Host-parasite and phylogenetic relationships of *Myxobolus filamentum* sp. n. (Myxozoa: Myxosporea), a parasite of *Brycon orthotaenia* (Characiformes: Bryconidae) in Brazil

Juliana Naldoni1, Suellen A. Zatti1, Kassia R.H. Capodifoglio1, Tiago Milanin1, Antônio A.M. Maia2, Marcia R.M. Silva2 and Edson A. Adriano1,3

1 Animal Biology Department, Institute of Biology, University of Campinas, São Paulo, Brazil; 2 Veterinary Medicine Department, College of Animal Science and Food Engineering, São Paulo University, São Paulo, Brazil; 3 Biology Science Department, Federal University of São Paulo, São Paulo, Brazil

**Abstract:** *Myxobolus filamentum* sp. n. was found infecting gill filaments of three of 39 *Brycon orthotaenia* Günther specimens examined (8%), which were taken from the river São Francisco in Minas Gerais state, Brazil. Plasmodia of the parasite were white and long, measuring 5 mm in length. Mature spores of *M. filamentum* sp. n. were oval from the frontal view and biconvex from the lateral view, measuring 7.5–9.7 µm (9.0 ± 0.3 µm) in length and 5.2–7.3 µm (6.2 ± 0.4 µm) in width. The polar capsules were elongated and equal in size, measuring 3.8–5.5 µm (4.7 ± 0.3 µm) in length and 1.3–2.2 µm (1.7 ± 0.1 µm) in width. The development of the parasite led to compression of the adjacent tissues and inflammatory infiltrate with granulocytic cells. Ultrastructural observation revealed that the plasmodia were delimited by two membranes, which had numerous and extensive pinocytotic channels extending into the wide ectoplasm zone. The plasmoidal wall exhibited abundant villi-like projections and a thin layer of granular material prevented direct contact between the plasmodial wall and the host tissue. Phylogenetic analysis, based on 18S rDNA, showed *M. filamentum* sp. n. as a sister species of *Myxobolus oliveirai* Milanin, Eiras, Arana, Maia, Alves, Carrierio, Ceccarelli et Adriano, 2010, a parasite of other fish species of the genus *Brycon* Müller et Troeschel from South America.

**Keywords:** taxonomy, 18S rRNA, ultrastructure, granulocytic cells, freshwater fish, Neotropical Region

Myxosporeans are among the most important pathogens for both wild and farmed fish. Most species use fish as host, although a small number have been described infecting amphibians, reptiles, birds and mammals (Eiras et al. 2005, Prunescu et al. 2007, Bartholomew et al. 2008). The genus *Myxobolus* Bütschli, 1882 is the most speciose group within the myxozoans, with more than 850 species described (Eiras et al. 2014). So far, 37 *Myxobolus* species have been described infecting fish species in South America (Eiras et al. 2005, 2014) and nine species have been reported for fish of the family Bryconidae (Characiformes) (Azevedo et al. 2010, Milanin et al. 2010, Carrierio et al. 2013).

The four following species were associated with *Brycon hilarii* (Valenciennes) in Brazilian Pantanal wetlands: *Myxobolus oliveirai* Milanin, Eiras, Arana, Maia, Alves, Silva, Carrierio, Ceccarelli et Adriano, 2010 and *Myxobolus brycon* Azevedo, Casal, Marques, Silva et Matos, 2011, which are parasites of gill filaments; *Myxobolus umidas* Carriero, Adriano, Silva, Ceccarelli et Maia, 2013, a parasite of the spleen; and *Myxobolus piratangarum* Carriero, Adriano, Silva, Ceccarelli et Maia, 2013, a parasite of the kidney (Milanin et al. 2010, Azevedo et al. 2011, Carrierio et al. 2013).

*Brycon orthotaenia* Günther, popularly known as ‘matrinxã’, is endemic to the river São Francisco basin in the southeast/northeast of Brazil, and is among the native species with a high risk of extinction (Godinho and Godinho 1994). This species has omnivorous feeding habits, can reach up to 7 kg in body weight and has great potential in regional aquaculture and fisheries (Sato et al. 2003, MPA 2012).

The aim of the present study was to describe, based on morphology, ultrastructure and sequencing of the 18S rRNA gene, a new *Myxobolus* species that infects the gill filaments of *B. orthotaenia* caught in the River São Francisco in the state of Minas Gerais, Brazil.

**MATERIALS AND METHODS**

Thirty-nine specimens of *Brycon orthotaenia* were collected from the River São Francisco (17°12′0.26″S; 44°50′0.45″W) in the municipality of Pirapora, state of Minas Gerais, Brazil. Sam-
amples were collected in July 2010 (n = 5), July (n = 5) and December 2011 (n = 12), December of 2012 (n = 12) and December 2013 (n = 5). After capture, the fish were immediately transported alive to a nearby field laboratory, where they were euthanised by a benzocaine overdose. The methodology of the present study was approved by the ethics research committee of the State University of Campinas under protocol number 2334-1, in accordance with Brazilian law (Federal Law No. 11.794, dated 8 October 2008 and Federal Decree No. 6899, dated 15 July 2009).

Plasmodia with mature spores were examined in fresh mounts with a light microscope. The morphological characterisation of the spores was based on mature spores obtained from three different host specimens. Measurements were taken from 34 spores using a computer equipped with Axivision 4.1 image capture software, coupled to an Axioplan 2 Zeiss Microscope. The dimensions of the spores were expressed as range with (mean ± standard deviation), all in μm. Smears containing free spores were air-dried and stained with Giemsa solution and mounted in a low-viscosity mounting medium (CytosealTM, Hatfield, PA, USA) on permanent slides for their deposition in the museum. For histological analysis, fragments of infected organs were fixed in 10% buffered formalin and embedded in paraffin. Serial sections with a thickness of 4 μm were stained with hematoxylin-eosin.

For transmission electron microscopy, the plasmodia were fixed in cold (4°C) 2.5% glutaraldehyde and 0.1 M sodium cacodylate buffer (pH 7.4) for 12 h, washed in a glucose-saline solution for 2 h and post-fixed in OsO₄. After dehydration using an acetone series, the material was embedded in EMBed 812 resin. Semithin sections were stained with toluidine blue solution and acetone series, the material was embedded in EMbed 812 resin. Ultrathin sections, double stained with uranyl acetate and lead citrate, were examined in an LEO 912 electron microscope at 60 kV. For sequencing, the plasmodia content was collected in a 1.5-ml microcentrifuge tube and the DNA was extracted using the DNeasy® Blood & Tissue kit (Qiagen, Valencia, CA, USA), following the manufacturer’s instructions. The product was quantified in a NanoDrop 2000 spectrophotometer (Thermo Scientific, Wilmington, USA) at 260 nm.

The polymerase chain reaction (PCR) was carried out using a final volume of 25 μL, which contained 10–50 ng of extracted DNA, 1 x Taq DNA polymerase buffer, 0.2 mmol of dNTP, 1.5 mmol of MgCl₂, 0.2 pmol of each primer, 0.25 μL (1.25 U) of Taq DNA polymerase (all reagents from Invitrogen by Life Technologies, Carlsbad, CA, USA) and ultrapure (MilliQ) water in an Eppendorf AG 22331 Hamburg Thermocycler (Eppendorf, Hamburg, Germany). The 18S rRNA gene was amplified with the primer pairs ERIB1 (Barta et al. 1997) – ACT1r (Hallett and Diamant 2001) and Myxgen4F (Diamant et al. 2004) – ERIB10 (Barta et al. 1997), which amplified two fragments of approximately 1000 bp and 1200 bp, respectively. An initial denaturation stage at 95°C for 5 min was followed by 35 cycles of denaturation at 95°C for 60 s, annealing at 58°C for 60 s, extension at 72°C for 120 s, finishing with an extended elongation stage at 72°C for 5 min. PCR products were electrophoresed in 1.0% agarose gel, stained with GelRed® and analysed by an FLA-3000 (Fugi) scanner. The size of the amplicons was estimated by comparison with the 1 kb DNA Ladder (Invitrogen by Life Technologies, CA, USA).

The PCR products were purified with QIAquick® PCR Purification Kit (Qiagen) and then were sequenced using the same primer pairs that were used in the amplification stage, plus primers MC5 and MC3 (Molnár et al. 2002) with the BigDye® Terminator v3.1 Cycle Sequencing Kit (Applied BiosystemsTM, Foster City, CA, USA) in an ABI 3730 DNA sequencing analyser (Applied BiosystemsTM Inc.). The sequence obtained was visualised, assembled and edited using BioEdit 7.1.3.0 software (Hall 1999). The ambiguous bases were clarified using corresponding ABI chromatograms. A standard nucleotide BLAST search was then conducted (Altschul et al. 1997). The BioEdit program (Hall 1999) was used to align the sequence obtained and to compare it with the sequences of species of Myxobolus and Henneguya Thélohan, 1892, recovered by BLAST, plus all sequences of South American species of these genera that did not appear in the BLAST search. Phylogenetic analyses were performed using the Maximum likelihood (ML) and Maximum parsimony (MP) methods.

In order to obtain the evolutionary model that best fit the sequences studied here, jModelTest analysis was performed using the Akaike information criterion (Posada 2008). ML analysis was performed with PhyML3.0 (Guindon et al. 2010) using the General Time Reversible substitution model and 4 gamma-distributed ration variations among sites. MP analysis was conducted with PAUP* 4.0b10 (Swoford 1998) using the heuristic search algorithm and tree bisection-reconnection (TBR) branch swapping. Myxidium anatidium Bartholomew, Atkinson, Hallett, Lowens-tine, Garner, Gardiner, Rideou, Keel et Brown, 2011 (EF602629) and Myxidium hardella Garner, Bartholomew, Whipple, Nordhausen et Raitt, 2005 (AY688957) were used as an outgroup in this analysis. In both analyses, clade support was assessed by bootstrapping with 1000 replicates. The trees obtained were initially visualised with FigTree v1.3.1 (Rambaut 2008) and edited with Adobe Illustrator (Adobe Systems Inc., San Jose, USA). Only bootstrap values above 50 were considered to be well-supported. The data of the family and/or order of host fish were obtained from Froese and Pauly (2013).

Distance analysis was performed to compare the new species with the most closely related species in the phylogenetic tree, using a p-distance model distance matrix for transitions and transversions in MEGA 5.0 (Tamura et al. 2011).

RESULTS

From total of 39 specimens of Brycon orthotaenia caught in the river São Francisco, the gill filaments of 3 (7.69%) had been parasitised by an unknown species of Myxobolus, which is described below.

Whitish plasmodia, elongated in shape and measuring up to 5 mm, were located in the gill filaments of B. orthotaenia. Histopathological analysis showed its development in the subepithelial connective tissue of the gill filaments. The development of the parasite led to compression of the adjacent connective and epithelial tissues and numerous granulocytic cells were observed at the infection site (Fig. 1A,B). The plasmodia exhibited asynchronous development, with mature spores in the central region and young sporogenous stages in the peripheral area (Fig. 1B).

Ultrastructure analysis revealed that the plasmodia were delimited by two membranes, which had numerous and
extensive pinocytotic channels of approximately 5 µm in length extending into the ectoplasm zone (Fig. 2A–D). The plasmodial wall exhibited abundant villi-like projections and a thin layer of granular material prevented direct contact between the plasmodial wall and the host cells of the adjacent connective tissue (Fig. 2A–C). Several mitochondria were observed in the ectoplasm. Early stages of sporogenesis and advanced spore developmental stages were noted below the ectoplasm, with sporoblasts containing two spores (Fig. 2A,B). Sporoplasm were binucleate and had small and scarce sporoplasmomes (Fig. 2D).

**Myxobolus filamentum** sp. n.  

**Description.** Mature spores pear-shaped in frontal view, measuring 7.5–9.7 (9.0 ± 0.3) in length, 5.2–7.3 (6.2 ± 0.4) in width and 4.8–5.7 (5.3 ± 0.3) in thickness (Figs. 1C, 3). In lateral view, spores biconvex, with symmetrical valves. Polar capsules elongated and equal in size, measuring 3.8–5.5 (4.7 ± 0.3) in length and 1.3–2.2 (1.7 ± 0.1) in width. Polar capsule occupies about half of spore (Figs. 1C, 3). Polar filaments with 10–11 coils arranged perpendicularly to longitudinal axis of polar capsule (Figs. 2D, 3).

**Type host:** *Brycon orthotaenia* Günther (Characiformes: Bryconidae), 37–42 cm (39 ± 4 cm).

**Type locality:** River São Francisco, municipality of Pirapora (17°12'8.26’S; 44°50'0.45’W), Minas Gerais state, Brazil. The type specimens of *Myxobolus filamentum* sp. n. was collected in December 2013.

**Site of infection:** Gill filament.

**Prevalence:** 3 of 39 specimens, i.e. 8%, infected.

**Specimens deposited:** One slide containing spores fixed with methanol, stained with Giemsa and mounted in a low-viscosity mounting medium, deposited in the collection of the Museum of Zoology, Institute of Biology, Universidade Estadual de Campinas, São Paulo, Brazil (Accession No. ZUEC Myx 46).

**DNA sequence:** 18S rRNA sequence deposited in the GenBank database under accession number KJ849240.

**Etymology:** The specific name refers to the infection site of the parasite, gill filaments, given that filamentum in Latin means filament.

**Remarks.** *Myxobolus filamentum* sp. n. was compared with all species of *Myxobolus* that have been described from South American freshwater fish and other continents (Eiras et al. 2005, 2014). So far, no species of myxosporean has been described from *B. orthotaenia*. Nevertheless, *M. oliveirai*, *M. brycon*, *M. piraputangae* and *M. umidus* have been reported from the congener *B. hilarii*. However, the spores of these four species differ morphometrically in some aspects from *M. filamentum* (Table 1), although *M. oliveirai* is the one that most resembles the species described herein, with both exhibiting a lightly-tapered anterior end and pear-shaped spores from the frontal view. Al-
Fig. 2. Electron micrographs of plasmodia of *Myxobolus filamentum* sp. n. from gill filaments of *Brycon orthotaenia*. A – Numerous pinocytic channels (pn) linking the ectoplasm (ec) to the plasmodial wall (large black arrows). Numerous mitochondria (m), generative cells (gc), young sporoblasts (sb) containing sporogenous cells (sc) and their nuclei (n). See immature spores (isp) with polar capsules (pc) in the innermost layer, in some cases containing polar filaments (white arrow). B – host parasite interface showing abundant villi-like projections of the plasmodial wall (large arrow), wide ectoplasm (ec) containing numerous and long pinocytic channels (thin arrows) and mitochondria (m). C – magnified part of B showing a layer of granular material (asterisks) surrounding the plasmodial wall composed by two membranes (black arrows). D – disporic sporoblasts (sp) sections containing immature spores (is) with polar capsules (pc) containing internalised (empty arrow) or still externalised polar filament (thin black arrow), sporoplasm (spl) with two nuclei (n) and few and small sporoplasmosomes (thin white arrow). Remaining abbreviations: H – host; P – plasmodium; Vm – valve-forming material; Ms – mature spores.

Fig. 3. Schematic representation of the mature spore of *Myxobolus filamentum* sp. n. infecting gill filament of *Brycon orthotaenia* in Brazil.

high enough to distinguish the two species. With regards to other species of *Myxobolus* from freshwater fish in South American or other continents, *M. filamentum* differs in at least one of the morphometric/morphological characteristics, such as the shape and size of the plasmodia, the shape and size of the spores, the number of polar coils, the host and the site of infection (Eiras et al. 2005, 2014). Therefore, based on all of the evidence above, we propose that the species studied herein is a new species of *Myxobolus*.

Molecular data: Sequencing of the 18S rRNA gene of *M. filamentum* resulted in a partial sequence of 1 885 bp. The most similar sequence to the species described here in according to BLAST was *M. oliveirai* (HM754633).

Maximum likelihood and Maximum parsimony phylogenetic trees showed species of *Myxobolus/Henneguya* clustering in two distinct clades. The smallest clade (A) was a cluster of parasites of characiform fishes (families Serrasalmidae and Bryconidae), all from South America. Clade B is further divided to form two clades: a small clade (B1), composed of *Myxobolus* species from salmonids, with *Myxobolus cordeiroi* Adriano, Arana, Alves, Silva,
Table 1. Comparison data of *Myxobolus filamentum* sp. n. with other species of *Myxobolus* from bryconid fish.

<table>
<thead>
<tr>
<th>Species of Myxobolus</th>
<th>Spore length</th>
<th>Spore width</th>
<th>Thickness</th>
<th>LPC</th>
<th>WPC</th>
<th>NCF</th>
<th>Site of infection</th>
<th>Host</th>
<th>Locality</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>M. filamentum</em> sp. n.</td>
<td>7.4–9.7 (9.0 ± 0.3)</td>
<td>5.1–7.3 (6.2 ± 0.4)</td>
<td>4.8–5.7 (5.3 ± 0.3)</td>
<td>3.7–5.4 (4.7 ± 0.3)</td>
<td>1.2–2.2 (1.7 ± 0.1)</td>
<td>10</td>
<td>gill filaments</td>
<td>Brycon orthotaenia Günther</td>
<td>River São Francisco, Brazil</td>
<td>Present study</td>
</tr>
<tr>
<td><em>M. aureus</em></td>
<td>12.6 ± 0.5</td>
<td>8.3 ± 0.3</td>
<td>5.5 ± 0.3</td>
<td>5.7 ± 0.3</td>
<td>2.9 ± 0.2</td>
<td>7–8</td>
<td>liver</td>
<td><em>Salminus brasiliensis</em> (Cuvier)</td>
<td>PW, Brazil</td>
<td>Carriero et al. 2013</td>
</tr>
<tr>
<td><em>M. brycon</em></td>
<td>6.9 (6.5–7.2)</td>
<td>4.2 (3.9–4.8)</td>
<td>2.5 (1.9–2.8)</td>
<td>4.2 (3.8–4.7)</td>
<td>1.9 (1.7–2.5)</td>
<td>8–9</td>
<td>gills</td>
<td><em>Brycon hilarii</em> (Valenciennes)</td>
<td>PW, Brazil</td>
<td>Azevedo et al. 2011</td>
</tr>
<tr>
<td><em>M. macroplasmodialis</em></td>
<td>10.5–12 (11)</td>
<td>8.9 (8.5)</td>
<td>5.5–5.5 (5.2)</td>
<td>4.5 (4.5)</td>
<td>2.3 (2.8)</td>
<td>6</td>
<td>abdominal cavity</td>
<td><em>Salminus brasiliensis</em></td>
<td>River Mogi, Brazil</td>
<td>Molnár et al. 1998</td>
</tr>
<tr>
<td><em>M. oliveirai</em></td>
<td>11.2 ± 0.4</td>
<td>7.4 ± 0.5</td>
<td>4.6 ± 0.6</td>
<td>5.6 ± 0.2</td>
<td>2.3 ± 0.2</td>
<td>6–8</td>
<td>gills</td>
<td><em>Brycon hilarii</em></td>
<td>PW, Brazil</td>
<td>Milanin et al. 2010</td>
</tr>
<tr>
<td><em>M. pantanalensis</em></td>
<td>9.3 ± 0.4</td>
<td>6.5 ± 0.4</td>
<td>N/A</td>
<td>4.2 ± 0.5</td>
<td>2.0 ± 0.1</td>
<td>4–5</td>
<td>gill filaments</td>
<td><em>Salminus brasiliensis</em></td>
<td>PW, Brazil</td>
<td>Carriero et al. 2013</td>
</tr>
<tr>
<td><em>M. paranensis</em></td>
<td>12–15</td>
<td>7–8</td>
<td>N/A</td>
<td>6–7</td>
<td>2.5</td>
<td>N/A</td>
<td>testes, ovary</td>
<td><em>Salminus brasiliensis</em></td>
<td>Argentina</td>
<td>Bonetto et Pignalberi 1965</td>
</tr>
<tr>
<td><em>M. piraputangae</em></td>
<td>10.1 ± 0.5</td>
<td>8.7 ± 0.5</td>
<td>6.7 ± 0.3</td>
<td>5.2 ± 0.4</td>
<td>3.0 ± 0.3</td>
<td>4–5</td>
<td>kidney</td>
<td><em>Brycon hilarii</em></td>
<td>PW, Brazil</td>
<td>Carriero et al. 2013</td>
</tr>
<tr>
<td><em>M. salminus</em></td>
<td>10.1 ± 0.4</td>
<td>6.1 ± 0.4</td>
<td>5.0 ± 0.6</td>
<td>4.6 ± 0.2</td>
<td>1.7 ± 0.1</td>
<td>7–8</td>
<td>gills</td>
<td><em>Salminus brasiliensis</em></td>
<td>PW, Brazil</td>
<td>Adriano et al. 2009</td>
</tr>
<tr>
<td><em>M. umidus</em></td>
<td>13.5 ± 0.7</td>
<td>7.8 ± 0.4</td>
<td>7.7 ± 0.1</td>
<td>5.1 ± 0.4</td>
<td>2.7 ± 0.3</td>
<td>4–5</td>
<td>spleen</td>
<td><em>Brycon hilarii</em></td>
<td>PW, Brazil</td>
<td>Carriero et al. 2013</td>
</tr>
</tbody>
</table>

LPC – length of polar capsules; N/A – no data; NCF – number of coils of polar filaments; PW – Pantanal wetland; WPC – width of polar capsules.

Ceccarelli, Silva et Maia, 2009, a parasite of siluriform pineliods in South America, as a basal species, and a large clade (B2), composed of species of *Myxobolus/Henneguya* from several families/families of fishes. In this large clade, *M. filamentum* clustered as a sister species of *M. oliveirai* in a clade that also contained species from percids and esocids (Fig. 4). The p-distance analysis showed that *M. oliveirai* was the species with the closest genetic similarity to *M. filamentum*, with a similarity of 95.6%.

DISCUSSION

The development of plasmodia of *Myxobolus filamentum* sp. n. caused compression of the adjacent connective and epithelial tissues, deforming the neighbouring structures, as previously recorded in *Henneguya mystusia* Salkar, 1985, *Myxobolus diversicapsularis* Slukhiai 1966, and *Henneguya pseudoplatsymena* Naldoni, Arana, Maia, Ceccarelli, Tavares, Borges, Pozo et Adriano, 2009. Inflammatory infiltrate with numerous granulocytic cells was observed in adjacent areas of the plasmodium, although the fish did not exhibit signs of sickness. Inflammatory infiltrate, composed of a mixed population of granulocytes including neutrophils and cells that resembled eosinophils, has been observed in proliferative gill disease (PGD) caused by species of *Henneguya* in the pond-reared channel catfish *Ictalurus punctatus* Rafinesque in North America (Lovet et al. 2011).

Ultrastructural analysis revealed that the plasmodial wall was delimited by two membranes, similar to *Myxobolus metynnis* Casal, Matos et Azevedo, 2006, *M. salminus* and *M. brycon*, which exhibited numerous and extensive pinocytic channels extending into the wide ectoplasmic zone (Casal et al. 2006, Adriano et al. 2009, Azevedo et al. 2010). The plasmodial wall also exhibited abundant villiform projections, similar to those reported for infection of host with *M. metynnis*, increasing surface area of the parasite (Casal et al. 2006). There was also a thin layer of granular material preventing contact between the plasmodium and host cells of the adjacent tissues, as observed in host infected with *Henneguya cuculator* Naldoni, Maia, Silva et Adriano, 2014 (see Naldoni et al. 2014). The plasmodial exhibited intense nutritional and metabolic activity, with numerous and extensive pinocytic channels in the ectoplasm, as well as a layer immediately below the ectoplasm with large numbers of mitochondria.

With regards to sporogenesis, this process followed the pattern of other species of *Myxobolus*, with the early stages of spore formation in the periphery of the plasmodium and mature spores located centrally (Current et al. 1979, Casal et al. 1997, 2002, Adriano et al. 2006). *Myxobolus filamentum* is the first myxosporean species reported from *Brycon orthotaenia*, the fifth in fish of the genus *Brycon* Muller et Troescher and the ninth of bryconid fish (Azevedo et al. 2010, Carriero et al. 2013, Moreira et al. 2014). Of
these eight myxosporean species previously reported from bryconids, 18S rRNA sequences of seven species are available in GenBank and were used in the phylogenetic analyses of the present study.

Maximum likelihood and maximum parsimony analyses using the sequences of *M. filamentum* and 58 species of *Henneguya* and *Myxobolus* and two of species *Myxidium* retrieved from GenBank produced phylograms with a nearly identical topology. Species of *Myxobolus* and *Henneguya* from bryconids appeared in two distinct clades. In the clade A, which was composed exclusively by species of *Myxobolus* from characiform fish, parasites of bryconids were dominant. Inside this clade, *Myxobolus cf. colossomatis* Molnár et Békési, 1993 and *Myxobolus macroplasmodialis* Molnár, Ranzani-Paiva, Eiras et Rodrigues, 1998 clustered as sister species, yet with low bootstrap val-
ues in ML analysis, whereas the former species appears as the basal species of the subclade composed by species of Myxobolus from bryconids in MP analysis (Fig. 4).

Clade B2 contained four other species, parasites of bryconids, which clustered in separate subclades. These were M. pantanalis from Brycon hilarii and Henneguya rotunda Moreira, Adriano, Silva, Ceccarelli and Maia, 2014 from Salminus brasiliensis, which clustered as sister species in a small subclade composed of parasites of characiform hosts, whereas M. filamentum appeared in another subclade as a sister species of M. oliveirai, a parasite of gill filaments of the congenic B. hilarii. However, clade B2 presents several internal nodes with low bootstrap value. Nevertheless, despite the fact that some studies show the tendency of myxosporeans to group according to the phylogenetic relationships of their fish hosts (Carriero et al. 2013, Shin et al. 2014), the results of the present study show polyphyletic clustering of the parasites of bryconids as well as of those from all characiform fish. This is in accordance with the results obtained by Moreira et al. (2014), who reported the polyphyletic clustering of the parasites of characiform hosts. In this context, future efforts aiming to describe new species of myxosporean parasites of characiform hosts, and their 18S rRNA sequencing, will be of great importance to clarify the evolutionary relationship of these parasites within the Myxozoa.

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