Zooplankton body-size structure and biomass in tropical floodplain lakes: relationship with planktivorous fishes

Claudia Costa Bonecker1, Fábio de Azevedo2 and Nadson Ressyé Simões1

1Núcleo de Pesquisas em Limnologia e Aquicultura – Nupélia, Universidade Estadual de Maringá – UEM, Av. Colombo, 5790 – Bloco H-90, Maringá, PR, Brazil
e-mail: bonecker@nupelia.uem.br; nadsonressye@yahoo.com.br
2Faculdade Estadual de Educação, Ciências e Letras de Paranavaí – FAFIPA, Brazil
e-mail: fabioazeve@yahoo.com.br

Abstract: Aim: This study evaluated whether the size structure of the zooplankton community, as analyzed from density and biomass, would be influenced by bottom-up and/or top-down mechanisms in isolated lakes from the Upper Paraná River floodplain during dry and rainy periods. Methods: zooplankton individuals were classified as: smaller-sized (<300 µm), intermediate-sized (301 to 600 µm) and larger-sized (greater than 601 µm). Fish abundance was indexed by the capture per unit effort (CPUE; number of individuals.100 m–2). Productivity was measured as chlorophyll-a. Results: The size structure of the community did not show significant relationship with chlorophyll-a concentration (p > 0.05), but it was associated to the increase in fish density during the dry season. The percentage of individuals with intermediate sizes (301 to 600 µm), in both biomass and density, was positively related to the fish density (R2 = 0.78 for biomass, p < 0.001; R2 = 0.58 for density, p = 0.02), indicating an increase numerical and mass of this size class with fish density. The percentage of larger-sized individuals (>600 µm), in both biomass (R2 = 0.86, p < 0.001) and density (R2 = 0.69, p = 0.02) was negatively associated to the fish density, reflecting a decrease of this size class with the increase of fish in the environment. Conclusions: This can implicate a direct and negative effect of the predation on larger-sized individuals, as well as indirect and positive effect on the intermediate-sized individuals, but it occurred only on dry season. Thus, different mechanisms act on the size structure of the zooplankton community between dry and rainy season. Keywords: predation, seasonal variation, fishes, shallow lakes, zooplankton body length.

Resumo: Objetivo: Este estudo avaliou se a estrutura de tamanho da comunidade zooplânctônica, analisada a partir da densidade e da biomassa, seria influenciada por processos ascendentes e/ou descendentes em lagos da planície de inundação do Alto Rio Paraná durante um período seco e outro chuvoso. Métodos: o zooplâncton foi classificado em: pequeno porte (<300 µm), porte intermediário (301 a 600 µm) e maior porte (maior que 601 µm). Abundância de peixes foi estimada em captura por unidade de esforço (CPUE; número de indivíduos.100 m–2). Produtividade foi definida como a concentração de clorofila-a. Resultados: A estrutura de tamanho da comunidade não apresentou relações significativas com a concentração de clorofila-a (p > 0,05), mas ela foi associada com o aumento da densidade de peixes no período seco. O percentual de indivíduos com tamanhos intermediários (301-600 mm), tanto em biomassa quanto em densidade, foi positivamente relacionado com a densidade de peixes (R2 = 0.78 para biomassa, p < 0,001; R2 = 0.58 para densidade, p = 0,02), indicando um aumento numérico e em massa dessa classe de tamanho com o aumento da densidade de peixes. A percentagem de indivíduos de maior porte (>600 mm), tanto em biomassa (R2 = 0.86, p < 0,001) quanto em densidade (R2 = 0.69, p = 0,02), esteve negativamente associado com a densidade de peixes, refletindo uma diminuição desta classe de tamanho com o aumento da densidade de peixes no ambiente. Conclusões: Isto pode implicar um efeito direto e negativo da predação sobre indivíduos de maior porte, bem como efeito indireto e positivo sobre os indivíduos de pequeno porte. No entanto, esta relação foi observada apenas no período seco. Assim, diferentes mecanismos atuam sobre a estrutura de tamanho da comunidade de zooplâncton entre os períodos analisados. Palavras-chave: predação, variação sazonal, peixes, lagos rasos, comprimento do corpo do zooplâncton.
1. Introduction

Brooks and Dodson (1965) showed that interactions of competition and predation influenced the structure of the zooplankton community in different lakes in the USA. The authors formulated the size-efficiency hypothesis, which indicates that filter-feeding zooplankton compete for a similar size of food (1-15 μm) and that the larger zooplankton filter particles more efficiently than the small zooplankton. But, when the predation pressure by fish is intense, larger zooplankton will be removed of the community and small zooplankton will dominate. On contrary, larger zooplankton will dominate due to their greater ability of filter-feeding overcoming competitively small ones. Dodson (1974) tested the size-efficiency hypothesis, showing that the larger zooplankton does not always exclude the smaller ones through competition for food. The author suggested an alternative hypothesis to extend the understanding of the importance of size-selective predators in order to include invertebrates selecting small prey. If the vertebrate predators are present, small zooplankton and invertebrate predators coexist. In the opposite, the latter predators coexist with large zooplankton.

This context suggests that large zooplankton should be a better competitor when resources become limiting due to their ability to use large and small particles and to survive at lower food levels, even if the partitioning of food resources among different sizes of zooplankton reduces competition between them: small zooplankton (rotifers and small cladocerans) ate fine particles (1-5 μm), whereas larger zooplankton (larger cladocerans and copepods) consumed larger particles (<5-15 μm) (Gliwicz, 1969).

However, Bogdan and Gilbert (1984) stated that there is no general relationship between zooplankton body length and the ability to ingest very small bacterial and algal cells. The efficiency rate of Crustacea, for example, on ultraplankton (planktonic organisms lesser than 2 micrometers in size) appears to be also correlated with the structural characteristics of the feeding appendages rather than with body size (Elmoor-Loureiro, 2005; Hart and Bychek, 2011). Body length certainly influences the maximal particle size a species can ingest, but it has little influence on the ingestion of smaller particles (Brooks and Dodson, 1965).

Body size is one of the most important parameters that determine the ecological and physiological characteristics of an organism (Peters, 1983). It also can reflect the competitive abilities of all organisms within a population and thus reflect the question of niche selection, predation relationship and the coexistence of similar species (Lampert and Sommer, 1997; Chase et al., 2002). Current studies point out that the predation on zooplankton species, especially by fish, has an important effect on the temporal variation of the community body size, resulting in an increase of small zooplankton dominance (Horn, 2003; Hambright, 2008). Nevertheless, this change in the size spectrum of the community also depends on the temporal dynamics from the involved populations, such as life cycle (Winder et al., 2003); and, on the smaller size of zooplankton species from tropical regions (Fernando, 1994); or, thermal influences (Hart and Bychek, 2011).

In this way, if the predation pressure is constant in tropical environments, the assessments concerning the size-selective predators hypothesis on the zooplankton community, in natural conditions, would be limited because the small-sized species numerically predominate in the community over time, since the zooplankton’s predators be present. Thus, we tested the size-selective predators hypothesis, and if size spectrum of the zooplankton community is associated with availability food. Therefore, the present study evaluated whether the organization of size structure in the zooplankton community would be influenced by bottom-up and/or top-down mechanisms in isolated lakes from the Upper Paraná River floodplain, during both a rainy and a dry periods. This implies that the size-efficiency hypothesis may also explain the community structure, and may further contribute to the differentiation of the size spectrum from the community between tropical and temperate lakes.

2. Material and Methods

2.1. Study area

The shallow lakes sampled (Capivara-A, Aurélio-B, Clara-C, Jacaré-D, Genipapo-E, Osmar-F, Traíra-G and Pousada-H) are located in the Upper Paraná River floodplain in the Paraná and Mato do Grosso do Sul States, Brazil (Figure 1). They have mean depth ranging 0.96 to 3.60 m and surface area ranging 0.006 to 7.20 ha (Table 1). The samples were carried out during the dry (August 2000) and rainy (February 2001) periods in order to verify if the results depend on the period. Maximum hydrometric level achieved 3.46 and 4.14 m, dry and rainy, respectively.
Table 1. Location and general characterization of the lakes’ size in the Upper Paraná floodplain.

<table>
<thead>
<tr>
<th>Location</th>
<th>Mean depth (m)</th>
<th>Area (ha)</th>
<th>Geographic coordinates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aurélio - B</td>
<td>1.95</td>
<td>0.43</td>
<td>22° 41' 34.68&quot; S; 53° 13' 50.58&quot; W</td>
</tr>
<tr>
<td>Capivara - A</td>
<td>3.60</td>
<td>7.20</td>
<td>22° 47' 56.52&quot; S; 53° 32' 5.4&quot; W</td>
</tr>
<tr>
<td>Clara - C</td>
<td>1.20</td>
<td>0.90</td>
<td>22° 45' 17.52&quot; S; 53° 15' 28.62&quot; W</td>
</tr>
<tr>
<td>Genipapo - E</td>
<td>0.96</td>
<td>0.06</td>
<td>22° 45' 33.24&quot; S; 53° 16' 5.94&quot; W</td>
</tr>
<tr>
<td>Jacaré - D</td>
<td>2.14</td>
<td>6.96</td>
<td>22° 47' 2.04&quot; S; 53° 29' 49.08&quot; W</td>
</tr>
<tr>
<td>Osmar - F</td>
<td>1.10</td>
<td>0.006</td>
<td>22° 46' 26.64&quot; S; 53° 19' 56.16&quot;</td>
</tr>
<tr>
<td>Pousada - H</td>
<td>2.30</td>
<td>3.80</td>
<td>22° 42' 1.14&quot; S; 53° 15' 23.52&quot; W</td>
</tr>
<tr>
<td>Traíra - G</td>
<td>2.10</td>
<td>0.47</td>
<td>22° 44' 45.6&quot; S; 53° 20' 21.66&quot; W</td>
</tr>
</tbody>
</table>

2.2. Field sampling

Water samples were collected with a van Dorn bottle from the subsurface in the pelagic region of each lake in order to analyze limnological features of water. Water temperature, concentration of dissolved oxygen (YSI oximeter), pH, electric conductivity (DIGIMED potentiometer), total alkalinity (Carmouze, 1994), water transparency (turbidimeter) was measured in situ. Sub-samplings of water were transported to laboratory for analyses of chlorophyll-a and nutrients.

Zooplankton samples were undertaken in the limnetic region from each lake at the subsurface (between 0.5 and 1.5 m) in the morning, in order to minimize possibility of diurnal vertical migration (even the lakes were shallow), using a motorized pump and a plankton net (68 µm) filtering 600 L of water per sample, and were
preserved in formaldehyde at 4% buffered with calcium carbonate. Samples were always obtained using a boat moving at constant speed to prevent a biased sampling.

Fish were captured on September 2000 and March 2001 using 20 m seining nets (internal mesh of 0.5 cm opposite knots) operated during the day in littoral zones. The species abundances in every sample were indexed by the capture per unit effort (CPUE; number of individuals.100 m$^{-2}$).

### 3. Laboratory Analyses

Water samples for chlorophyll-$a$ concentration were stored in an ice chest, and then they were filtered through GF 52-C 130 membranes (<10 hours after sampling) and immediately frozen (−20 °C) for further analyses according Golterman et al. (1978). The ions nitrate, nitrite, ammonium and reactive dissolved phosphate were measured in a spectrophotometer, according to Bergamin et al. (1978), Mackereth et al. (1978) and Golterman et al. (1978). Some studies have used chlorophyll-$a$ concentration as surrogate to primary productivity (Declerck et al., 2007; Thackeray, 2007).

Zooplankton abundance was estimated from counting 2 mL sub-samples using Sedgewick-Rafter chambers and Hensen-Stempell pipette. At least three sub-samples were counted, containing at least 80 individuals within each sub-sample (Bottrell et al., 1976). These sub-samples were then summed, and the organism's density for the total sample volume was then estimated and expressed in individuals.m$^{-3}$.

Biovolume calculation for rotifers was determined according to Ruttner-Kolisko (1977), and the biovolume values were converted to wet weights. Assuming that 10$^6$ µm$^3$ corresponds to 1 µg of wet weight (Bottrell et al., 1976), the dry weight was estimated as 10% of wet weight (Pace and Orcutt Junior, 1981). The cladoceran and copepods biomass were estimated through calculated length-weight relationships (from weighing in a micro-analytical balance of 10$^{-7}$ g) (Mc Cauley, 1984; Wetzel and Likens, 2000).

The organisms' length was obtained under optical microscope, using an objective with micrometer reticle, considering (i) for the rotifers, the distance between the superior and inferior margin of carapace, without spines; (ii) for the cladocerans, the distance from the head until the end of the carapace, without the helmet and spine; and (iii) for the copepods, the distance from the head until the genital segment. These measures were taken from at least 30 individuals of each species. The individuals were classified as: smaller-sized, for those belonging to size class lesser than 300 µm; intermediate-sized, for those belonging to size class ranging from 301 to 600 µm; and larger-sized, for those individuals belonging to size class greater than 601 µm.

According to Bottrell et al. (1976) and Dumont et al. (1975), the weight differences between organisms preserved in formaldehyde, as used in our study, and non-preserved organisms are negligible or showed a little influence on the biomass results. Thus, mathematical corrections for the obtained weight results were not performed.

All individuals of fishes were identified, enumerated, weighed (g) and classified according to Britski et al. (1999). Voucher specimens were deposited in the Ichthyological Collection of Nucleus of Research in Limnology, Ichthyology and Aquaculture (Nupelia). In order to indicate fishes species potentially consumers of zooplankton, we consulted bibliography for tropical fishes (Casatti et al., 2003; Cassemiro et al., 2002; Crippa et al., 2009; Russo and Hahn 2006; Pelicice and Agostinho, 2006; Santana-Porto and Andrian, 2009) and a list available in http://www.fishbase.org/.

#### 3.1. Data analyses

Mann-Whitney tests were realized in order to verify differences in limnological features of the water between seasons.

Principal Component Analysis (PCA) was used to summarize the relationship between samples locales and limnological features. Data were standardized through a correlation matrix.

Linear regression analyses were carried out, separately, to verify as fish density and chlorophyll-$a$ concentration are associated with aggregate community properties (density and biomass) and with percentage of size classes on the community structure. Values of fish density, chlorophyll-$a$ concentration, zooplankton biomass and density were log-transformed to minimize data variability and assist the visualization of the results. The regression assumptions were tested through residual visualization.

The significant level adopted in the analysis was 0.05, meaning that $p$-values lower than 0.05, the relationships recorded are rare under a normal probability distribution. Even the results obtained arose from an empirical study with sampling
effort limited to eight lakes, hence lacking any manipulation of the descriptor variables and preventing a cause-effect approach.

4. Results

4.1. Limnology

Electric conductivity, dissolved oxygen and pH were significantly higher in the dry season, while ammonium and phosphate were higher in the rainy season (Table 2). In the dry season, turbidity, nitrate, nitrite and phosphate had most loadings for axis 1 (PCA), while chlorophyll-a and dissolved oxygen were most important for axis 2 (Figure 2a). During rainy season, chlorophyll-a, pH and phosphate forms characterized the axis 1, while turbidity and total alkalinity were important to axis 2 (Figure 2b). In both seasons, the locales E, H, C and F (Genipapo, Pousada das Garças, Clara and Osmar Lakes, respectively) had lower levels of nutrients and chlorophyll-a concentration, while D (Jacaré lake) showed higher levels of nutrients and chlorophyll-a concentration.

4.2. Zooplankton

Zooplanktonic community was represented by 101 species in the dry and 103 species in rainy season. During dry seasons, Lecane proiecta Hauer, 1956, Bosmina hagmanni Stingelin, 1904, and Notodiaptomus iheringi Wright, 1935, were the species most important, of each group (rotifer, cladoceran and copepod), in abundance and biomass. While in rainy season the species most important, of each group, were: Asplanchna sp., Diaphanosoma spinulosum Herbst, 1967, and Notodiaptomus amazonicus Wright, 1935.

4.3. Fishes

Thirty-four species of fish were recorded, but only eight of these (Table 3) ever had individual’s zooplankton as food item. The density of fishes ranged from four (Capivara - A) to 38,624

Table 2. Limnological features variation in the dry and rainy seasons in eight shallow lakes from Upper Paraná Floodplain. *Significant differences (p < 0.05) between seasons (Mann-Whitney test).

<table>
<thead>
<tr>
<th></th>
<th>Dry</th>
<th>Rainy</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH*</td>
<td>6.0</td>
<td>5.7</td>
</tr>
<tr>
<td>Electric conductivity (µS.cm⁻¹)*</td>
<td>23.9</td>
<td>20.3</td>
</tr>
<tr>
<td>Total alkalinity (mEq.L⁻¹)</td>
<td>75.5</td>
<td>107.7</td>
</tr>
<tr>
<td>Turbidity (NTU)</td>
<td>3.4</td>
<td>3.0</td>
</tr>
<tr>
<td>Dissolved Oxygen (mg.L⁻¹)*</td>
<td>3.5</td>
<td>7.2</td>
</tr>
<tr>
<td>Chlorophyll-a (µg.L⁻¹)</td>
<td>11.1</td>
<td>9.6</td>
</tr>
<tr>
<td>Nitrate (µg.L⁻¹)</td>
<td>0.1</td>
<td>7.8</td>
</tr>
<tr>
<td>Nitrite (µg.L⁻¹)</td>
<td>1.0</td>
<td>0.7</td>
</tr>
<tr>
<td>Ammonium (µg.L⁻¹)*</td>
<td>0.1</td>
<td>3.4</td>
</tr>
<tr>
<td>Phosphate (µg.L⁻¹)*</td>
<td>0.1</td>
<td>10.3</td>
</tr>
<tr>
<td>Dissolved phosphorus (µg.L⁻¹)</td>
<td>2.0</td>
<td>18.8</td>
</tr>
</tbody>
</table>

Zooplankton abundance ranged from 8,728 ind.m⁻³ to 4,622,461 ind.m⁻³. In dry season, a greater number of individuals was observed in the Osmar (F) (420,654 ind.m⁻³) and Genipapo lakes (E) (456,977 ind.m⁻³) (Figure 3a). On the other hand, in the rainy season, the highest abundance was registered in Capivara Lake (A) (4,622,461 ind.m⁻³) (Figure 3b).

Zooplankton biomass varied between 7,128 µg.dw.m⁻³ and 265,740 µg.dw.m⁻³. As observed for the density, the highest value of biomass was recorded in rainy season at Capivara Lake (A) (265,740 µg.dw.m⁻³), and in the dry season the highest values was in Osmar Lake (F) (252,993 µg.dw.m⁻³) (Figure 3b). Zooplankton density and biomass presented a positive association during both dry (p = 0.0154) and rainy seasons (p = 0.0001).

The individuals’ sizes in the lakes ranged from 70 to 1,260 µm. The distribution pattern of the size classes within each environment was distinct between the seasons (Figures 4 and 5). In general, the individuals of intermediate size (300-600 µm) were more abundant in the dry season (Figure 4b), whereas smaller-sized individuals (<300 µm) presented a greater numerical contribution to the community in the rainy season (Figure 5b). Considering the overall biomass, larger-sized organisms (>600 µm) were dominant in the dry season (Figure 4c).
Community abundance and biomass were positively associated to fish density during the dry season (Figure 7a and b), although only the abundance has been significant relationship, and were negatively associated in the rainy season (Figure 7c and d).

4.5. Relationship between size structure of the zooplankton and fishes

The size structure of the community did not show significant relationship with chlorophyll-a concentration in the dry season (Figure 6a and b). Otherwise, they were positively associated to chlorophyll-a concentration during rainy season (Figure 6c and d), suggesting that the increase in resource amount promotes a significant increase in the aggregate properties of the zooplankton community (biomass and density).
Figure 4. Representation of the simple regression analyses between zooplankton and fish abundance obtained in the lakes during the dry season (August-2000). a) Density of fishes (CPUE); b) Relative abundance; and c) Relative biomass. $R^2$ - coefficient of determination resultant of linear regression. Black, gray and dotted percent are >600, 301-600 and <300 size classes, respectively.

Figure 5. Representation of the simple regression analyses between zooplankton and fish abundance obtained in the lakes during the rainy season (February-2001). a) Density of fishes (CPUE); b) Relative abundance; and c) Relative biomass. $R^2$ - coefficient of determination resultant of linear regression. Black, gray and dotted percent are >600, 301-600 and <300 size classes, respectively.
community size classes was not associated to the fish densities (Figure 5).

5. Discussion

The present study evidenced the positive significant association of the density and biomass of zooplankton with fish density, as seen during the dry period (Figure 7a and b). This could indicating a numerical and mass increase of this size class with fish density (Figure 4). The percentage of larger-sized individuals, in both biomass ($R^2 = 0.86$, $p < 0.001$) and density ($R^2 = 0.69$, $p = 0.02$) was negatively associated to the fish density, reflecting a decrease of this size class with the increase of fish in the environment (Figure 4).

During rainy season, the regressions were not significant, indicating that the proportion of community size classes was not associated to the fish densities (Figure 5).

Table 3. Zooplanktivorous fishes species sampled lakes from Upper Paraná Floodplain during dry (September 2000) and rainy seasons (March 2001). The fonts are scientific records of zooplankton as food item of fish species.

<table>
<thead>
<tr>
<th>Species</th>
<th>Font</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Astyanax altiparanae</em> Garutti and Britski, 2000</td>
<td>Casatti et al., 2003; Cassemiore et al., 2002</td>
</tr>
<tr>
<td><em>Aphyocharax anisitsi</em> Eigenmann and Kennedy, 1903</td>
<td>Crippa et al., 2009; Russo and Hahn 2006; Santana-Porto and Andrian, 2009; fishbase</td>
</tr>
<tr>
<td><em>Bryconamericus stramineus</em> Eigenmann, 1908</td>
<td>Crippa et al., 2009; Santana-Porto and Andrian 2009.</td>
</tr>
<tr>
<td><em>Hyphessobrycon eques</em> Steindachner, 1882</td>
<td>Casatti et al., 2003; Crippa et al., 2009; Pelice and Agostinho, 2006; fishbase</td>
</tr>
<tr>
<td><em>Hemigrammus marginatus</em> Ellis, 1911</td>
<td>Casatti et al., 2003; Pelice and Agostinho, 2006; fishbase</td>
</tr>
<tr>
<td><em>Moenkhausia aff. Intermedia</em> Eigenmann, 1908</td>
<td>Santana-Porto and Andrian, 2009; fishbase</td>
</tr>
<tr>
<td><em>M.aff.sanctaefilomenae</em> Steindachner, 1907</td>
<td>Crippa et al., 2009; fishbase</td>
</tr>
<tr>
<td><em>Serrapinnus notomelas</em> Eigenmann, 1915</td>
<td>Pelice and Agostinho, 2006; Santana-Porto and Andrian, 2009; fishbase</td>
</tr>
</tbody>
</table>

Figure 6. Association of the total density and biomass zooplanktonic with chlorophyll-α concentration in the lakes during the dry (a and b) and rainy (c and d) seasons.
disguise the predation effect on the individuals, once this effect was remarkable only in the class of larger size (Figure 4), consequently decreasing the density and biomass larger-sized individuals and increasing density and biomass intermediate-sized individuals. The predation reduces the competitive effect between different size classes, promoting the development of populations with small-sized individuals (Wang et al., 2007) and corroborating the predation effect on the size-efficiency hypothesis (Brooks and Dodson, 1965; Dodson, 1974). This implicates a direct and negative effect of the predation on larger-sized individuals, as well as indirect and positive effect on the small-sized individuals. Several studies have registered the numerical reduction of larger-sized zooplankton individuals, mainly with sizes greater than 500 µm, as a function of a predation effect (Bramm et al., 2009; Manca et al., 2008; Masson et al., 2004; Rettig, 2003).

The positive association between the density and biomass of zooplankton with the fish density may also suggest that the predator density does not exert a predation pressure on the community; but other effect would act positively on the structure of these two communities, promoting a concordance between them, or that the predation effect is not enough to negatively influence the structure of the zooplankton community. Thus, the explanation for the increase in zooplankton density and biomass total would be the direct influence of autotrophic production, or an indirect influence, as for fish excretion, that would also influence the same production (Attayde and Hansson, 2001).

On the other hand, in the rainy period, when we did not verify a significant and direct relationship between the density of fish and the size classes of the zooplankton community (Figure 5), the biomass and density of the community were also associated with local productivity, represented by the chlorophyll-a concentration (Figure 6c and d). This association evidenced that the increase in resource availability favored increment of individuals. The productivity has been reported as the most important factor regulating the zooplankton biomass (because it reflects the increase in resource availability), but
it also has been associated to the size structure, i.e., more productive environments will present a predominance of small-sized individuals, and the inverse will be observed in less productive environments (Masson et al., 2004). Thus, in the rainy period a bottom-up effect may be more important than top-down one, on the size structure of the zooplankton community.

Masson et al. (2004) considered bottom-up and top-down processes as complementary strengths on the zooplankton structure and may act temporally on the community properties, in this case, the biomass and size structure. Similarly, Winder et al. (2003) suggest that the combination of predation and resource limitation defines the temporal variability of zooplankton size structure. Our results only indicated evidence to top-down effect in the dry period, mainly affecting the size structure of the community, in which there was a removal of larger-sized individuals.

Under the spatial perspective (among lakes), the size structure of the zooplankton community has been related to physical, chemical and biotic aspects (Cottenie et al., 2001; Masson et al., 2004; Drenner et al., 2009). The obtained results corroborated previous studies and indicate that even in tropical environments, in which zooplankton communities are compounded by small-sized species (Fernando, 1994), the size structure of the zooplankton community may be related to top-down effect. Drenner et al. (2009) ascribed the differences found in the zooplankton size structure to the presence of fish, since the greater efficiency of predation on adult individuals from larger-sized species prevent the reproduction and, in consequence, the recruitment of new individuals with larger-size in the population. Productive environments, with higher fish densities, produces a higher selection pressure on the larger-sized individuals of zooplankton, while a lower density of fish would decrease the predation effect, allowing a greater survival of these larger-sized individuals (Cottenie et al., 2001).

Small zooplankton generally ate fine particles (1-5 μm), whereas larger zooplankton generally consumed larger particles (Gliwicz, 1969). Large populations of smaller-sized individuals are related to a greater efficiency of these organisms in the assimilation of smaller organisms, as bacteria, nano-phytoplankton and small planktonic protozoan, suggesting a greater importance of the "microbial loop" in the energy transfer. On the other hand, when there is a numerical predominance of larger-sized individuals, it can mean that producer organisms with larger size are present in the environment, as filamentous algae, suggesting that the base of the food chain is herbivory. In this way, the seasonality differences of our results would enable to indicate other scientific investigation process, where the climatic periods would modify the energy transfer process; an alternative to support this hypothesis is the evaluation of the nano- and micro-planktonic communities. It is clear that the evaluation of these hypotheses depends on a better understanding of the relationship between zooplankton size and food size.

The results add information about biotic mechanisms (which influence the zooplankton size structure in temperate lakes, where the size gradient among the species is noticeable) that may also be extrapolated to tropical environments, which are compounded by zooplankton communities with a mean size inferior to that in temperate regions. In summary, we verified that the main hypothesis of this study was partially confirmed because it was supported only one sampling season; however this suggests that different mechanisms act on the size structure of the zooplankton community between the analyzed periods.

Acknowledgements

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