The incubation period (Hours) range of
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20.66±40.00-23.00±25.00 hours gave a good hatchability rate in all the eggs that were induced with both varying doses of ovaprim and ovatide.

ACKNOWLEDGMENTS

Our appreciation goes to the Head, Department of Biological Sciences, Benue State University, Makurdi for lending us some equipment that were used in the course of this work.

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