THE USE OF PROTOZOA, ROTIFIERS AND NEMATODES AS LIVE FOOD FOR SHRIMP RAISED IN BFT SYSTEM

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ABSTRACT

The Protozoan and Nematodes Culture Medium – PNCM, enable inexpensively production of microorganism to be used as live food in shrimp culture. It was tested the ciliates produced by PNCM as live food for shrimp larvae in an experiment with four treatments: (BFT) only biofloc medium; (BFT 10) 10 mL of PNCM culture added to biofloc; (BFT 100) 100 mL of PNCM culture added to biofloc; and (BFT 1000) 1000 mL of PNCM culture added to biofloc. The microorganisms were analyzed at the beginning, intermediate and final phases of the experiment which lasted for 30 days. The analysis of shrimp gut contents indicated a predation of shrimps on the ciliates. In most treatments, the initial period was marked by an increase in the abundance of the ciliates and rotifers. Nematodes were absent at the end of the final phase in all treatments, suggesting that these organisms were effectively preyed by shrimps. This study indicate that ciliates, rotifers and nematodes play an important role as live food in hatcheries mainly due to their small size, nutritional value and attractivity exerted on shrimp post-larvae.

INTRODUCTION

In the Biofloc-Technology (BFT), with zero water-exchange, macroaggregates or “bioflocs” are formed during the shrimp production cycle with the objectives of reducing wastewater discharge and environmental impact caused by the input of dissolved nutrient.

The microbial flocs are comprised primarily by bacteria, cyanobacteria, algae, protozoans, small metazoans, invertebrates larval forms, feces and dead organisms remains, which could serve as complementary food to shrimps. Some of these microorganisms (heterotrophic bacteria) are able to assimilate the nitrogen compounds from the shrimp excretion and food decomposition, improving water quality (McIntosh et al. 2000, Decamp et al. 2002, Burford et al. 2003, Wasielesky et al. 2006a, Wasielesky et al. 2006b).

The species composition of the natural microbial community, influences the biochemical composition of the floc and therefore, its nutritional value and possibly, palatability to shrimp. Protozoan like flagellates and ciliates contain sterols in their chemical composition, and a large portion of these sterols is converted to cholesterol or other lipid forms. Moreover, dry matter of eukaryotic organisms is composed of 50% protein, 10% lipids, 3 - 4% of RNA and DNA. Similarly, nematodes biochemical analysis indicated high levels of protein and fat (Biedenbach et al. 1989) but these levels are significantly dependent on the environment where the nematodes are cultured.

Thus, the addition of cultured microorganism in BFT, especially ciliates and nematodes could increase the protein and lipid availability, improving the cultured organisms growth (Phillips 1984; Stolp 1988, Biedenbach et al. 1989, Decamp et al. 2002, Focken...
et al. 2006) and providing a reduction of crude protein used in commercial feed (Samocha et al. 2004).

A technique of massive production of protozoan and nematodes, based on the Martinez-Cordova (2002) method, enabled an inexpensive enrichment of super-intensive shrimp culture with ciliates and nematodes. The method uses alfalfa hay and sugarcane molasses as sources for the growth of microorganisms (Loureiro et al. submitted). The bacteria will act in the decomposition of organic matter using carbon as an energy source, and nitrogen to form the cell structure through protein synthesis. The bacterial growth will allow the establishment of a microbial food web composed by flagellates, ciliates, rotifers and nematodes.

This study aims to determine the use of ciliates, produced by the PNCM, as live food to enrich the BFT system.

**MATERIAL AND METHODS**

The study was carried out in the facilities of the Aquaculture Marine Station - EMA of the Federal University of Rio Grande - FURG, at Cassino beach - Rio Grande – RS (Brazil). The experiments started on February 1st, 2010, and lasted for 30 days.

*Protozoan and Nematodes Culture Medium – PNCM*

In order to stimulate the growth of heterotrophic microorganisms, an intensive bacteria, protozoan and nematodes culture was developed for 10 days before the beginning of the trial. Six cylinder-conical tanks were filled with 50 L of filtered seawater (5 μm Cuno filter). In each tank, 20 g L\(^{-1}\) of sugarcane molasses were added during three consecutive days. Besides, cod liver oil was used at a rate of 18 mL L\(^{-1}\) as well as 2 g L\(^{-1}\) of yeast (*Saccharomyces cerevisiae*), 0.2 g L\(^{-1}\) of ascorbic acid and 100 g L\(^{-1}\) of alfalfa hay (Martinez-Cordova 2002 modified). The inoculum of microbiota was obtained from 50 mL of brackish water (20 psu) from Patos Lagoon estuary. All ingredients were mixed up and remained under continuous aeration and temperature around 38 °C.

A 10-mL aliquot was sampled, fixed and stained with Lugol’s iodine solution of 2% (v:v) for the determination microorganism abundance into the PNCM tanks. Further the identification and quantification of microorganisms, such as flagellates, ciliates, rotifers and nematodes, were done according to Utermöhl method (Throdsen 1978), using an inverted microscope Axiovert 135 ZEISS at 200X magnification.

**Experimental design**

Sixteen 5-L plastic containers with 4 liters of final volume were divided into four treatments as follow: a) with only biofloc (BFT treatment); b) 10 mL of PNCM culture added to biofloc medium (BFT 10 treatment); c) 100 mL of PNCM culture added to biofloc medium (BFT 100 treatment) and d) 1000 mL of PNCM culture added to biofloc medium (BFT 1000 treatment).

The biofloc used in the experiment was originated from a tank production of juvenile *Litopenaeus vannamei* at Marine Aquaculture Station - EMA. During the study the containers remained in a water-table with controlled temperature (28 °C) and constant aeration. Every container received 50 L. *vannamei* post-larvae (PL25) (0.006 ± 0.003 g) per liter.

The PLs used in this study were obtained at Aquaculture Marine Station hatchery and their survival rate was estimated by counting the PLs at the beginning and the end of study period.

**Maintenance of the experimental units**

Water in the tanks was renewed every 48 hours (renewal cycle) by siphoning of almost 90% of total volume. The original volume was replaced by the addition of water with bioflocs from production tanks located at EMA/FURG. The same amount of pre-established PNCM was added in every container.

**Analysis of microorganisms**

The microorganisms were analyzed at the beginning, intermediate and final phases of the experiment. Water samples were collected twice in each phase. For this, after each renewal cycle, 30-mL water samples were taken from containers for quantification of flagellates, ciliates, rotifers and nematodes (initial period sample). Same procedure was repeated 48 hours after water renewal. Differences of microorganism abundance in both sampling periods indicated possible microorganisms consumption by shrimps. Thus initial sample occurred between February 7th (initial period sample) and at February 9th, 2010 (final period sample), in the intermediate phase.
Sampling was conducted at February 17th (initial period sample) and at February 19th, 2010 (final period sample) and the final phase, sampling was conducted at February 27th (initial period sample) and at March 1st, 2010 (final period sample).

All samples were preserved in 2% Lugol’s iodine solution (v:v) and stored in amber glass flasks (50 mL) for posterior analysis of the microorganism community. Aliquots of 2.1 mL from every sample were analyzed in sedimentation chambers (Throndson 1978). The quantification was performed using an inverted microscope Axiovert 135 (ZEISS) at 100X, 200X and 400X magnifications (Utermöhl 1958).

**Physicochemical parameters**

An YSI™ mod. 556 sensor was used for daily measurements of water temperature, salinity, pH and dissolved oxygen. The suspended solids and total ammonia (TAN) concentration were determined at the end of each phase (initial, intermediate and final). For this, Imhoff’s cone was used to determine the volume of suspended matter accumulated on the bottom of the cone 15 min after sampling (Avnimelech 2007). The TAN was determined following the method proposed by UNESCO (1983).

**Feed supply**

In all treatments the initial feed supply (Guabi - Active 38% PB) was equivalent to 50% of the shrimp biomass (0.6 g day⁻¹). From the fourth day of the experiment period the amount of feed had to be daily adjusted, regarding the surplus of the last time of feeding.

**Shrimp gut content analysis**

The analysis was performed on 5 shrimps collected from all treatments on the initial, intermediate and final phases of culture. The dissection was conducted under a stereo microscope using dissection needles. The stomach was removed and its content extruded on a slide, the material was stained with Lugol’s iodine solution and identified using an inverted microscope Axiovert 135 (ZEISS) at 100X, 200X and 400X magnifications.

**Statistical analysis**

A one-way analysis of variance (ANOVA) was applied to differentiate the survival rate, final weight of shrimps, the dissolved oxygen, the ammonia-TAN concentration and the initial and final abundance of microorganisms. Significant differences at (p<0.05) were determined using the Tukey’s test (Zar 1999).

**RESULTS**

**Zootechnical Indexes**

Survival rates and final weights of the shrimps are presented in Table 1. The shrimps had better survival rates in the BFT 100 and BFT 1000 than in the control (BFT). The final weight of PL in the BFT 10 indicated a higher weight increment than observed in the other treatments.

<table>
<thead>
<tr>
<th>BFT</th>
<th>BFT 10</th>
<th>BFT 100</th>
<th>BFT 1000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Survival (%)</td>
<td>44±2.6 a</td>
<td>42±3.2 a</td>
<td>53.5±4.2 b</td>
</tr>
<tr>
<td>Final weight (g)</td>
<td>0.068±0.003 a</td>
<td>0.085±0.003 b</td>
<td>0.059±0.002 c</td>
</tr>
</tbody>
</table>

**Water Quality Parameters**

As showed in table 2, there were much higher DO concentrations in the BFT and BFT 10, which were significantly different from the other treatments, as well as there also were higher ammonia - TAN values in those treatments. Apart from this, there were no significant differences among the treatments for temperature, salinity, pH and suspended matter during all the experiment period (Table 2).
TABLE 2 – Water quality parameters (MD±SD) of *Litopenaeus vannamei* juvenile tanks, exposed to: BFT, BFT 10, BFT 100 and BFT 1000 treatments during the culture. Different letters indicate significant difference between treatments (*p* <0.05).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>BFT</th>
<th>BFT 10</th>
<th>BFT 100</th>
<th>BFT 1000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dissolved Oxygen (mg L⁻¹)</td>
<td>7,29±0,32 a</td>
<td>7,24±0,41 a</td>
<td>6,35±0,36 b</td>
<td>4,60±1,35 c</td>
</tr>
<tr>
<td>Temperature (°C)</td>
<td>28,02±0,37</td>
<td>28,12±0,26</td>
<td>28,02±0,37</td>
<td>28,12±0,27</td>
</tr>
<tr>
<td>Total ammonia-TAN (mg L⁻¹)</td>
<td>0,79±0,19 a</td>
<td>0,30±0,14 b</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>Salinity (%)</td>
<td>32,6±0,5</td>
<td>32,6±0,5</td>
<td>32,6±0,5</td>
<td>32,6±0,5</td>
</tr>
<tr>
<td>pH</td>
<td>7,9±0,18</td>
<td>7,9±0,22</td>
<td>8,1±0,17</td>
<td>8,2±0,14</td>
</tr>
<tr>
<td>Cone Imhoff (ml L⁻¹)</td>
<td>8,7±4,2</td>
<td>9,9±3,8</td>
<td>9,6±5,8</td>
<td>7,7±5,5</td>
</tr>
</tbody>
</table>

Analysis of microorganisms

The abundances of flagellates, ciliates, rotifers and nematodes in the initial phase (February 7th) are shown in the Figure 01. In this period of the experiment there was a significant increase of flagellates in most treatments, except in the BFT 100 (Fig. 01B). Overall, there was a decrease in the concentration of larger organisms such as rotifers and nematodes, mainly in the BFT 100 and BFT 1000 treatments, respectively (Fig. 01F and 01H).

![Figure 1](image-url)
A significant increase in the flagellates abundance 48 hrs after the addition of PNCM was also observed in the intermediate phase of the experiment (February 17th) in BFT 10 and BFT 1000 treatments (Fig. 02B), as well as a significant decrease in the population of ciliates in BFT 100 and BFT 1000 treatments (Fig. 02D). Likewise observed for the initial phase, there were a few rotifers left (Fig. 02F) and the nematodes were absent at the end of that intermediate phase (note the lack of the picture for this).

By contrast, the final period (February 27th) was represented by the increase in the abundance of the ciliates in BFT 10 and BFT 1000, the rotifers abundance increased in most treatments, except in the BFT treatment (Fig. 03F). Despite the presence in the BFT 1000, nematodes were absent at the end of the period of that final phase. There was an increase in the number of flagellates at the end of this final period only in the BFT 10 (Fig. 03B).
**Shrimp gut content analysis**

Ciliates and rotifers were seen in the shrimp gut contents mainly in the initial and intermediate phases of culture, indicating that the shrimp grazed preferentially on these microorganisms especially in the early stages of life. Flagellates and nematodes fragments were not seen at any stage.

**DISCUSSION**

The Zooplankton Promoter – ZP was developed by Martinez-Cordova (2002) as a strategy to increase the abundance of zooplanktonic organisms in extensive shrimp culture. The ZP consisted of bunches of alfalfa with sugarcane molasses, vitamins and fish oils, these bunches were bundled and put into plastic net bags, subsequently placed in the ponds at a rate of 4-kg (alfalfa) per hectare. It increased the microorganism growth that benefited the raised shrimp (Martinez-Cordova 2002; Martinez-Cordova *et al.* 2002). Some improvements of this technique led to the development of the PNCM, which promote the development of a microbial food chain, originally represented by flagellates species, ciliates like *Uronema sp.*, *Litonotus* sp. followed by *Euplotes* sp. and finally by nematodes (*Rhabditis* sp.) (Loureiro *et al.* submitted to Aquaculture).

These microorganisms are important source of lipids, proteins and vitamins to aquatic cultured organisms (Abreu *et al.* 1988; Biedenbach *et al.* 1989;...
is likely that the huge numbers of the microorganisms in the BFT 100 and BFT 1000 treatments may have contributed to the lower DO levels, due to supposedly increased microbial respiration rates (Moriarty 1987). Some studies indicate that the dissolved oxygen is the most limiting factor for the growth of penaeid shrimp, and show that decreasing DO levels in the tanks can inhibit their growth and consequently reduce ecdysis (Clark 1986). The oxygen consumption is an important part of the bioenergetic balance of shrimp, because it reflects the energy that is directed to metabolic work. Although it does not act directly on the growth or feed digestibility, it may limit the food consumption and shrimp growth rates (Lucas 1993; Rosas et al. 1998). In contrast, culture environments with low dissolved oxygen levels have been resulted in lower weight of shrimp at the harvest (Rosas et al. 1998; Vinatea 1997; Ribeiro 2001b, Li et al. 2006). However, we believe that the lower final body weight of shrimp in the BFT 100 and BFT 1000 treatments was due to the lack of commercial feed in these treatments where survival rate was higher.

The results of this study indicate that protozoan, nematodes and rotifers produced massively in the Protozoan and Nematodes Culture Medium can be an effective source of live food for cultures shrimp. However, further studies must be conducted in order to improve the survival of shrimp reared at high densities using the PNCM.

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