Biochemical Changes in the Rumen Infecting Paramphistome, *Gastrothylax crumenifer* during Miracidial and Intramolluscan Developmental Stages

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**Abstract**

We hypothesize that the level of various biomolecules is variable and remain in a dynamic state during the course of development in *Gastrothylax crumenifer*. The aim of the present study is to determine whether these molecules are in dynamic state or their levels are constant in all the developmental stages. The present study was designed to investigate level of various macromolecules e.g. protein, glycogen, lipid, lipid fractions and nucleic acids by standard Spectrophotometric methods. All the biomacromolecules are expressed in terms of mg/g wet weight. A minimum of three separate replicates were run for each experiment. To investigate similarities/differences in polypeptides of different developmental stages namely fresh eggs, eggs containing mature miracidia (Em), cercariae, metacercariae, immature and adult stages, SDS-gradient PAGE was performed. In the present study we have investigated biochemical alterations during the larval stages of *Gastrothylax crumenifer* (a rumen infecting paramphistome) which revealed marked differences during the miracidial development as well as in the intra-molluscan larval stages. We observed that the protein, glycogen and lipid of freshly laid eggs were utilized during the development. Concentration of protein, glycogen and lipid contents significantly elevated in the cercariae as compared to the non-feeding metacercarial stage, indicating an adaptation of the parasite to build up the nutrients for the metacercariae which is a dormant stage. Very little amount of lipid was detected in the fresh eggs (E0) which further decreased in the eggs containing mature miracidia (Em) and then increased sharply in the cercarial and metacercarial stages indicating the substantial buildup of lipid reserves. Nucleic acid contents decreased from E0 to the subsequent developmental stages. Such changes could be of intrinsic significance for the cellular differentiation and organogenesis in larval paramphistomes. Further, we analyzed polypeptide profile of developmental stages which revealed heterogeneous mixture of polypeptides. A total of 27, 15, 14, 15 and 34 polypeptides in E0, Em, cercariae, metacercariae, immature and adult stages respectively were resolved. Eight and nine characteristic polypeptides (17-90 kDa) were observed in E0 and adult stages respectively.

**Introduction**

Paramphistomes are commonly occurring digenetic trematodes causing the disease ‘paramphistomosis’ in ruminants. The disease paramphistomosis causes high morbidity and mortality in tropical and subtropical countries resulting in great economic losses [1]. *Gastrothylax crumenifer* is an elongated circular paramphistome found in the rumen and reticulum in sheep, cattle and buffalo. The available literature and studies from our laboratory have indicated that 60-80% of the buffaloes slaughtered have paramphistome infections in various parts of India [2,3]. The problem of paramphistomosis has been comprehensively reviewed by Horak [4] and Dutt [5]. In a survey from our laboratory on the prevalence of paramphistomosis, 71.4% of buffaloes examined were infected with adults of *Paramphistomum Epiclitum* (51.9% prevalence), *G. crumenifer* (32.7%), *Orthocoelium solicoeloid* (16.1%), *Fischoederius elongatus* (6%), *Calicophoron papillosum* (5%), *C. calicophorum* (2%), *Olveria indica* (0.34%) and *Gigantoctyle explanatum* (19.6%) [6]. The adult worms inhabiting the rumen have low pathogenicity while the migrating immature stages cause severe pathological disturbances [4,7,8] including hemorrhagic inflammation in the alimentary tract, edema and anemia. The damage caused in ruminants badly influences production, since these parasites cause a lower feed conversion, weight loss, decrease in milk production and responsible for major economic losses [9].

During the completion of life cycle paramphistomes pass through a succession of environments i.e. free living to tissues of intermediate and definitive hosts undergoing multiplication for genome tract, edema and anemia. The damage caused in ruminants badly influences production, since these parasites cause a lower feed conversion, weight loss, decrease in milk production and responsible for major economic losses [9].

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To obtain metacercariae, the infected snails were kept in a beaker containing water and grass blades. The emerging cercariae were allowed to encyst on grass blades. Soon after encystation the metacercariae were detached with the help of a small brush by applying gentle pressure and finally collected in an Eppendorf and stored at -20 °C until use.

**Biochemical estimations**

The alkali soluble glycogen was extracted and estimated by the method of Roe and Dailey [27] using rabbit liver glycogen (Sigma USA) as standard. Protein content was determined following the method of Spector [28]. RNA and DNA were estimated according to the method of Dische [29] and Giles and Myers [30] respectively. Total lipids were extracted by the method of Folch, et al. [31] as modified by Misra [32] and estimated by the method of Zöllner and Krisch [33]. The total lipid fractions such as free fatty acids and cholesterol were estimated by the method of Lowry and Tinsley [34], Sackett [35] respectively. The phospholipids were determined by estimating the phosphorus following the method of Rouser, et al. [36]. The values of biochemical components of each developmental stage were subjected to statistical analysis with respect to each other using t-test and the levels of significance were determined as described in Sokal and Rohlf [37].

**Electrophoresis**

SDS-PAGE was performed following the method of Laemmli [38] in a separating 7-15% gradient slab gel and 4% stacking gel using a vertical slab gel system. The soluble protein sample containing about 70 µg protein was mixed with an equal volume of Laemmli's sample buffer (0.625 M Tris-HCl, pH 6.8) containing 20% SDS and 5% (v/v) β-mercaptoethanol and the sample was boiled for 8 min at 100 °C. Standard high molecular weight marker proteins ranging from 29 kDa to 205 kDa were purchased from Pharmacia (LKB, Sweden). Electrophoresis was carried out at 6 °C by applying a constant current of 30 mA/slab gels until the marker tracking dye reached 1 cm above the gel before the end of the electrophoretic run. The electrophoresed gel was Coomassie stained and analyzed.

**Results**

Results obtained during the present study revealed differences in the concentration level of biomolecules in fresh eggs (E0), eggs containing mature miracidia (Em), cercariae, metacercariae and immature stages of G. crumenifer. The values obtained for various biochemical components are summarized in Tables 1 and 2.

<table>
<thead>
<tr>
<th>Biochemical components</th>
<th>Fresh eggs (0 day)</th>
<th>Eggs containing mature miracidia</th>
<th>Cercariae</th>
<th>Metacercariae</th>
<th>Immature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein</td>
<td>±0.138</td>
<td>±0.139</td>
<td>±0.119</td>
<td>±0.14</td>
<td>±0.105</td>
</tr>
<tr>
<td>Glycogen</td>
<td>±0.118</td>
<td>±0.118</td>
<td>±0.118</td>
<td>±0.118</td>
<td>±0.118</td>
</tr>
<tr>
<td>Lipid</td>
<td>±0.118</td>
<td>±0.118</td>
<td>±0.118</td>
<td>±0.118</td>
<td>±0.118</td>
</tr>
<tr>
<td>Nucleic Acids</td>
<td>±0.118</td>
<td>±0.118</td>
<td>±0.118</td>
<td>±0.118</td>
<td>±0.118</td>
</tr>
</tbody>
</table>

All the values are mean of three separate replicates, expressed in mg/g wet weight.

± SEM: indicates standard error of mean.

### Table 1: Shows variation in the level of major biochemical components during the developmental stages of Gastrothylax crumenifer.

### Table 2: Shows variation in the level of major lipid fractions during the developmental stages of Gastrothylax crumenifer.

Higher protein concentration was recorded in the fresh eggs followed by Em, cercariae, metacercariae and immature worms (Table 1). A similar trend was observed for the glycogen in the developmental stages. Differences in the lipid content were also observed in the developmental stages under study. Very little amount of lipid was detected in fresh eggs which further decreased in the eggs containing mature miracidia (Em) but increased sharply in the cercarial and metacercarial stages (Table 2). Comparative analysis of lipid fractions revealed that phospholipid concentration was higher in fresh eggs while cholesterol and free fatty acid fractions were predominant in cercarial stage. The concentration of phospholipids was observed in decreasing order starting from fresh eggs to subsequent developmental stages. However, the level of cholesterol increased up to cercarial stage followed by a decline in the subsequent developmental stages. A similar trend in the free fatty acid level was observed from E0 to immature developmental stages (Table 2). Further, the nucleic acid contents were found to be in decreasing order from E0 onwards to the later developmental stages (Table 2). An overall comparison of RNA and DNA concentration in the developmental stages of *G. crumenifer* revealed higher concentration of RNA in all the developmental stages.

Analysis of electrophoresed polypeptides revealed that the individual proteins of each developmental stage separated into a heterogeneous mixture of polypeptides of varying molecular weights (Table 3, Figure 1). A total of 27, 15, 14, 15 and 34 distinct protein bands were resolved in fresh eggs (E0), eggs containing mature miracidia (Em), cercariae, metacercariae, immature worm and adult *G. crumenifer* respectively. Moreover, some stage specific polypeptides were also observed in the developmental stages under study. A total of 8 and 9 specific polypeptides were observed in E0 and adult worm with an apparent molecular weight of 17, 36, 40, 45, 53, 57, 70 and 80 kDa for fresh eggs and 35, 38, 46, 49, 58, 73, 85, 87 and 90 kDa for adult fluke. Em, cercariae, metacercariae and immature worms showed only one specific polypeptide with a molecular weight of 18.5, 17.5, 16 and 34 kDa respectively. Further, Em and metacercarial stages did not reveal any specific polypeptide. Besides some fundamental similarities during the development of different larval stages under investigation, the protein profile revealed discrete qualitative and quantitative differences. On the basis of present data, the polypeptides can be grouped into three categories, the first category includes ‘conserved’ polypeptides comprising of those polypeptides which are present throughout the development of parasite, the second category consisted of ‘stage specific’ polypeptides which are present only in a particular stage of development while the third category includes ‘variable’ polypeptides which showed inconsistent presence during the course of development. During the present study, polypeptides with apparent molecular weight of 18-20, 25-29, and 55-57 kDa were considered as ‘conserved’ polypeptides, since these proteins were present in all the stages of *G. crumenifer* during the course of development. The Em and adult stages did not contain any characteristic polypeptide. The polypeptides with an apparent molecular weight of 30, 40, 43 kDa were considered as ‘variable’ polypeptides because of their inconsistent appearance in all the developmental stages.

**Discussion**

The life cycle of digenetic trematodes consist of different larval stages, which have strikingly different morphology, live in extremely different environments and express widely different functions. Such diversity can obviously be expected to have a reflection in the biochemical composition of the various forms [39].

The results of present study reveal that levels of major biochemical components like protein, glycogen, total lipid, lipid fractions and nucleic acids are in a dynamic state during the development of larval stages of rumen paramphistome, *Gastrothylax crumenifer*. The protein, glycogen and lipid contents of *G. crumenifer* eggs seem to be utilized as a source of nutrient during the course of development. Very little amount of lipids is found in the fresh eggs (E0) and it further decreases in the eggs containing mature miracidia (Em) but increases sharply in the cercarial and metacercarial stages indicating the substantial build up of lipid reserves. Further, the nucleic acid contents were on decrease from eggs to the subsequent developmental stages. The biochemical concentration of protein, glycogen and lipid contents was found to be higher in the free living mature cercaria as compared to the non-feeding dormant metacercarial stage, during the present study. The higher level of these biochemical components in the cercariae of *G. crumenifer* may be an adaptation because the subsequent metacercarial stage remains in a dormant state until it infects the definitive host. It is well known that the intramolluscan

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**Table 3**: Shows SDS-PAGE profile of different developmental stage of *Gastrothylax crumenifer*.

<table>
<thead>
<tr>
<th>Developmental stages</th>
<th>Total no. of polypeptides</th>
<th>Total no. of specific polypeptides</th>
<th>Apparent molecular weight of specific polypeptides (kDa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresh eggs (0 day eggs)</td>
<td>27</td>
<td>8</td>
<td>17.36,40,45,53,57,70 and 80</td>
</tr>
<tr>
<td>Eggs containing mature miracidia</td>
<td>15</td>
<td>1</td>
<td>18.5</td>
</tr>
<tr>
<td>Cercariae</td>
<td>14</td>
<td>1</td>
<td>17.5</td>
</tr>
<tr>
<td>Metacercariae</td>
<td>15</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>Immature</td>
<td>14</td>
<td>1</td>
<td>34</td>
</tr>
<tr>
<td>Mature</td>
<td>34</td>
<td>9</td>
<td>35,38,46,49,56,73,85,87 and 90</td>
</tr>
</tbody>
</table>
larval stages show preference towards the hepatopancreas which is a nutritionally rich organ of the snail and during multiplication process the increased feeding by the larvae may help in food storage for the later developmental stages. Protein synthesis undoubtedly proceeds at a higher rate in the larval parasites because the larvae not only build their own tissues but also those of the individuals of subsequent generation [40]. Studies concerned with the protein synthesis have also been carried out in other developmental stages like rediae, sporocysts or cercariae of a number of trematode species, e.g. Gyphophilmins spp., Gorgoderamphistomum and Echinoparyphium sp. [41]. The gradual decline in the protein content of the eggs during the miracidial development in the present study suggests that proteins may be involved in cellular differentiation, organogenesis and other metabolic processes as also reported in a similar study by Wilson [42] who observed a significant decline in the protein and carbohydrate reserves of the eggs during the development of F. hepatica miracidia. Marginal decrease in the energy reserves during the later phase of miracidial development of G. crumenifer reflects a slow rate of metabolism which may be due to glycogenolysis [43]. Rogers and Petronijevic [44] suggested that the development of an organism is controlled by sets of genes which are activated or suppressed on receiving an appropriate stimulus involving complex interaction between intrinsic and extrinsic factors at a specific stage of the life cycle. Such controlled genes may also be responsible for contributing the differential antigen/protein turnover in parasites. The proteins are of obvious interest because they are the first conceivable product of gene activity involving development of new structures and physiological activities which are linked with these molecules besides immunologically linked activities.

The presence of a large amount of glycogen in the freshly collected eggs of G. crumenifer and a gradual decline in their levels during the later developmental stages clearly indicates that these components are being actively utilized for somatic differentiations of the miracidia and other developmental stages. Horstmann [45] and Wilson [42] have also reported a decline in the glycogen level and oxygen consumption during the miracidial development of F. hepatica. The glycogen content of transforming G. crumenifer cercariae into immature stage revealed nearly 45% decline under present investigation. In earlier studies it has been reported [46] that 50% decrease in carbohydrate/protein ratio, during the transformation from cercariae to schistosomule. The loss of carbohydrate material was also reflected in the intensity of PAS-stained electrophoretic bands of schistosome larval extracts. Gazzinelli, et al. [47] reported the loss of total carbohydrate and protein from the transforming cercariae. They (loc. cit) determined that two polysaccharide bands obtained from cercarial extracts were much reduced in schistosome preparations. Studies on the carbohydrate metabolism of the intramolluscan stages have been generally restricted to a few species e.g. Schistosoma spp. [48-50], Cercaria helvetica [51,52], Cercaria linearis and F. hepatica [53,54]. Histochemical studies by Cheever and Weller [55] and Cheng and Snyder [56] have revealed that the cercariae and rediae were rich in glycogen. In schistosome cercariae the tail acts a major reserve in glycogen. In schistosome cercariae the tail acts a major reserve and Snyder [56] have revealed that the cercariae and rediae were rich in glycogen. In schistosome cercariae the tail acts a major reserve of nutrients comprising the glycogen and cytochrome oxidase [40]. Little is known about the carbohydrate metabolism of metacercariae which is considered to be a dormant stage. Studies on metacercariae are more or less fragmentary and primarily of descriptive nature [57,58]. However, a number of aspects on carbohydrate metabolism such as glucose uptake, glucose leakage and the effect of starvation on glycogen content of the non-dormant progenetic metacercariae Clonostomum complanatum have been observed by Siddiqui and Nizami [59]. These authors (loc. cit.) reported high levels of glycogen in the C. complanatum metacercariae which may be because this parasite inhabits a semi-anaerobic environment and being a transitional stage in the development, requires more energy for growth and metabolism. Occurrence and utilization of glycogen in Cyclocoelum aculeum [60], Schistosoma mansoni and S. japonicum [61] has also been reported. Variations in the biochemical contents have also been observed during the miracidial development of an amphistome Gigantocotyle explanatum[43].

The level of RNA concentration constitutes a marker for over all metabolic activity [62] in an organism. The elevation of RNA concentration in the larval stages of G. crumenifer during the present study suggests the possibility of activation of development regulatory genes as well as somatic growth and differentiations.

During the present study, changes in lipid level of developing miracidia were significant (p<0.05). Further, significantly high (p<0.001) level of lipid was recorded in cercariae and metacercariae. It can be suggested that the lipids in developing miracidia are possibly stored for utilization in the subsequent stages of the life cycle [42]. In the present study, increased level of lipids in cercarial stage as compared to the metacercariae indicates rapid synthesis. The possible reason for the higher concentration of lipid could be because the cercariae occupy a nutritionally rich habitat, the hepatopancreas which might be a contributing factor in a nutritionally rich habitat and building the nutritional reserve for the dormant metacercarial stage. Higher lipid content was also reported in the fish infecting C. complanatum metacercariae which could be an adaptation for long survival strategies [59]. This may also be true for the higher concentration of lipid contents in G. crumenifer metacercariae under study, because the stored lipids not only help in the survival of the metacercariae but they are also utilized by the adult worms during the egg production [57,63,64]. The difference in lipid and their fractions in the developmental stages under study can be correlated with the metabolic and physiological differences of their microhabitat which are influenced by the effects of various environmental factors. Further, Vykrestyuk and Yarygina [65] have suggested that the inconsistence in the lipid composition of parasites may depend on factors like parasite species, sex, age and environmental factors like habitat and diet of the host. Among the major lipid fractions in G. crumenifer developmental stages the phospholipids concentration was observed higher followed by cholesterol and free fatty acids. The major lipid fractions were triglycerides followed by cholesterol, free fatty acids and phospholipids. Fairbairn [66] and Barrett [67] have reported the utilization of esterified fatty acids and triacyl glycerol for energy release during the embryonation of Ascaris eggs. Higher level of free fatty acids in the eggs containing mature miracidia (Em) stage of G. crumenifer can be suggested as a biochemical adaptation for the subsequent free living developmental phase. This is significant because, fatty acids provide more metabolic water upon oxidation than other metabolic fuels [68]. High concentrations of free fatty acids have also been reported from a number of nematode larvae [69,70]. Further, Lee and Song [70] reported an increase in the cholesterol of A. suum during the development of eggs. Decline in the level of phospholipids from fresh eggs to the subsequent developmental stages of G. crumenifer, indicates their involvement in molecular reorganization and metabolism. Among various lipid

fractions of *C. complanatum* metacercariae, triglycerides account for high percentage while the cholesterol and free fatty acids were observed in approximately similar amounts [59]. A similar trend was reported for adult *Dugesia dorotocephala* [71], *S. mansoni* [72] as well as *G. crumenifer*, *Cotylphorophor cotylphorum*, *G. explanatum*, *P. buski*, *Isparorchis hypselobagri* and *Gastrodiscoides hominis* [73,74]. Thus it can be suggested that levels of total lipids and their fractions in developmental stages of *G. crumenifer* during present investigation are controlled by both external and internal environmental conditions which the parasite encounters during thermal and nutritional acclimatization.

Analysis of the protein profile of various developmental stages of *G. crumenifer* revealed quantitative as well as qualitative variations. Apart from some fundamental similarities, proteins of ‘conserved’, ‘stage specific’ and ‘variable’ nature were also identified. The conserved polypeptides like 18-20,25-29 and 55-75 kDa were present throughout the developmental stages, viz. fresh eggs (E0), eggs containing mature miracidia (Em), cercariae, metacercariae, immature (size, 2mm) and adult worms under study while the stage specific polypeptides were characteristic of a particular stage only. Similarly, Anderson [75] observed a total of 20 conserved polypeptides (MW 23-220kDa) in the soluble proteins of 4.5, 8, 11 and 18 week old F. hepatica. The surface protein of schistosomula and adult stages also revealed a number of conserved polypeptides in the molecular weight range of 16-150 kDa [76]. In the cestode parasite *Echinococcus granulosus*, the conserved group of polypeptides included a wide range of varying molecular weight (~29 to 153 kDa) polypeptides [77]. On the other hand some variable polypeptides were also observed during the present study. Unfortunately, no previous report on the protein turn over during the course of development is available from any other amphistomes; therefore the present work is just a beginning to open further avenues for research in this direction. The SDS-PAGE results resolved a minimum of 14 and a maximum of 34 protein bands in the protein sample of developmental stages under study and it was observed that three polypeptides of 24, 28 and 43 kDa in E0 and Em stages, six polypeptides of 16, 20, 21.5, 23, 25 and 28 kDa in cercariae and metacercariae while eight polypeptides of 18, 19, 20, 25, 28, 30, 40 and 50 kDa in immature and mature worms were of similar intensity. A similar pattern was recorded by Ruppel and Cioli [78] who observed some dissimilarity in the protein profiles of mature and immature schistosomes as well as male and female worms. During the present study, polypeptides of 58 and 23 kDa molecular weight were observed specific to the adult *G. crumenifer*, which was conspicuously absent in all the other developmental stages. The stage specific protein with a molecular weight of 58 and 23 kDa might be associated with appearance of some important adult structure like the intestine and its contents as has also been suggested by Ruppel and Cioli [78] for schistosomes. It seems that in spite of great morphological differences during the development, specialized structures characteristic of each stage emerge as differences in protein composition, if they represent a sizable portion of the total parasite tissues. In addition to this, we reported earlier some conserved and species specific polypeptides in the fresh eggs of five amphistomes species [79]. Moreover, variations in the soluble egg polypeptide profile of three isolates of *S. mansoni* of different geographical regions are on record [80-83]. Occurrence of ‘conserved’, ‘stage specific’ and ‘variable’ polypeptides in the developmental stages of *G. crumenifer* during the present study requires further studies to ascertain their functional role as well as biochemical nature so that the polypeptides of interest may be used in chemotherapeutic modulation to block reproduction and development of the parasite.

Taken together, it can be said that the fluctuations in the biochemical components like protein, glycogen, lipid, lipid fractions, RNA and DNA of developmental stages of the rumen amphistome *G. crumenifer*, may either be a characteristic of a particular developmental stage under study or else may be due to the influence of various physico-chemical factors of different micro-habitats. It could also be a biochemical adaptation to meet the metabolic requirements. Thus, it can be concluded that the variation in the metabolic turnover accompanying the transition from one larval stage to the next involves quantitative as well as qualitative changes in components which may be greatly influenced by various physico-chemical factors of different micro-habitats.

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**References**


