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Initial Observations in Himri (Barbus lutes, Heckel) Propagation

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Abstract

Reaction of himri (Barbus lutes) to carp pituitary extract injections tested month and a half before its natural spawning. Females treated with two consecutive injections resulted higher relative fecundity (7464.96 ± 481 eggs) than those treated with only one injection (5286.62 ± 392.5 eggs). No significant differences noticed between treated (22.5 ± 0.4 ml) and non-treated males (23.2 ± 1.6 ml). High fertilization (98.6-97.8 %) could not attain satisfactory hatching (57 %). Colors modifications and acoustic acts have been observed during ripping. Spawned egg has 1.57±0.3×10³ g weight and 1.10-1.18 mm diameter. Spermatozoon motility increased significantly (45%) in hypotonic solution (150 mOsm.kg⁻¹). Embryonic development was similar to other cyprinids in durations, but the important stage of eye pigmentation has not been noticed even after hatching which occurred after about 74 h.

Key Words : Barbus lutes, propagation, genital products, embryo, Euphrates-Tigris.

Introduction

Himri (Barbus lutes, Heckel) is indigenous cyprinid in the basin of Mesopotamia. Beckman (1962) reported its presence in Orontes basin, but it hasn’t proved again.

Their tasty scented flesh raises popular demand on this fish. Some signs and observations indicate to promising possibility of being used for aquaculture in polyculture ponds, since it is herbivorous and detritus feeder (Epler et al., 2001a). Adaptation to earthen ponds with common carp Cyprinus carpio and other cyprinids has been noticed when it entered accidentally with water flux to ponds nearby Euphrates River (Al - Daham et al., 1991) which could consider it as a new species for the aquaculture practices in spite of clear decrease in body growth after the second year of life (Szupla et al., 2001)

Few investigations have been performed to recognize himri’s reproductive biology. It matures within 1 - 2 years of age at 14 cm average total length (Epler et al., 2001b). Its spawning in the middle and lower Euphrates occurs between June and July among waterweeds and roots of the floating plants. It has high fecundity and extended ovulation period for many days, so Al-Daham and Bhatti (1979) describe it as partial-spawner. Histo logical gonad analysis in both sexes showed annual changes lead to extended spawning season (Epler et al., 2001b; Al-Daham and Bhatti, 1979; Bhatti and Al-Daham, 1978) whereof increasing the total fecundity of himri.

The aim of this study is to examine the response of himri to exogenous hormonal induction by common carp pituitary extract (CPE) for simple propagation practices to be used in fish hatcheries, some specifications of gametes to improve gamete management, and the embryonic development. All these studies performed with non-complicated techniques can be used in field.

Materials and Methods

Experiments performed in Mreaiya fish farm hatchery on the left bank of Euphrates River, Deir ez Zor, Syria with river water heated by solar energy from 4 to 11 May.

Animals

Sexually mature females (n=10) [355-395 g body weight (W), 23.2-27.5 cm standard length (SL)] and males (n=9) [110-413 g (W), 21.8-26.5 (SL)] caught by gill net (4x4) cm from Euphrates River and transported to the hatchery farm by hauler. Acclimation with hatchery water was adjusted to (20-22°C) temperature in (6.5-7.5) pH.

Induced spawning

Sex and weights of broods were determined, tagged and divided into two groups. Broods in group I (G. I) injected with CPE at 3 mg.kg⁻¹ Body weight
(BW) in 1 ml saline as preparatory hormone treatment. After 10.5 h of the injection of G. I broods, genital papillae of all females sutured and the decisive injection at 8 mg.kg⁻¹ BW in 1 ml saline applied just for females in G. I. Females of G. II were given 8 mg.kg⁻¹ BW in single injection. Males of G. II treated just with saline. All injections were intraperitoneal under pectoral fin's base. Broods were stocked together at 20-22°C and 2.5 L min⁻¹.kg⁻¹ BW. After 520-540 degree-hours (dh) since first injection; fish stripped by slight abdominal pressure. Eggs fertilized in dry conditions at ratio of 25 ml milt/1kg eggs.

Fertilized eggs swelled, desticked (Al Hasszaa and Hussein, 2003), then put into Zouger jars of 8 l. Water flow adjusted to 0.5-1 L min⁻¹, temperature and dissolved oxygen (DO) measured at hourly basis by using the oxygen meter W. T. W. type OXI-92. Turbidity and salinity of water estimated by Yellow Springs 6600 multi parameter at starting. Malachite green used as fungicide.

**Embryonic development**

Developmental stages of fertilized eggs observed and examined under stereomicroscope (4-7×) and microscope (10, 20, 40×). Observations documented in hourly basis and categorized according to distinguished development intervals.

**Gametes**

Egg (n=40) diameter measured by micrometer eyepiece attached to stereomicroscope and weight by quantitative analysis on electronic balance. Milt volume measured for each male. Sperm viability estimated in distilled water and saline (3g NaCl.l⁻¹ dist. water). This saline raised osmotic pressure of water to 150 mOsm.kg⁻¹.

Duration of motility and stages of viability examined under microscope on 60× and 100× after 10 seconds of motility initiation. Viability divided into three stages of mobility: fast forward (FF), rotational waves (RW), and then the last stage starting by tail immobility and vanished swinging in head till quite immobility (VD).

**Statistical analysis**

Student's T-test used to compare mean values of fecundity, fertilization, and hatching rates in the two treatments.

**Results**

**Induced spawning**

By watching spawning behavior, ripping irritations of broods noticed gradually about 10 h after second injection (the single injection in G. II) with some chafes among males and females. Males turned to reddish brown in anterior body part and greenish around caudal peduncle. Females were less pigmented than males. Acoustic behavioral changes observed in males during ripping. Ripping males emitted short and sharp pulse trains accompanied with springs over pond's water. These sounds were not aggressive behavior within or between sexes. Fertilization rate rose over 98.6 %, but hatching still less than 57.3 %.

Statistical analysis showed significant effect of two injections on female fecundity (P<0.01) since female's mean fecundity estimations were 11.72 g.kg⁻¹ BW in G. I and 8.3 g.kg⁻¹ BW in G. II in spite of the high quantities of unspent ova noticed in ovaries after post-stripping dissections. No differences in milt production observed between induced and non-induced males (P>0.01) as shown in (Table 1). Additional milt could be collected for second and third time after 6-10 h but decreased in volumes gradually.

**Embryonic development**

Embryonic development defined in Table (2) where embryos developed to final formation outward after 64 h of fertilization. Hatching started after 74 h of fertilization in 20-22°C; 6-11 mg.L⁻¹ DO; and neutral pH, but embryo's eyes still without pigmentation (Figure 1).

**Table 1.** Effect of hormonal stimulation by common carp pituitary extract on fecundity, fertility, and hatching of himri *Barbus luteus* at 20-22°C.

<table>
<thead>
<tr>
<th>Group</th>
<th>1st dose mg.kg⁻¹ (BW)</th>
<th>2nd dose mg.kg⁻¹ (BW)</th>
<th>Relative fecundity Mean ± S.D.</th>
<th>% Fertilization</th>
<th>Degree-day Mean ± S.D.</th>
<th>% Hatching</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Females (n=5) 3</td>
<td>8</td>
<td>7464.96 ± 481†</td>
<td>98.6 ± 1.2</td>
<td>62.5 ± 2</td>
<td>57.3 ± 1.2</td>
</tr>
<tr>
<td></td>
<td>Males (n=5) 3</td>
<td>-</td>
<td>22.5 ± 0.4†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Females (n=5) -</td>
<td>8</td>
<td>5286.62 ± 392.5†</td>
<td>97.8 ± 0.7</td>
<td>62.5 ± 2</td>
<td>56.8 ± 0.8</td>
</tr>
<tr>
<td></td>
<td>Males (n=4) -</td>
<td>-</td>
<td>23.2 ± 1.6†</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

†: Eggs.kg⁻¹ BW, ‡: ml milt.kg⁻¹ BW.
<table>
<thead>
<tr>
<th>Time</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 to 2.5 h</td>
<td>Segmentation, first cleavage followed within 30 min by second cleavage up to 8-cell stage</td>
</tr>
<tr>
<td>3.5 h</td>
<td>16-cell Morula, cells are medium in dimensions</td>
</tr>
<tr>
<td>6 h</td>
<td>Morula, small and many cells</td>
</tr>
<tr>
<td>10 h</td>
<td>Blastula</td>
</tr>
<tr>
<td>13 h</td>
<td>Gastrula, epiboly increased. Cells ring closing yolk sac.</td>
</tr>
<tr>
<td>26 h</td>
<td>Completion of yolk invasion, blastopore, and embryonic rudiments</td>
</tr>
<tr>
<td>29 h</td>
<td>3, 6 then 12 myotomes. Yolk sac elongation. Formation of intestine, notochord, and caudal bud.</td>
</tr>
<tr>
<td>31 h</td>
<td>Head and tail elongated, tail separating from yolk sac. Head is clearly developed and bent toward yolk. Colorless, slowly-circulating blood in rudiments. Sudden movements turning embryo around.</td>
</tr>
<tr>
<td>63 h</td>
<td>Development of Myotomes. Well developed eye lens present but unpigmented.</td>
</tr>
<tr>
<td>64 h</td>
<td>Final outward, tail moving freely. Continuous movement. 32 myotomes. Regular heart contractions and red blood circulation in vessels.</td>
</tr>
<tr>
<td>74 h</td>
<td>Hatching continued 6 h in our experiment. Final formation of hatching larva is straight, head is slightly bent towards yolk. Single chamber swim bladder. Total length 5 mm (Fig. 1)</td>
</tr>
</tbody>
</table>

*: approximately

Figure 1. Newly hatched himri ‘Barbus luteus’ larva, total length 5 mm. No pigmented optical vesicle could be noticed.

Gametes

Unfertilized eggs were demersal, spherical, adhesive and dark yellow in color with mean weight of $1.57 \pm 0.3 \times 10^3$ g and size range of 1.10-1.18 mm.

Milt look homogenous milky in color with mean pH value of 8.5. Viability of sperm rose up to 45% in hypotonic solution (150 mOsm.kg$^{-1}$) comparing with distilled water (Table 3).

Discussion

Himri showed positive response to exogenous stimulation by pituitary extract which appeared by accelerating gametogenesis and ability to spawn one and a half month before its natural spawning season. Emission of vocalizations by males near the incoming spawning time referring to acoustic mating behaviour in himri, but it has failed to make females release eggs due to suturing genital papilla. Such behaviour has been noticed in other species during spawning interactions in sand gopies Pomatoschistus marmoratus, P. canestrini, and Knipowitschia panizzae (Lugli and Torcelli, 1999), mating in Hypoplectrus unicolor (Lobel, 1992), before and during oviposition (Bisazza et al., 1989). This act may be positively affecting gamete production of both brood genders. It is recommended to observe the natural spawning of himri in natural habitats or mimicking conditions to discover the details of spawning acoustic behavior and its relation with female fecundity and spawned eggs. These acts could be manipulated to enhance total gamete production in propagation practices. However, Lugli et al. (1996) found that not all prespawning sounds may be a functional component of the spawning behaviour such as the male Arno goby Podogobius migricans.

Significant differences in fecundity between females received two doses of CPE of the priming dose lesser than one third of the last one and females received single dose indicates to use the CPE in two doses to attain good results in oocyte maturation, ovulation, and fecundity. Priming dose induces a significant GH increase one hour after the injection and then GH levels continue to increase for 3 hours and remain high during the next 12 hours (Weil et al.,

<table>
<thead>
<tr>
<th>Treatment</th>
<th>FF</th>
<th>RW</th>
<th>VD</th>
<th>Viability (min.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hatchery water</td>
<td>Min.</td>
<td>22</td>
<td>27</td>
<td>1.983</td>
</tr>
<tr>
<td></td>
<td>Max.</td>
<td>22.5</td>
<td>30</td>
<td>70</td>
</tr>
<tr>
<td>3g NaCl¹ water</td>
<td>Min.</td>
<td>28</td>
<td>47</td>
<td>98</td>
</tr>
<tr>
<td></td>
<td>Max.</td>
<td>32</td>
<td>54</td>
<td>100</td>
</tr>
<tr>
<td>Total increment %</td>
<td></td>
<td>27-42</td>
<td>74-80</td>
<td>33-40</td>
</tr>
</tbody>
</table>

1980). Main exogenous stimulation to induce the spawning in some cyprinids could be the second injection, in other cyprinids it could be unique and massive (some Chinese carps). We found that application of consecutive CPE doses as injected stimulator in himiri raised female fecundity more than single dose. Similar results have been reported in some other cyprinids such as Indian major carps *Catla catla*, *Labeo rohita*, *L. calbasu Cirrhinus mrigala* (Jhingran, 1986); Chinese carps *Ctenopharyngodon idella*, *Hypopthalmichthys molitrix*, and *Aristichthys nobilis*, (Shigang, 1989); tench *Tenca tenca*; (von Lukowicz et al., 1986); nase, *Chondrostoma nasus* (Szabó et al., 2002).

Himiri eggs are slightly different in size and weight which could be interspecific differences observable between individuals according to their age and weight. Differences in egg size and weight in many fishes have been related to spawning season (Bagenal, 1971; Papala et al., 1998), fish individual size (Bartel et al., 1999; Bonishawska et al., 2000; Marteinsdottir and Steinarsson, 1998), or brood protection (Tilney and Hecht, 1993).

Sperm production in males was not effected by CPE injection. However, males have produced in a single stripping more milt than Horváth et al. (1985) described for the common carp.

Collection of additional milt quantities and the high quantities of spent ova indicates that application of CPE in himiri propagation were not highly satisfactory. Thus, all future attempts to propagate wild himiri broods should consider that it is a partial-spawning species like bunni barbel *Barbus sharpeyi* (Al-Daham and Jasim, 2000) from the same habitat of himiri.

Embryonic development outlines in himiri look identical to other cyprinids reported by Peňáž (2001) from first cleavage to gastrula and organs formation, but appearance of optical vesicle without pigmentation till several days after hatching seems to be specific. Embryonic development calendar of himiri was almost same as bunni barbel described by Pyka et al. (2001) at same temperature, but absence of pigments in optical vesicle were exclusive in himiri.

Fertilized eggs of himiri hatched after about 64 degree-days (dd) in well oxygenated water (7-9 mg.l⁻¹). This is close to incubation period of bunni: 60 dd (Al-Nasih, 1992) to 70 dd (Pyka et al., 2001); common carp: 60-70 dd (Coch et al., 1998); and tench 60-70 (von Lukowicz et al., 1986).

Low hatching rate of himiri’s eggs couldn’t be explained. Similar problem noticed in bunni and gattan *Barbus xantopterus* (Pyka et al., 2001). Using other incubation techniques with investigating microbial loading of eggs and other ambient incubating conditions could raise hatching rates. Variations in egg dimensions could affect developmental strategy and hatching percentage in himiri.

Himiri’s spermatogenesis is generally similar to other teleost species (Bhatti and Al-Daham, 1978). Spermatozoa of many fish species including himiri are immotile in the testes and the genital tracts. They are activated only after release into external medium and have a short period of motility (Scott and Baynes, 1980). It was shown that a simple hypotonic shock of 60-100 mOsm.kg⁻¹ amplitude could activate common carp’s sperm (Billard et al., 1995). But dissimilarities of osmotic pressures in aqueous media according to variant concentrations of dissolved particles originate different responses of motility and viability of fish sperms. Billard (1978) also ascribed these variants to original structure of the spermatozoon. Intermix of teleost spermatozoa with ionic factors in aqueous media could depolarize sperm membrane which initiate motility, but it appears that freshwater is not the best media for the survival of the spermatozoa of freshwater fish (Billard, 1978). So, there is no use of this practice in artificial insemination. Osmotic pressure seems to be the major controlling factor in cyprinids sperms motility (Linhart et al., 1991; Morisawa et al., 1983). So, when himiri’s sperm activated by hypotonic Na⁺ solution (150 mOsm.kg⁻¹); we accomplished longer viability comparing with freshwater, which can raise fertilizing rates.

Acknowledgment

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