Determination of breeding season of endangered riverine catfish *Rita rita* (Hamilton, 1822) by studying ovarian development and Gonado-Somatic-Index

M. L. Rahman* and M. F. A. Mollah
Department of Fisheries Biology & Genetics, Bangladesh Agricultural University; Mymensingh-2202, Bangladesh
*E-mail: lrgolap_bau@yahoo.com

Abstract

*Rita rita* is an endangered riverine catfish having high market and nutritive values. The histological process of ovaries of *Rita rita* were performed to determine the breeding season for facilitating the artificial breeding in controlled environment. Fish were sampled monthly from the Brahmaputra River adjacent to Bangladesh Agricultural University (BAU), Mymensingh. The average Gonado-Somatic Index (GSI) ranged from 0.14 to 0.60 with the highest GSI value of 0.60 in July. The highest fecundity of 60114±3207.88 was found in the fish having length of 38.91±0.59 cm and weight of 1081.35±74.52 g. In contrast, the lowest fecundity was 37307.2±154.57 having length of 31.98±1.63 cm and weight of 810.87±49.75 g. The fecundity was found to increase with the increase of total length, body weight and the ovary weight and the relationship was linear and significant among them. Six developmental stages of ovaries were observed and categorized as the synchronous group. Vitellogenesis began in April with full development of ovaries in July. From this study, it can be presumed that gonadal development of *R. rita* peaked in July and breeding time within the months of June and July.

Keywords: Histology, *Rita rita*, Gonad, Gonado-somatic index

Introduction

The catfish *Rita rita* (Hamilton, 1822) is greenish brown carnivorous fish widely distributed in South Asia. It is one of the critically endangered species in Bangladesh (IUCN, 2003). The abundance of this species has been reported to decline day by day due to over-exploitation and various ecological changes in the natural aquatic ecosystem (Devi et al. 1991).

Knowledge on gonadal development and the spawning season of a species is necessary to determine the spawning frequency of its population as well as to establish a suitable induced breeding technique. The study of Gonado-Somatic Index (GSI) determines the state of maturity in terms of gonadal development and onset of spawning season. However, very few literatures are available on fecundity and GSI of *R. rita*. Although, monthly changes in the ovaries have been examined in some related species including brackish water catfish *Plotosus canius* (Khan et al., 2002), *Mystus gulio* (Sarker et al., 2002), *Clarias batrachus* (Faruq et al., 1996). A detail understanding of the gonadal development of a fish species is also considered an important step for the fish culturists. Histological observation may indicate the gonadal maturation and spawning season of a species.

Fecundity and gonadosomatic index have been useful tools to determine suitable spawning season of a particular species. The fecundity of fish can be estimated by actual counting method which is supposed to be the most accurate, but is very tedious, time consuming and to certain extent rather impossible for highly fecund fishes. Approximate fecundity is other alternative, which can be obtained by one of the following methods a) Volumetric method b) Gravimetric method c) Von vayer method as outlined by Lagler (1956). Gravimetric method was found to be more efficient than those of other methods and provided fairly accurate results (Islam and Hossain, 1990).

This species could be additional source of animal protein for local consumption, but it would have to be grown in controlled area like hatcheries, using low cost production methodology. To establish the induced breeding technique and culture in controlled environments of this riverine catfish, it is essential to know about its breeding biology and season.

In this present study fecundity and GSI was examined and breeding season was determined observing the gonadal development using histological process.
Materials and Methods

Study area and duration

The samples of *R. rita* (Hamilton, 1822) were collected from the old Brahmaputra river adjacent to BAU campus, Mymensingh. At least five females were collected every month during first year study period. The total length and body weight of individual fish were measured first and then the ovary of each fish was taken out carefully and weighed using a sensitive portable electronic balance (Denver Instrument, XP-300; 3000×0.1g) to estimate GSI. Finally ovaries were preserved in 10% buffered formalin for further study. The GSI is frequently applied to determine the spawning frequency of fishes and calculated according to the following formula:

$$\text{GSI} = \frac{\text{Weight of gonad}}{\text{Weight of body}} \times 100$$

Fecundity estimation

In this method, 20 mg sample of ovary was taken separately from anterior, middle and posterior portions of each ovarian lobe accurately. The number of mature and maturing eggs from each portion was found out separately by actual counting due to small amount. The mean number of eggs in 20 mg sample was determined and then multiplied by the total weight of the ovary, which gave the total number of eggs i.e. the absolute fecundity of respective fish. The relative fecundity also are calculated by absolute fecundity/body weight.

Histology of gonad

The preserved ovaries were taken out in a perforated plastic holder covering by perforated steel plates. Cleaning, infiltration and dehydration process were carried out in an automatic tissue processor unit using a series of alcohol of increasing concentrations, two times changes of xylene and finally through molten wax (three series).

Paraffin embedded blocks were cut by microtome knife at 4-5 µm size and left the sections into a water bath at a temperature of 40°C. The sections were placed on a glass slide and kept overnight on a slide drier hot plate at a temperature of 20°C. Then the sections were stained routinely with haematoxyline and eosin (Humason, 1972).

Microscopic examination of the gonadal tissue

The stained sections were mounted on the glass slide with Canada balsam and covered by cover slips and studied under a compound microscope (SWIFT M 4000-D). The photographic records were done simultaneously to study the different maturation stages oocyte and monthly developmental variations of ovary.

Data analysis

Microsoft Excel (2003) was used to determine linear relationship and correlation co-efficient (r) between total length and fecundity, body weight and fecundity, gonad weight and fecundity following Zar (1984).

Results

The female reproductive system

**The morphology of the ovary:** The ovaries of *R. rita* were paired tubular organs lying dorsal to the alimentary canal and ventral to the swim bladder. They were attached to the body cavity by the mesovarium. A posterior extension of tunica albuginea united both ovaries forming an oviduct opened to the exterior via the oval shaped urogenital papilla. They were usually equal in size but occasionally one was larger than the other.
Gonado-Somatic Index of *R. rita*: The GSI, the indicator of the status of gonadal development and maturity of individuals of experimental species, was calculated for female *R. rita* from January to December. Month wise changes in mean GSI values of female *R. rita* are presented in Fig. 1. Values of GSI ranged from 0.14±0.02 to 0.57±0.04. The higher values of GSI were observed during May to July ranging from 0.46±0.01 to 0.57±0.04. The GSI increased in every month from August to July.

Fecundity of *R. rita*: The fecundity was found to vary from 37307.2±154.57 to 60114±3207.88 in fish samples weighing 810.87±49.75 to 1081.35±74.52g with ovaries weight ranging from 3.28±0.02 to 6.17±0.78 g (Table 1).

### Table 1. Fecundity counts at various lengths ranges and number of ova per kg body weight of *R. rita*

<table>
<thead>
<tr>
<th>Frequency</th>
<th>Length range(cm)</th>
<th>Mean total length (cm)</th>
<th>Mean body weight (g)</th>
<th>Mean ovary weight (g)</th>
<th>Mean fecundity</th>
<th>No. of ova/kg body wt.</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>30-33</td>
<td>31.98</td>
<td>810.87</td>
<td>3.28</td>
<td>37307.62</td>
<td>46058.02</td>
</tr>
<tr>
<td>10</td>
<td>33-36</td>
<td>34.73</td>
<td>964.51</td>
<td>4.51</td>
<td>46737.38</td>
<td>48684.37</td>
</tr>
<tr>
<td>11</td>
<td>36-39</td>
<td>37.72</td>
<td>1058.19</td>
<td>5.74</td>
<td>56267.21</td>
<td>53587.62</td>
</tr>
</tbody>
</table>

Relationship between fecundity and total length: The linear relationship between log of fecundity and total length showed a positive correlation ($r = 0.9787$) (Fig. 2). The regression equation of fecundity with total length of fish could be expressed as:

$$\log_{10}Y = 1.0546 + 2.3429\log_{10}X$$

[Where, $Y =$ Fecundity, $X = $ Total length]

Relationship between fecundity and body weight: The logarithmic relationship of fecundity against body weight (Fig. 3) provided a positive linear correlation ($r = 0.9545$). The equation can be stated as:

$$\log_{10}Y = 0.3749 + 1.443 \log_{10}X$$

[Where, $Y = $ Fecundity, $X = $ Body weight]

A straight line through the origin would fit the points well, showing that the number of egg were directly proportional to the weight of the fish.

Relationship between fecundity and ovary weight: The fecundity increased progressively with ovary weight of the fishes (Fig. 4). The linear relationship between log fecundity and log ovary weight was positively correlated ($r = 0.9874$):

$$\log_{10}Y = 4.196 + 0.7274\log_{10}X$$

[Where, $Y = $ Fecundity, $X = $ Ovary weight]

Annual changes in the ovary

Development of ovary: Ovaries were internal, longitudinal and paired. A pair of mesentaries (mesovaria) suspended them dorso-laterally to the body cavity. This paired organ consisted of two ovarian lobes, which were separated by a septum. They were covered with an ovarian membrane and numerous ovarian lamellae, protected into ovarian cavity. The ovarian cavity connected the oviducts and the oviducts from each bilateral ovary joined to lead to the genital pore. The ovarian lamellae consisted of connective tissue lined by germinal epithelium, which contained cell nests of oogonia. Ovarian follicles developed along the lamellae and the vitellogenic oocytes were ovulated into the ovarian cavity. The follicles of vitellogenic and fully mature oocytes consisted of a transparent theca, granulosa and zona radiata. The size and extent of occupany of the body cavity varied with the stage and condition of sexual maturity of the female. In initial stage, the ovaries were thin, elongated, slightly flattened and semi transparent in appearance. Gradually they took characteristic and specific colour and the middle portion of the two ovarian lobes became broader than the anterior and posterior region that was determined by monthly observation of ovaries. Immediately prior to breeding season, especially from May to June the ovaries became much expanded and occupied almost the whole cavity. The two ovarian lobes in *R. rita* were equal in size. The anterior part of the ovaries of *R. rita* was more or less triangular. The colour of the developing and maturing ovaries of *R. rita* was creamy, brownish and yellowish respectively.
Fig. 1 Monthly variation of gonadosomatic index of female *R. rita*

\[ y = 1.4434x + 0.3749 \]
\[ r = 0.9545 \]

Fig. 2. Relationship between log of fecundity and log of total length of female *R. rita*

\[ y = 2.3429x + 1.0546 \]
\[ r = 0.9787 \]

Fig. 3. Relation between log of fecundity and log of body weight of female *R. rita*
Histological observation of ovary: From the histological study of ovaries at successive months, it was observed that oocytes did not develop synchronously and oocyte at various maturation stages were observed in paired ovaries in the month of early year where most of the ova were developed and full of yolk in the middle. In the present study, maturation stages of oocytes that indicates ovarian development in female *R. rita* shown in Figs. 5 (A, B, C, D, E, F).

Fig. 5A. Photomicrograph of a cross section of a rita ovary Early (PO₁) and Late (PO₂) perinucleolar stage of oocytes in the ovary of *Rita rita* in January and February. N = Nucleus with nucleoli (Nu) [Haematoxyline and eosin (H&E×200)]
Fig. 5B. Late perinucleolar stage of oocytes (PO\textsubscript{2}) in the ovary of *Rita rita* in March and April. N = Nucleus with nucleoli (H&E×200)

Fig. 5C. Yolk-vesicle stage of oocytes (YVO) in the ovary of *Rita rita* in May. T: theca, G: granulosa, Z: zona radiate (H&E×200)

Fig. 5D. Early yolk-granule stage of oocytes (EYGO) in the ovary of *R. rita* in June
The mean GSI values throughout the year during the study period showed the existence of one breeding season in *Rita rita*. The histological observations of monthly samples of ovary confirmed this finding. The GSI was calculated from January to December. There was a gradual rise in the values from April to July. The highest gonado-somatic index of *R. rita* was 0.57±0.03 in July and the second highest was 0.51±0.00 in June, indicating that *R. rita* may breed in June and July.

Fecundity may be expressed in terms of the number of eggs produced per brood fish in a breeding season (Lagler, 1949). It is sometimes referred to as total or absolute fecundity or more usually just as fecundity (Heese, 1990). The fecundity of *R. rita* varied from 37307±154 to 60114±3207 for the fish length from 31.97±0.63 to 38.91±0.59 cm and weighing from 810.87±49.75 to 1081.35±74.51 g. Azadi et al. (1987) recorded the fecundity to vary from 5,683 to 21,922 for *Heteropneustes fossilis* and relationship of fecundity-body length, fecundity-body weight and fecundity-ovary weight was found to be linear relationship. The present study also indicates that *R. rita* belonging to the same size group had varying number of eggs in their ovaries.

Azadi and Siddique (1986) also reported a linear relationship between fecundity-body length, fecundity-body weight, fecundity-standard length and fecundity-ovary weight for catfish *Heteropneustes fossilis* which is consistence with the present study. The relation between fecundity and ovary weight were found to be the most prominent among all the relationships and the correlation coefficient, r (0.9874) between fecundity and ovary weight was highly significant (P<0.01). These findings agree with the findings of Das et al. (1989) for *Heteropneustes fossilis*.

Histological observation shows the evidence of gonadal maturation and spawning season. Reproductive prospective of a population is one of the basic needs to designate the individuals of that population in respect to their gonadal conditions (Jhingran & Verma, 1972).

Ovarian development of *R. rita* was examined to study the pattern and timing of growth phase and maturation stages of germ cells in the gonad of female. It was observed that fish exclusively in immature stages (early and late perinucleolar stage oocytes) of maturity was mostly available in the months from January to April. Yolk- vesicle stage oocytes appeared in the month of May. Both yolk vesicle and yolk granular stages were found in the month of June but most of the eggs in the month of July were in yolk granule stage. These findings agree with the findings of Mollah (1986) reported for another catfish,
Determination of breeding season of endangered riverine catfish *Rita*

*Clarias macrocephalus*. Karamchandi & Motwani (1955) concluded from the larvae and juveniles collected during July and August that the fish most probably breed in the river Ganga from March to August. Finally it may be assumed that *R. rita* may breed in the months of June and July in the river old Brahmaputra in Bangladesh.

**Conclusion**

This study will be helpful for an aquaculturist who will try to breed *R. rita* by artificial or induced breeding technique. However, a complete picture of breeding season of *R. rita* can be obtained if samples from other river systems hosting *R. rita* are collected and studied similarly.

**Acknowledgements**

This work was made possible by the financial assistance of the University Grants Commission (UGC). The authors are also grateful to the Department of Aquaculture, Bangladesh Agricultural University, Mymensingh, Bangladesh for providing the laboratory facilities to the preparation of histological slides.

**References**


