Comparative breeding performances of indigenous climbing perch (Anabas testudineus) populations in a newly developed semi-artificial condition

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Abstract. The indigenous Anabas testudineus populations from two different regions were bred in a newly developed semi-artificial breeding condition in modified 500L plastic tanks, with inlet and outlet systems having continuous water supply facilities. The induced breeding was performed through synthetic hormone “Flash” and a single dose was used as 0.2 ml/kg for females and 0.1 ml/kg for male fishes. The experiment was set with four treatments such as T₁ Purbred: (Netrokona ♂ × Netrokona ♀), T₂ Purbred: (Sylhet ♂ × Sylhet ♀), T₃ Crossbred (Sylhet ♂ × Netrokona ♀), and T₄ Crossbred (Netrokona ♂ × Sylhet ♀) with three replications each. The ovulation rate was higher in T₃ for the Sylhet female population (445.358 ± 14.18/gram body weight). The fecundity of Sylhet female (T₂ and T₃) was 24000-30000, and the fecundity of Netrokona female (T₁ and T₄) was 15000-20000. The highest and lowest fecundity was found in T₃ (29444±1988) and T₁ (16888±346), respectively. The highest fertilization rate was recorded (90.41±2.10%) in T₂ and (89.46±41.50%) T₄, while the lowest fertilization rate was 84.58±2.40% in T₃. The highest hatching rate was 77.16±2.33% in T₂ and the lowest hatching rate was 42.23±1.58% in T₃. Also, a higher survival rate was found in T₂. The present study indicated that the breeding performances were better in the Sylhet population than the Netrokona population in the developed semi-artificial condition.

Keywords: Anabas testudineus, Induced breeding, Plastic tank incubator,

Introduction

The climbing perch, Anabas testudineus (Bloch, 1792) of the family Anabantidae, is popularly known as “koi” in Bangladesh is a small-sized food fish inhabiting both freshwater and brackishwater in the most tropical or subtropical areas (IUCN 2015, Talwar and Jhingran 1991). Among SIS (small indigenous species), A. testudineus has been considered as one of the potential candidates for aquaculture and captive breeding (Ponniah and Sarkar 2000). Once upon a time, indigenous climbing perch was very much abundant in almost all freshwater systems of Bangladesh (Mahmood 2003) but its availability has been declined in recent years. Sarkar et al. (2005) and Bhattacharyya and Homechaudhuri (2009) optimized induced breeding technique of this species using different doses of a synthetic hormone, Wova-FH, and Ovaprim, respectively. Seed production technology of A. testudineus through artificial propagation was developed in captive conditions (Kohinoor 1991, 2008), but the growth rate of native strain is very slow in the pond’s ecosystem.

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In 2002, a fast growing strain of A. testudineus known as Thai koi was introduced in Bangladesh for culture purposes. Due to the failure of maintaining a proper hatchery protocol in the fry production phase, inbreeding has resulted in reducing the high yielding characteristics of Thai koi (Kohinoor and Zaher 2006). To overcome this situation, Sharnalata Agro Fisheries Limited introduced another very fast growing Vietnam koi in Bangladesh in 2010, for its higher production and growth than the other variety of climbing perch (Shafiquzzoha et al. 2018, Hafijunnahar et al. 2016). It was observed that Vietnam koi grows as much as 250-300 g within 120 days of culture period, and the body color is almost similar to native koi (Amin et al., 2015). Nevertheless, consumers do not widely accept this fish due to lack of taste and natural odour compared to native koi. Consequently, induced breeding of indigenous climbing perch needs to be prioritized, and breeding technique needs to be improved.

One of the major constraints in breeding this species is the collection of quality brood fish from nature without any stock information. In the previous experiment (Alam et al., unpublished data), quality brood fish of native koi was identified, and broodstock was developed for seed production. The population from Netrokona and Sylhet in Bangladesh was considered good brood fish for breeding performances. Thus, this study aimed to study comparative breeding performances of two different populations of Bangladeshi A. testudineus and develop a semi-artificial breeding technique for captive conditions.

Materials and Methods

Broodfish rearing and breeding facilities development: Broodfish rearing and breeding facilities were developed before starting the experiment, besides the faculty building of Bangabandhu Sheikh Mujibur Rahman Agricultural University, Gazipur. The mini hatchery has six cemented cisterns (5’ × 3’ × 4’) for brood fish rearing, two (6’ × 5’ × 3’) cisterns with bottom mud for hatchling rearing of indigenous climbing perch. A plastic tank (500L) was used for supplying water holding in the rooftop in a house was used as an incubator for spawning and hatching fish after modification. The upper side of the tank (6 inches) was cut and removed, and it was then open on the upper side. Inside the tank outlet, a short piece of pipe (with a diameter of 0.5-inches and a length of 15 inches) was horizontally set and came to the center of the tank and then using another short piece of pipe (15 inches) used vertically with the elbow. The outside outlet of the tank was also connected with another short piece of pipe having a height of 15 inches. Thus, the water level was maintained by lowering both the inside and outside vertical pipe upward and downward. The mouth of the inlet and outlet pipe was covered by a fine mesh size net so that the egg cannot come outside of the tank along with the water flow. The upper opening mouth side of the tank was also covered by a fine mesh size net so that fish can’t escape from the tank. The tank was then set under a horizontal water supply system by a short porous piece of pipe (36 inches) with the dead-end mouth. Thus, twelve plastic tanks were prepared and used for breeding in the present experiment. Digital electric balance was used for measuring fish weight. A plastic bowl, petri dish, and beaker were used to count the number of eggs, fertilization, and hatchling.

Brood fish collection and rearing: Five hundred brood fish of native koi from each of the two populations Netrokona (Bandha beel, Mohonganj) and Sylhet (Murier haor, Golapganj) were collected in January 2017, and the fish were reared and kept in the cemented tank (5’ × 3’ ×
having a water height of around 3 feet. Approximately 120-150 fish were kept in each
cemented tank, and the fishes were fed with a commercially available feed twice daily at 5%
body weight. The broodfish were fed a VitE supplement with the feed. The excess amount of
feed and excreta was cleaned every day, and the whole water of the tank was changed twice a
week.

**Experimental design:** The experiment was designed with four treatments T₁, T₂, T₃, and T₄
(treatments T₁ - Netrokona ♀ × Netrokona ♂, T₂ - Sylhet ♀ × Sylhet ♂, T₃ - Sylhet ♀ × Netrokona
♂, T₄ - Netrokona ♀ × Sylhet ♂) for two populations, and each treatment has three replications
(R₁, R₂, and R₃). Temperature and dissolved oxygen were recorded a similar (p >0.05) in all
treatments throughout the research period. The average temperature and dissolved oxygen with
standard deviation (SD) was 28.8±0.5 and 5.4±0.3, 29.3±0.5 and 5.1±0.3, 30.1±0.5 and
5.2±0.3, and 30.2.8±0.5 and 5.4±0.3, in T₁, T₂, T₃, and T₄, respectively.

**Broodfish response in captivity and breeding trial:** Broodfish maturity was checked weekly
since March 2017 to observe whether eggs and sperm come out (Fig. 1) after gentle pressure in
the abdomen. Thus, the maturity check continued, and the first maturity response was found in
the first week of April. One breeding trial, including broodfish selection, hormone injection,
ovulation, incubation, fertilization, hatching, all steps took around two weeks. Thus, six
breeding trials were done from April to June. The breeding response was found the best in the
middle of this period. However, the best three breeding trials were used in the present
experiment (Table I). The male and female fish weights varied from 27-51 g and 37-95 g. The
average weight (g) of brood fishes used in different trails is given in Table I. In each treatment,
three pairs of male and female fishes were used, and thus, in each trial, 72 fish were used in the
four treatments (T₁, T₂, T₃, and T₄).

![Fig. 1. Milt and eggs come out upon gentle pressure on the abdomen of male (A) and female (B) koi.](image)
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Table I. Average weight (g) of brood fish koi, *A. testudineus* used in different trails (N = 216)

<table>
<thead>
<tr>
<th>Date</th>
<th>Weight (g)</th>
<th>T₁</th>
<th>T₂</th>
<th>T₃</th>
<th>T₄</th>
</tr>
</thead>
<tbody>
<tr>
<td>27-04-17</td>
<td>Male</td>
<td>41.0</td>
<td>38.0</td>
<td>37.6</td>
<td>41.6</td>
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<tr>
<td></td>
<td>Female</td>
<td>70.5</td>
<td>63.0</td>
<td>73.4</td>
<td>50.3</td>
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<tr>
<td>16-05-17</td>
<td>Male</td>
<td>49.6</td>
<td>36.2</td>
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<tr>
<td></td>
<td>Female</td>
<td>80.0</td>
<td>54.0</td>
<td>62.6</td>
<td>49.8</td>
</tr>
<tr>
<td>07-06-17</td>
<td>Male</td>
<td>38.0</td>
<td>34.5</td>
<td>39.0</td>
<td>40.0</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>64.4</td>
<td>51.3</td>
<td>62.6</td>
<td>62.6</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Mean body weight (±SD)</th>
<th>T₁</th>
<th>T₂</th>
<th>T₃</th>
<th>T₄</th>
</tr>
</thead>
<tbody>
<tr>
<td>27-04-17</td>
<td>Male</td>
<td>42.87 ± 6.02</td>
<td>36.23 ± 1.75</td>
<td>37.73 ± 1.20</td>
<td>39.07 ± 3.10</td>
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<tr>
<td></td>
<td>Female</td>
<td>71.62 ± 17.01</td>
<td>56.08 ± 12.31</td>
<td>66.22 ± 10.38</td>
<td>54.22 ± 10.59</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Mean fecundity (±SD)</th>
<th>T₁</th>
<th>T₂</th>
<th>T₃</th>
<th>T₄</th>
</tr>
</thead>
<tbody>
<tr>
<td>27-04-17</td>
<td>16888.67 ± 346b</td>
<td>24291.67 ± 355a</td>
<td>29444 ± 199a</td>
<td>18555 ± 376b</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Ovulation rate (No. of egg released/g of fish)</th>
<th>T₁</th>
<th>T₂</th>
<th>T₃</th>
<th>T₄</th>
</tr>
</thead>
<tbody>
<tr>
<td>27-04-17</td>
<td>238.54 ± 31.07c</td>
<td>431.00 ± 16.81a</td>
<td>445.36 ± 14.18a</td>
<td>344.44 ± 80.63b</td>
<td></td>
</tr>
</tbody>
</table>

(T₁-Netrokona ♀ × Netrokona ♂, T₂-Sylhet ♀ × Sylhet ♂, T₃-Sylhet ♀ × Netrokona ♂, T₄-Netrokona ♀ × Sylhet ♂)

Different superscripts in the same column indicate significant differences among different treatments (p<0.05);
One-way ANOVA followed by LSD multiple range test.

*Inducing agent and injecting the broods:* The broodfish were selected by observing external morphology. Synthetic commercial hormone Flash (Synthetic Gonadotropin Releasing Hormone Analogue, SGnRH, 10ml vial) was used. Each 10ml hormone contains analogue 0.002%, Domperidone 0.998%, Propylene glycol 99%. The single dose was used as 0.2 ml/kg for females and 0.1 ml/kg for male fishes. At first, female fishes were selected and weighed by the electric balance (Fig. 2A), and kept in a plastic bucket for a short time; and at the same time, male fishes were selected and kept in a separate plastic bucket before hormone injection. The intramuscular injection of hormone was administered at the base of the pectoral fin gently (Fig. 2A).

![Fig. 2. Hormone injection to *A. testudineus* (A) and newly hatched larvae (B).](image)
**Hatching and rearing:** Eggs and milt were released from the injected fish naturally, and fertilization occurred in the tank. It was taken around 24 to 30 hours for incubation and hatching of fertilized eggs (Fig. 2B). The hatchlings did not take food from outside until the absorption of their yolk sac, and it took around three days to start feeding on the outside as the first feed. The hard-boiled chicken egg was given as the first feed for around three days. The hard-boiled yellow part of the egg was smashed with water and sieved by fine microjorjet cloth and supplied at the rate of one egg/20 tank and twice daily. After 5-6 days of hatching, commercially available nursery feed (Mega nursery powder feed) containing 40% protein was given in each tank. The dead hatchlings were removed carefully from the tank, and a continuous water supply was maintained; and thus, fish spawns were reared in the tank for about two weeks. Uneaten particles and dirty materials were removed periodically, and the bottom was cleaned by siphoning. After 14 days, the fry was transferred into the bottom mud cemented tank (6’ × 5’ × 3’) and reared for one month.

**Assessment of breeding parameters of A. testudineus:** The total number of eggs and the number of fertilized eggs were calculated by the volumetric method after ovulation and fertilization in the 500L tank. At first, water containing fertilized eggs from 500L tank were taken in a 1 Liter biker and counted total eggs and fertilized eggs and recorded. This way, it counted ten times, and the average number was multiplied with the total volume of the water (average 50L) in the tank, and approximate total egg number and fertilized egg were estimated. However, fertilized eggs were transparent, while unfertilized eggs were opaque in appearance. The fertilization rate was determined by the following formula:

\[
\text{Fertilization rate (\%)} = \frac{\text{No. of fertilized eggs}}{\text{Total no. eggs (fertilized and unfertilized)}} \times 100
\]

The hatched number was counted in the same way mentioned above, and the hatching rate was determined by the following formula:

\[
\text{Hatching rate (\%)} = \frac{\text{No. of eggs hatched}}{\text{Total no. of eggs}} \times 100
\]

The survival number of spawns was recorded after 4, 7, 14 days after hatching following the method described above, and the survival rate was determined by the following formula:

\[
\text{Survival rate (\%)} = \frac{\text{No. of survived}}{\text{Total no. of eggs}} \times 100
\]

**Statistical analysis:** The gathered data were summarized and scrutinized consciously before actual tabulation. After data entry, the data were analyzed by Statistics 10.0, and the results found in the experiment were subjected to statistical analysis, ANOVA (one way), that showed the significance (\(p<0.05\)) level of differences between treatments. Significant results (\(p<0.05\)) were further tested by the LSD multiple range test.

**Results**

**Fecundity and ovulation:** Fecundity is one of the important factors of the biology and population dynamics of fish. In the study, induced breeding was achieved successfully in all four
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treatments and, the latency and incubation period was 8-12 and 24-30 hrs, respectively. The highest number of the egg was found in T₃, and the lowest was found in T₁. In the present study, fecundity varied from 15000-30000. The fecundity was always found higher in Sylhet females. The mean fecundity of Sylhet females was 24000-30000 in treatment T₂ and T₃. On the other hand, the mean fecundity of Netrokona females is 15000-20000 in treatment T₁ and T₄. Also, the ovulation rate was higher in T₃ (445.358±14.182) compared to ovulation rates found in T₁ (238.54±31.07), T₂ (431.00±16.81), and T₄ (344.44±80.63), respectively (Table II).

**Table II. Showing details of induced breeding performances of koi *A. testudineus***

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Fertilization rate (%) (±SE)</th>
<th>Hatching rate - Day 1 (%) (±SE)</th>
<th>Day 4 (%) (±SE)</th>
<th>Day 7 (%) (±SE)</th>
<th>Day 14 (%) (±SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>T₁</td>
<td>88.07 ± 1.48ᵇ</td>
<td>56.25 ± 0.19ᵇ</td>
<td>23.03 ± 0.86ᵇ</td>
<td>14.75 ± 0.31ᵇ</td>
<td>9.32 ± 0.31ᵇ</td>
</tr>
<tr>
<td>T₂</td>
<td>90.41 ± 2.10ᵃ</td>
<td>77.16 ± 2.33ᵃ</td>
<td>32.79 ± 0.90ᵃ</td>
<td>22.57 ± 0.67ᵃ</td>
<td>11.9 ± 0.45ᵃ</td>
</tr>
<tr>
<td>T₃</td>
<td>84.58 ± 2.41ᵇ</td>
<td>42.23 ± 1.58ᵇ</td>
<td>21.66 ± 1.01ᵇ</td>
<td>13.50 ± 0.43ᵇ</td>
<td>7.19 ± 0.34ᵇ</td>
</tr>
<tr>
<td>T₄</td>
<td>89.47 ± 1.51ᵇ</td>
<td>58.34 ± 3.97ᵇ</td>
<td>31.95 ± 0.62ᵇ</td>
<td>19.49 ± 1.23ᵇ</td>
<td>11.68 ± 1.05ᵇ</td>
</tr>
</tbody>
</table>

(T₁-Netrokona ♀ × Netrokona ♂, T₂-Sylhet ♀ × Sylhet ♂, T₃-Sylhet ♀ × Netrokona ♂, T₄-Netrokona ♀ × Sylhet ♂)
Different superscripts in the same column indicate significant differences among different treatments (*p*<0.05); One-way ANOVA followed by LSD multiple range test.

**Fertilization and hatching rate:** From the experiment, the fertilization rate was recorded as 88.00±1.49%, 90.41±2.10%, 84.58±2.41%, and 89.47±1.51% in the treatments of T₁, T₂, T₃, and T₄, respectively (Table II). The highest fertilization rate, 90.41±2.10%, was recorded in T₂ whereas the lowest fertilization rate, 84.58±2.41%, was found in T₃. The hatching rate was found 56.25±0.19%, 77.16±2.33%, 42.23±1.58%, and 58.34±3.97% in treatments of T₁, T₂, T₃, and T₄, respectively (Table II). The highest hatching rate was recorded (77.16±2.33%) in T₂ and the lowest hatching rate was recorded (42.23±1.584%) in treatment T₃ (Table II).

**Survival rate:** The survival rate of koi larvae in four different treatments (Table II) were 23.03±0.86%, 32.79±0.90%, 21.66±1.012%, and 31.95±0.62% in T₁, T₂, T₃, and T₄, respectively after four days of hatching. The number of koi larvae gradually decreased after seven days in all treatments. The survival rate of seven-day-old koi larvae was 14.75±0.31%, 22.57±0.67%, 13.50±0.43 %, and 19.49±1.23% in T₁, T₂, T₃, and T₄, respectively. After 14 days of hatching, the survival rate was 9.32±0.31 %, 11.9±0.45 %, 7.19±0.34%, and 11.68±1.05% in T₁, T₂, T₃, and T₄, respectively (Table II).

**Discussion**

Quality brood fish is the most important for effective breeding performance (Ingram and Nguyen 2014). Kohinoor (2008) collected native koi (*A. testudineus*) from natural sources, but there was no detailed information on the origin of broodfish. On the other hand, in the present experiment, the brood fish was collected from the haor area of Netrokona and Sylhet as comparative genetic variation was found better in these two populations in the previous molecular work (Alam et al., unpublished). According to Kohinoor (2008), induced breeding of native koi was conducted from March to July, keeping the Male and female ratio as 1:1, whereas at present, the best-
induced breeding period was observed from April to June, maintaining the male and female ratio as 1:1 in 500L plastic tank. Sarkar et al. (2005) said that the brood fishes of both males and females were collected from the river Punarbhava, located in Maldah district, West Bengal, from December to February. Chaturvedi et al. (2015) collected mature and healthy brood fish of A. testudineus from a private fish farm for breeding research. Samarendra et al. (2016) conducted induced breeding of A. testudineus, but no detailed information was found about the brood fish source. In the case of induced breeding of Vietnamese koi, brood fish were collected from Mohalakhsmi Fish Hatchery, Bogra, and another from its own farm’s pond (Amin et al., 2015).

The rearing of brood fish has a strong impact on breeding response. In the present experiment, 120-150 broodfish were stocked in a cemented tank (5×3×4 feet) with around 2 feet water height, whereas Kohinoor et al. (1991) reared two hundred brood fish in a 280 m² earthen rearing pond. Chaturvedi et al. (2015) kept brood fish in a cemented cistern size (3×2×1 m) with a water depth of 10-12 inches. Sarker et al. (2005) said that the brood fish were maintained in the earthen small-sized pond (0.05 ha, average depth 70-80 cm). Suraiya et al. (2012) broodstocks were reared in rectangular ponds of size 18 ×14 m² and an average depth of 1.3 m. Sarkar et al. (2005) conducted captive breeding of A. testudineus in separate nylon hapa. Akhtar et al. (2014) carried out induced breeding in a cemented tank filled with 30 cm of water (10ft /8ft). A 500 L modified plastic tank was used for ovulation and fertilization, where induced breeding successfully occurred in the present experiment. There is an effect of maturation of the brood on ovulation, fertilization, hatching rate, and also on the survival rate of larvae. Ananda (1973) observed that though many adult females of carp attain the age of maturity, their gonads were still in the immature stage with a lot of fat accumulated in the body cavity. Though the external feature can determine the maturation, the acentric or peripheral location of nuclei is a definite indication of fish readiness for spawning. If the ova nuclei are found to be centrally located, then such a fish would not respond to hormonal breeding treatment (Jhingran and Pullin 1990). Immature eggs fail to get fertilized, and while fertilized, eggs sometimes produce deformed spores that do not hatch out. In the present experiment, since March, weekly checking of the male and female maturity was observed after gentle pressure on the abdomen whereas sperm and egg were come out or not and thus maturity was confirmed.

The administration of the appropriate dose of hormone is the basic to the success of induced breeding; the condition of brood fish and the environmental conditions are also equally important (Pillay 1993). The experiment was carried out at the constant dose of synthetic hormone dose of 0.2 ml/kg of body weight for females and 0.1 ml/kg of body weight for males following the information provided by the company. On the other hand, Kohinoor (2008) used the pituitary gland (PG), and the dose was 8-10 mg/kg for females and 4mg/kg for male koi fish. Samarendra et al. (2016) used synthetic hormone WoVA-FH, and the dose was 0.4 ml/kg for males and 0.9ml/kg female. Fecundity is the number of ripening eggs in the female and varied from species, age, length, weight, and environment (Sivashanthini et al., 2008). Fecundity increases with the increase of length, weight, and egg size (Ghafari and Jamili 2010, Lawson, 2011). The early report showed that A. Testudineus, a freshwater fish, had high fecundity (Amornsakun et al., 2005). In this study, the mean fecundity of Sylhet female was 24000-30000 in fish weight range 44-89 g, and the mean fecundity of Netrokona female was 15000-20000 in fish weight range 37-95 g. Sarkar et al. (2005) reported that the number of eggs
released by the female ranged from 52000 to 130000, indicating high fecundity. Khan and Mukhopadhyya (1972) observed fecundity ranging from 10002 to 36477 in the fish size range of 99 to 169 mm. However, Banerjee and Prasad (1974) reported the fecundity of 4588-34998 in the Bihar region in the fish size range 84-100.2 g. Chanchal et al. (1978) reported a minimum of 3481 to a maximum of 42564 in the fish weight range of 9.0-53.1 g. Banerjee and Thakur (1981) reported 2000-13000 eggs in seven sets of induced bred A. testudineus (24.8-40.1 g).

According to Sarkar et al. (2005), the fertilization rates in A. testudineus were highest (98.50±3.5%) treated with Wova-FH at female 0.3 ml/kg and male 0.3 ml/kg dosage, respectively. In this experiment, the highest fertilization rate was 90.41±2.10%, and the lowest fertilization rate 84.58±2.41%. The highest rate of fertilization was occurred by Sylhet ♀ × Sylhet ♂ and the lowest by Sylhet ♀ × Netrokona ♂. Though Sylhet females had the highest fecundity in Sylhet ♀ × Netrokona ♂ found the lowest fertilization. It might be due to the poor quality of sperm. Therefore, further research on sperm quality may be undertaken. Furthermore, Zalina et al. (2012) reported the highest rate of fertilization (98.47±0.45%) in A. testudineus injected LHRHa 2 µg/kg whereas, lower fertilization rates tabulated than the present study as 71.00±3.97% and 78.11±3.24% in A. testudineus injected with PG at 2 mg/Kg for male and 7 mg/Kg for female (Amin et al., 2015). Similarly, Sarkar et al. (2005) reported the highest hatching rate, which was 90.50±3.65 using Wova-FH at 0.3 ml/kg for both males and females. Chaturvedi et al. (2015) also reported a hatching rate of 90% for A. testudineus injected with ovatide at 0.6 ml/kg for females and 0.4 ml/kg for the male. Zalina et al. (2012) reported the lowest hatching rate of 56.52±1.35, 59.61±2.18, and 65.33±2.69 in A. testudineus injected with different doses of LHRHa. Bhattacharyya and Homechauhdhuri (2009) conducted captive breeding of A. testudineus with the synthetic hormone Ovaprim with dose as injected 2 ml/kg body weight.

In the present experiment, the highest hatching rate was recorded (77.16±2.330%), and the lowest hatching rate was recorded (42.23±1.58%). Sarkar et al. (2005) reported the highest hatching rate found 90.5 ± 3.65 using Wova-FH at 0.3 ml/kg for both males and females. Chaturvedi et al. (2015) also reported a hatching rate of 90% for A. testudineus injected with ovatide at 0.6 ml/kg for females and 0.4 ml/kg for the male. Zalina et al. (2012) reported the lowest rate of hatching 56.52±1.35, 59.61±2.18, and 65.33±2.69, respectively, in A. testudineus injected with different doses of LHRHa. Bhattacharyya and Homechauhdhuri (2009) stated that hatching was 68.57% in the breeding experiments with Ovaprim. The best survival rate of 14 days old koi (A. testudineus) was recorded in male and female combinations of the Sylhet population, indicating the best broodfish source for induced breeding. However, in most previous reports, there was no data about the survivability of koi fry, and present research might help increase the survivability of koi fry during nursing management in future studies.

Native koi fish (A. testudineus) has a high market price but still could not be brought into commercial culture due to lack of fish seed through induced breeding as well as slow growing pattern. These results showed that the fertilization, hatching, and survival are comparatively higher in T2. The fecundity and egg/gram of body weight are also higher in the Sylhet population rather than the Netrokona population. Finally, these results suggest that the breeding performances of the Sylhet population are better than the Netrokona population. In the present experiment, the breeding of two natural populations of koi fish was compared, and the source of
quality brood fish was tried to identify, and low-cost breeding techniques were also developed. However, a semi-artificial breeding technique using a 500L plastic tank and further rearing of hatching in the cemented tank was used, but the survival rate of fish fry was unsatisfactory, which may be due to a lack of live food and a new environment in the plastic tank. Therefore, further monitoring of these issues and continuous domestication with live food in the new environment may reduce the mortality of the native *A. testudineus* in the newly developed incubation system.

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**Literature Cited**


BREEDING PERFORMANCES OF A. TESTUDINEUS IN A MODIFIED CONDITION


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