Myxozoan infections in fishes of the Tasik Kenyir Water Reservoir, Terengganu, Malaysia

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ABSTRACT: During a survey on fishes of the Tasik Kenyir Reservoir, Malaysia, 5 new *Myxobolus* spp. and 2 known *Henneguya* spp. were found. The specific locations for 2 *Myxobolus* spp. were the host's muscles, while 2 other *Myxobolus* spp. were found to develop in the host's kidney and gills, respectively. Of the species developing intracellularly in muscle cells, *M. terengganuensis* sp. nov. was described from *Osteochilus hasselti* and *M. tasikkenyirensis* sp. nov. from *Osteochilus vittatus*. *M. csbai* sp. nov. and *M. osteochili* sp. nov. were isolated from the kidney of *Osteochilus hasselti*, while *M. dykovae* sp. nov. was found in the gill lamellae of *Barbonymus schwanenfeldii*. *Henneguya shaharini* and *Henneguya hemibagri* plasmodia were found on the gills of *Oxyeleotris marmoratus* and *Hemibagrus nemurus*, respectively. Description of the new and known species was based on morphological characterization of spores, histological findings on locations of plasmodia and DNA sequence data.

KEY WORDS: New species · Myxosporea · *Myxobolus* · *Henneguya* · Histology · 18S rDNA

INTRODUCTION

Research on myxosporean fish parasites is a fast-developing field of ichthyoparasitology. Numerous myxosporeans, among them highly pathogenic species, are known from freshwater and marine fishes. In terms of freshwater fishes, the majority of studies both in freshwater and culture systems deal with fishes of the northern hemisphere (Eurasia and North America), while studies on myxosporeans of marine fishes have mostly been done worldwide on cage-cultured species. Among myxosporeans, the genus *Myxobolus* includes the highest number of species. In a synopsis of this genus, Eiras et al. (2005) reported 751 valid species, while Lom & Dyková (2006) counted 792 known species. *Henneguya* spp. also belong to the commonest parasites of fishes, and in his review of this group Eiras (2002) listed 146 *Henneguya* spp. mostly from fishes of the temperate climate zones. The genus *Thelohanellus* is a less well studied myxosporean group but several of the known species have been precisely described from South Asia. Most of the papers written on myxosporean parasites of the tropical and subtropical countries have been presented by Indian authors (Chakravarty 1939, Tripathi 1952, Lalitha-Kumari 1965, 1969, Seenappa & Manohar 1981, Sarkar 1985, 1986, 1995, Sarkar et al. 1985, Basu & Haldar 2003, 2004). Data on myxosporean infections were also presented by Ky (1971) from Vietnam and a review of the known fish parasites of this country was written by Arthur & Te (2006). Studies on several species were also done in Thailand. From that country, first Lom et al. (1991) described a new genus, *Hennegoides*, with *Hennegoides longitudinalis* as type species, then Supamattaya et al. (1991) reported on *Sphaerospora epinepheli* from the grouper *Epinephelus malabaricus* cultured in marine cages. A third species is *S. ojiroveci* from an aquarium fish originating from Thailand, but described by Lom & Dyková (1997) in Europe. More recently, *M. balantiocheili*, a species causing neurological symptoms, was described from the brain of *Balantiocheilos melanopterus* in Thailand (Levsen et al. 2004). The first paper on a myxosporean infection of Malaysian fishes was written by Shariff (1982), who presented a complex morphological and pathological study on the new species *Henneguya shaharini*. 
Recently, additional myxosporeans have been found by Molnár et al. (2006a,b) from Pangasius hypophthalmus and Hemibagrus nemurus. They described 3 Hennegeois, 2 Myxobolus and 4 Henneguya species from these fishes.

Lake Tasik Kenyir was formed by the construction of 2 major dams in 1982. The total water surface area of the lake is about 36 900 ha, with a surrounding catchment area of 260 000 ha. The shoreline of this lake is rather steep since it was formed in a hilly terrain. The average depth of the lake is about 100 m. Surface temperatures of the lake do not fluctuate very much (around 28 to 30°C), while in the upstream portions of the feeding rivers, water temperatures as low as 23.0°C have been recorded. The fish fauna that once dominated the upstream and downstream rivers has presumably adapted to the lacustrine habitat. At least 38 fish species are known to occur in the lake, 17 of which are economically important for their market price or aquarium trade value (Zulkafli & Ashhar 2000).

The aim of the present study was to get information on myxosporean infections of fishes in the southeast Asian region. Five new Myxobolus spp. are described and the occurrence of 2 known Henneguya spp. is recorded from fishes of Tasik Kenyir, and the site selection in the infected organs and phylogenetic positions of these parasites are discussed.

MATERIALS AND METHODS

Study location and sample preparation. Fishes were collected with the help of gill nets in the Tasik Kenyir Water Reservoir on 20 and 21 April 2007 (area ‘Anglers Paradise’, 5° 12.057’ N, 102° 40.159’ E). The following fishes of various size and age classes (a total of 15 species) were examined for myxozoan infections on the spot or 1 to 3 d later in the laboratory of the Institute of Tropical Aquaculture (AQUATROP), University Malaysia Terengganu (UMT); Neolissochilus (Acrosocheilus) hexagonolepis (copper mahseer, tengas) (n = 2), Cyclocheilichthys apogon (Indian river barb, temperas) (n = 6), Epalzeohynchos kalopterus (selimang) (n = 2), Hampala macrolepidota (hampala barb, sebarau) (n = 5), Labiobarbus lineatus (kawan) (n = 4), Mystacoleucus marginatus (sia) (n = 5), Osteochilus vittatus (bonylip barb, rong) (n = 4), Osteochilus hasselti (silver shark minnow, terbol) (n = 10), Barbonymus schwanenfeldii (tinfoil barb, lampam sungai) (n = 10), Oxyleotris marmoratus (marble goby, ketutu) (n = 2), Pristolepis fasciatus (patung) (n = 1), Notopterus notopterus (bronze featherback, belida) (n = 3), Hemibagrus nemurus (Asian redtail catfish, baung) (n = 2), Channa micropeltes (snakehead, toman) (n = 2) and Osphronemus gouramy (giant gouramy, kalui) (n = 1).

After surveying the skin, the fins and the gills in toto, the gills were cut out, the abdominal cavity opened, the intestine, spleen, liver, gonads and kidneys separated and checked mostly with the unaided eye and in some cases by Zeiss Stemi DV4 stereomicroscope and Motic DMBA300 digital microscope. Detailed (microscopic) investigations were performed only on the gills and the muscles. The hemibranchial area was studied in detail. From the muscles only, several cubes of about 1 g were cut out from 3 different parts of the body.

The gill filaments of each hemibranchium were checked for myxozoan plasmodia under a Zeiss Stemi DV4 stereomicroscope. Plasmodia in filaments, lamellae or in the gill arches were differentiated according to Molnár (2002) into epithelial, vascular, muscular and chondroid locations. Plasmodia were carefully removed from the tissues and opened with a fine needle in a slide. Some of the spores obtained from mature plasmodia were studied as fresh preparations under a Motic DMBA300 digital microscope, and other spores obtained from the same plasmodia were fixed in 70% ethanol in vials for further morphological and molecular biological examination. Pieces from the muscles were compressed between 2 glass slides and studied as squash preparations under a compound microscope. Photos of fresh spores were taken with a Motic DMBA300 digital microscope. Infected organs (hemibranchia of gills, kidney and muscles) were fixed in Bouin’s solution for 4 h, washed in 80% ethanol several times, embedded in paraffin wax, cut into 5 to 8 µm thick sections and stained with hematoxylin and eosin. Photos of histological sections were taken in Hungary with an Olympus BH-2 microscope equipped with a DP-10 digital camera. Measurements were taken of 25 photographed fresh spores according to the guideline suggested by Lom & Arthur (1989).

Molecular methods. For DNA extractions, samples preserved in ethanol were centrifuged at 5000 × g for 5 min to pellet the myxospores, then the ethanol was removed. The DNA was extracted using a QIAGEN DNeasy tissue kit (animal tissue protocol; Qiagen) and eluted in 50 µl AE buffer.

The 18S rDNA was amplified using the primers 18e and 18g’ (Table 1) in a 25 µl reaction mixture comprising 1 µl of extracted genomic DNA, 5 µl of 1 mM deoxyribonucleotide triphosphates (dNTPs, MBI Fermentas), 0.25 µl of each primer (50 pmol µl⁻¹), 2.5 µl of 10x Taq buffer (MBI Fermentas), 2 µl of 25 mM MgCl₂, 1 µl of Taq polymerase (2 U) (MBI Fermentas) and 13 µl of distilled water. The PCR cycle consisted of an initial denaturation step of 94°C for 4 min, followed by 35 cycles of 94°C for 30 s, 56°C for 30 s, 72°C for 80 s, and
was finished with terminal extension at 72°C for 7 min, then rested at 4°C. This was followed by a second round of PCR with the SphF-SphR primer pair (Table 1). The total volume of the nested PCR reactions was 50 µl, which contained 1 µl of amplified DNA, 10 µl of 1 mM dNTPs, 0.5 µl of each primer, 5 µl of 10× Taq buffer, 2.5 µl of 25 mM MgCl₂, 2 µl of Taq polymerase (2 U) and 28.5 µl of DEPC water. Amplification conditions in the second round were 94°C for 50 s, 56°C for 50 s, 72°C for 60 s for 35 cycles, and the cycle was terminated with an extension period at 72°C for 10 min, then rested at 4°C. Both PCR cycles were performed in a PTC-200 thermocycler (MJ Research). The PCR products were electrophoresed in 1.0% agarose gels in tris-acetate-EDTA (TAE) buffer gel stained with 1% ethidium bromide and then purified with the PCR-M Clean Up System (Viogene).

Purified PCR products were sequenced in both directions with the primers listed in Table 1, using the ABI BigDye Terminator v3.1 Cycle Sequencing Kit with an ABI 3100 Genetic Analyser.

The various forward and reverse sequence segments were aligned in BioEdit (Hall 1999) and ambiguous bases clarified using corresponding ABI chromatograms. Nucleotide sequences were aligned using the software CLUSTAL W (Thompson et al. 1994). The alignment was corrected manually using the alignment editor of the software MEGA 4.0 (Tamura et al. 2007). DNA sequence similarities were calculated with the Sequence Identity Matrix of the software BioEdit. Phylogenetic calculations were performed with PAUP 4.0b10 (Swofford 2000). The data were analyzed with maximum parsimony (close-neighbor-interchange [CNI] level 3, random addition trees 100). Also neighbor-joining was calculated by MEGA 4.0 using the Tamura-Nei model and pairwise deletion for gaps. Clade support was assessed with bootstrapping (100 replicates for maximum likelihood and maximum parsimony and 1000 replicates for neighbor-joining). *Ceratomyxa shasta* was chosen as an outgroup.

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### RESULTS

Myxozoan infection was found in 5 out of the 15 fish species examined. Plasmodia with mature *Myxobolus* spp. spores were found in the musculature of *Osteochilus vittatus* and *O. hasselti*. In *O. hasselti*, *Myxobolus* spores of 2 species were found in the kidney, while in *Barbonymus schwanenfeldii*, plasmodia of *Myxobolus* spp. were found in the gills. Besides *Myxobolus* spp., the occurrence of 2 known *Henneplagia* spp. was demonstrated in the gills: *H. shaharini* in marble goby and *H. hemibagri* in Asian redtail catfish.

*Myxobolus terengganuensis* sp. nov.

**Type host:** Silver shark minnow *Osteochilus hasselti* (Valenciennes, 1842) (Cyprinidae). Local name: ikan terbol.

**Type locality:** Lake Tasik Kenyir, Malaysia.

**Site of tissue development:** Muscle. Mature disporoblastic plasmodia located intracellularly in muscle cells were elongated, up to 500 to 700 µm in length. Only old plasmodia filled with spores were found.

**Type material:** Digitized photos of syntype spores and histological sections were deposited in the parasitological collection of the Zoological Department, Hungarian Natural History Museum, Budapest, coll. no. HNHM-70080. The 18S rDNA sequence of *Myxobolus terengganuensis* sp. nov. was deposited in GenBank under accession no. EU643629.

**Prevalence of infection:** 1/10 of 10 to 22 cm sized fish.

**Etymology:** The species was named after Terengganu State, Malaysia.

**Spores:** The spores (Figs. 1a & 2a) were ellipsoidal in frontal view. In sutural view they had an elongated-ellipsoidal shape (Fig. 1b). Length of the spores was 12.7 ± 0.51 (range, 12 to 13.4) µm (n = 50), width 7.4 ± 0.53 (range, 6.7 to 8.3) µm (n = 50), thickness 5 ± 0.21 (range, 4.8 to 5.2) µm (n = 15). Polar capsules were elongated, different in size, slightly converging anteri-
orly. Larger capsule was $6.8 \pm 0.40$ (range, 6.2 to 7.3) µm long ($n = 50$) and $3.2 \pm 0.14$ (range, 2.9 to 3.4) µm wide ($n = 50$). Smaller capsule was $5 \pm 0.42$ (range, 4.4 to 5.5) µm long ($n = 50$) and $2.3 \pm 0.13$ (range, 2.2 to 2.6) µm wide. Filament coils could not be observed in the capsules. No intercapsular appendix was seen. The width of the suture was $5.4 \pm 0.40$ (range, 5 to 6) µm. Sutural edge markings were not seen. A single binucleated sporoplasm with round iodinophilous vacuole was asynchronous in shape. Mucous envelope was not found.

**Molecular data:** The 18S rDNA of *Myxobolus terengganuensis* sp. nov. was related to that of another muscle-infecting species described from European habitats (see Figs. 10 & 11). Based on the nucleotide sequences, *M. pseudodispar* (GenBank accession no. AF380145) appears to be its closest relative with 95.4% similarity. A 94.9% homology with *M. musculi* and a 94.1% homology with *M. cyprini* were observed.

**Histology:** Spores were found in elongated intracellular plasmodia inside the muscle cells. In more progressed cases of plasmodial development, the infected muscle cells became disrupted and masses of spores were located among the neighboring muscle cells surrounded by histiocytes and melanin-containing macrophages (Fig. 3). Very often spores were associated with metacercariae of a closely non-identified trematode species infecting the musculature of this fish (Fig. 4a). *Myxobolus* spores were often found inside metacercarial cysts (Fig. 4b). In some cases the cysts formed by closely non-identified metacercariae could no longer be recognized, but among the muscle cells nodules containing mature spores infiltrated and surrounded by melano-macrophage cells were observed (Fig. 4c).

**Remarks:** By its elongated spores of varying size and by its intracellular location, this species most closely resembles the muscle-dwelling species of the *Myxobolus cyprini* group (*M. musculi, M. pseudodispar*) which is supported by the relatively high similarity of the 18S rDNA sequences. However, the members of the *M. cyprini* group infect genetically different hosts.

**Myxobolus tasikkenyirensis** sp. nov.

**Type host:** Bonylip barb *Osteochilus vittatus* (Valenciennes, 1842) (Cyprinidae). Local name: ikan rong. **Type locality:** Lake Tasik Kenyir, Malaysia. **Site of tissue development:** Plasmodia and spores of this species were found in the muscles. **Type material:** Digitized photos of syntype spores and histological sections were deposited in the parasitological collection of the Zoological Department, Hungarian Natural History Museum, Budapest, coll. no. HNHM-70081. The 18S rDNA sequence of *Myxobolus tasikkenyirensis* sp. nov. was deposited in GenBank under accession no. EU643626.

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**Fig. 1.** Schematic drawings of spores of *Myxobolus terengganuensis* sp. nov., (a) frontal and (b) sutural view; *M. tasikkenyirensis* sp. nov., (c) frontal and (d) sutural view; *M. csabai* sp. nov., (e) frontal and (f) sutural view; *M. osteochili* sp. nov., (g) frontal and (h) sutural view; *M. dykovae* sp. nov., (i) frontal and (j) sutural view; *Henneguya shaharini*, (k) frontal view, (l) sutural view, (m) with complete caudal extensions; *H. hemibagri*, (n) frontal and (o) sutural view. Scale bars = 10 µm.
Prevalence of infection: 2/4 of 16 to 18 cm sized fish.

Etymology: The species was named after the habitat of the host fish.

Spores: The spores (Figs. 1c & 2b) were oval in frontal view, tapering at the anterior poles. In sutural view they were elongated ovals, tapering at both ends (Fig. 1d). Length of the spores was 12.8 ± 0.72 (range, 11.8 to 13.8) µm (n = 50), width 9.2 ± 0.55 (range, 8.3 to 9.9) µm (n = 50), thickness 6 (range, 5.5 to 6.5) µm (n = 3). Polar capsules were elongated, equal in size, slightly converging anteriorly. Capsules were 6.4 ± 0.42 (range, 5.9 to 7.2) µm long (n = 50) and 3 ± 0.25 (range, 2.7 to 3.4) µm wide (n = 50). Five to six loosely wound filament coils were arranged perpendicularly to the capsule length in the polar capsule. Intercapsular appendix was absent. The width of the suture was about 5.5 µm. Sutural edge markings were not seen. A single binucleated sporoplasm with round iodinophilous vacuole was present. Mucous envelope was not found.

Histology: Cysts of larval trematodes and plasmodia were observed in close association in the musculature of the fish. Metacercariae seemed to be readily encysted in or close to plasmodia (Fig. 5).

Molecular data: Myxobolus tasikkenyirensis sp. nov. was on the same branch on the phylogenetic tree as M. csabai sp. nov. described in the next section (see Figs. 10 & 11). The similarity between their sequences was 96.5%. The closest species available in GenBank were M. pellicides (92.5%) and M. pendula (91.9%).

Remarks: The spores lacking intercapsular appendix resemble many muscle-dwelling Myxobolus spp. but differ from the latter by their elongated, anteriorly converging polar capsules and by their DNA sequences. In its affinity to metacercarians, this species resembles M. uvuliferis infecting the pumpkinseed Lepomis gibbosus in North America, but differs from the latter species by its more elongated shape and by their genetically different hosts.

Myxobolus csabai sp. nov.

Type host: Silver shark minnow Osteochilus hasselti (Valenciennes, 1842) (Cyprinidae). Local name: ikan terbol.

Type locality: Lake Tasik Kenyir, Malaysia.

Site of tissue development: Round-shaped plasmodia with a diameter of 160 (range, 150 to 180) µm were found in the renal interstitium (Fig. 6). Aged spores were located in the melanomacrophage centers of the kidney.

Type material: Digitized photos of syntype spores and plasmodia were deposited in the parasitological collection of the Zoological Department, Hungarian Natural History Museum, Budapest, coll. no. HNHM-70082. The 18S rDNA sequence of Myxobolus csabai sp. nov. was deposited in GenBank under accession no. EU643628.

Prevalence of infection: 5/10 of 10 to 22 cm sized fish.
**Etymology:** The species is named after György Csaba, the well-known Hungarian fish pathologist.

**Spores:** The spores (Figs. 1e & 2c) were oval in frontal view, tapering at the anterior poles. In sutural view they were spindle-shaped (Fig. 1f). Length of the spores was $12.9 \pm 0.52$ (range, 12.1 to 14) µm ($n = 50$), width $8.6 \pm 0.49$ (range, 8.2 to 9.6) µm ($n = 50$), thickness $5 \pm 0.44$ (range, 4.7 to 5.6) µm ($n = 15$). Polar capsules were pyriform, different in size, slightly converging anteriorly. Larger capsule was $6.1 \pm 0.29$ (range, 6.4 to 6.8) µm long ($n = 50$) and $3.8 \pm 0.14$ (range, 3.7 to 4.0) µm wide ($n = 50$). Smaller capsule was $3.6 \pm 0.54$ (range, 3.3 to 4.8) µm long ($n = 50$) and $2.3 \pm 0.36$ (range, 2.0 to 2.9) µm wide. Six densely wound filament coils were arranged obliquely to the capsule length in the polar capsule. A large intercapsular appendix located asymmetrically was present. The width of the suture was $6.6 \pm 0.19$ (range, 6.4 to 6.8) µm. Sutural edge markings were not seen. A single binucleated sporoplasm with round iodinophilous vacuole was present. Mucous envelope was not found.

**Molecular data.** *Myxobolus csabai* sp. nov. and *M. tasikkenyirensis* sp. nov. were closely related, with 96.5% similarity. Likewise, *M. pellicides* and *M. pen-
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*dula* showed relatively high correspondence in their 18S rDNA with 91 to 92% homology.

**Histology:** No plasmodia were found in histological sections.

**Remarks:** By its large, asymmetrically located intercapsular appendix and by its DNA sequences this species differs from all known *Myxobolus* spp.

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**Myxobolus osteochili** sp. nov.

**Type host:** Silver shark minnow *Osteochilus hasselti* (Valenciennes, 1842) (Cyprinidae). Local name: ikan terbol.

**Type locality:** Lake Tasik Kenyir, Malaysia.

**Site of tissue development:** Round plasmodia

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Figs. 6 to 9. Fig. 6. *Myxobolus csabai* sp. nov. plasmodium (p) in the interstitial part of the kidney (k). Some spores (s) were released from the compressed plasmodium. Fresh mounts. Scale bar = 25 µm. Fig. 7. Kidney of *Osteochilus hasselti* infected with groups of *M. osteochili* sp. nov. plasmodia (p). Plasmodia are located in the fibrous capsule (arrow) covering the parenchymal part of the kidney (k). Inset: enlarged part of the plasmodium with spores. Histological section with H&E staining. Scale bar = 100 µm for the picture and 10 µm for the inset. Fig. 8. A young *M. dykovae* sp. nov. plasmodium (p) in the gills of *Barbonymus schwanenfeldii*. The plasmodium is located inside a gill lamella. Around the plasmodium, serum fills the space inside the lamella. The tip of the lamella is undamaged. Histological section with H&E staining. Scale bar = 50 µm. Fig. 9. A mature *M. dykovae* sp. nov. plasmodium (p) filled with spores (inset) in the gills of *B. schwanenfeldii*. Histological section with H&E staining. Scale bar = 50 µm for the picture and 10 µm for the inset.
arranged in large groups in the fibrous capsule of the kidney.

**Type material:** Spores in glycerin-gelatin and histological sections were deposited in the parasitological collection of the Zoological Department, Hungarian Natural History Museum, Budapest, coll. no. HNHM-70083. Sequence data for this species are being further studied by Szilvia Marton (Veterinary Medical Research Institute, Hungarian Academy of Sciences), and have not yet been submitted to GenBank.

**Prevalence of infection:** 6/13 of 10 to 22 cm sized fish.

**Etymology:** The species was named after the generic name of the fish host.

**Spores:** The spores (Figs. 1g & 2d) were oval in frontal view, tapering at the anterior poles. In sutural view they were lemon shaped (Fig. 1h). Length of the spores was 13 ± 0.63 (range, 12 to 14) µm (n = 50), width 8.7 ± 0.45 (range, 8 to 9) µm (n = 50), thickness 5 ± 0.45 (range, 4.5 to 5.5) µm (n = 15). Polar capsules were elongated, similar in size, 6.6 ± 0.52 (range, 6 to 7) µm long (n = 50) and 2.4 ± 0.42 (range, 2 to 3) µm wide (n = 50). Eight to 9 densely wound filament coils were arranged perpendicularly to the capsule length in the polar capsule. In sutural view the thickness of the suture rim measured about 0.5 µm. The sutural rim protruded over the surface of the spore 1 to 1.2 µm at the anterior pole, and 0.5 to 0.7 µm at the posterior pole. No intercapsular appendix was found. Sutural edge markings were not seen. A single binucleated sporoplasm with round iodinophilous vacuole was present. Mucous envelope was not found.

**Molecular data:** Comparison of Myxobolus osteochili in a pairwise alignment in BioEdit showed no close similarity with any of the Malaysian myxospores described here.

**Histology:** Plasmodia of this species were found in the fibrous capsule covering the kidney. In the infected portions of this capsule, large nodules were formed by groups of round-shaped plasmodia (Fig. 7). Free spores were also found in the melanomacrophage centers.

**Remarks:** Myxobolus osteochili corresponds to M. csabai in size and shape but differs from the latter species by its equal and elongated polar capsules and by the lack of an intercapsular appendix.

**Myxobolus dykovae** sp. nov.

**Type host:** Tinfoil barb *Barbonymus schwansenfeldii* (Bleeker, 1853) syn. *Puntius schwansenfeldii* Bleeker, 1853 (Cyprinidae). Local name: lampam sungai.

**Type locality:** Lake Tasik Kenyir, Malaysia.

*Henneguya shaharini* Shariff, 1982

**Host:** Marble goby, *Oxyeleotris marmoratus* (Bleeker). Local name: ketutu.

**Locality:** Lake Tasik Kenyir, Malaysia.
**Site of tissue development:** Gills.

**Material:** Digitized photos of syntype spores and histological sections were deposited in the parasitological collection of the Zoological Department, Hungarian Natural History Museum, Budapest, coll. no. HNHM-70085. The 18S rDNA sequence of *Henneguya shaharini* was deposited in GenBank under accession no. EU643630.

**Prevalence of infection:** 1/2 of 21 to 26 cm sized fish.

**Molecular data:** The 1033 bp long 18S rDNA fragment of *Henneguya shaharini* was very similar to those from other *Henneguya* spp. such as *Henneguya lesteri*, *Henneguya pagri*, *Henneguya lateolabracis*, *Henneguya doori*, and *Henneguya gurlei*, although even the highest similarity value did not exceed 75%. The 18S rDNA nucleotide sequence of *Hennegoides mekongiensis* is available in GenBank and is 85.2% similar to *H. shakarini*; however, it is only 435 bp long. *Henneguya mekongiensis* could not be placed on the phylogenetic tree (Figs. 10 & 11) because its extremely short sequence negatively affected the alignment and the construction of the tree.

**Histology:** Plasmodia with mature spores were found intralamellarly.

**Remarks:** *Henneguya shaharini* seems to be common on marble goby, as in addition to the present survey, similar infections were found also during a previous survey in 2004 from River Terengganu (K. Molnár & Cs. Székely unpubl.). Round or ellipsoidal small plasmodia 100–120 × 80–100 µm in size developed intralamellarly in the gills. Spores found in both surveys corresponded to spores described by Shariff (1982).

**Henneguya hemibagri** Tchang et Ma, 1993

**Host:** Asian river (redtail) catfish *Hemibagrus nemurus* (Valenciennes). Local name: baung.

**Location:** Lake Tasik Kenyir, Malaysia.

**Site of infection:** Plasmodia developed intralamellarly in the gills.

**Prevalence of infection:** 2/2 of 23 to 27 cm sized fish.

**Type material:** Spores in glycerol-gelatine were deposited in the parasite collection of the Hungarian National History Museum, coll. no. HNHM-70086.

**Remarks:** In morphology and size the spores found during the present survey correspond to the spores observed by Molnár et al. (2006b) from another locality (net-cage farms on Terengganu River).

**DISCUSSION**

Due to the great similarity of *Myxobolus* spp. spores, the identification of different species exclusively on morphological basis is rather difficult. Considering host specificity and location of vegetative developmental stages in organs and tissues is a great help in differentiating morphologically identical or similar spores, but a final answer to these questions can only be obtained by studying spores with molecular biological methods.

Little is known about the host specificity of myxosporeans, but earlier studies (Molnár 1994) suggest that some genera of myxosporeans have relatively

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**Fig. 10.** Phylogenetic tree generated by neighbor-joining analyses of the 18S rDNA sequences of myxosporeans, rooted at *Ceratomyxa shasta*. Numbers at nodes indicate bootstrap confidence values (1000 replications). GenBank accession numbers are given in parentheses. Myxosporeans examined in the present study are in bold. *M.: Myxobolus; H.: Henneguya; T.: Thelohanellus*
strict host specificity and a given myxosporean species has only 1 or a few closely related fish hosts. The existence of relatively strong host specificity in the Myxobolus genus has also been supported by molecular biological studies. Eszterbauer & Székely (2004) reported that DNA sequences can clearly differentiate between morphologically similar species occurring in cyprinid fishes. In a similar way, clear molecular differences were found among Myxobolus spp. collected during the present study. Accepting this general rule, most of the endemic Malaysian species might have specific Myxobolus spp. which should be described as new ones. Myxosporean species common in other biotopes of the world can occur in southeast Asia, mostly in introduced fishes such as common carp, silver carp, goldfish or Tilapia and Batrachus spp.

The organ and tissue specificity of these myxosporeans also seems to be common (Molnár et al. 2002). Two of the Myxobolus spp. studied here (M. terengganuensis sp. nov. and M. tasikkenyirensis sp. nov.) had definite tissue tropism towards the muscle cells, while in the case of M. csabai sp. nov. and M. osteochilii sp. nov., the specific location of plasmodium stages was the kidney. M. dykovae sp. nov. formed cysts in the gills of its host, and its plasmodia developed in the capillary network of the secondary lamellae in a vascular intralamelar position (see Molnár 2002). Both species found in the muscles had an intracellular route of development. Although M. terengganuensis and M. tasikkenyirensis are typical intracellular parasites of the muscle cells, the common occurrence of their spores inside the cysts of larval trematodes needs further explanation. The affinity of myxosporean spores to larval trematodes is already known; Cone & Anderson (1977) described Uvulifer from the cysts of Uvulifer ambloplitis metacercariae infecting the pumpkinseed Lepomis gibbosus in Canada. Those authors surmised that the cercariae invaded portions of the muscle which already contained myxosporeans and that the capsule might have originally been a myxosporean cyst. Our findings support this theory and our pictures (Figs. 3 & 4) show that the metacercarial stages are actually formed inside a M. tasikkenyirensis cyst. Cone & Anderson (1977) found both spores and young plasmodia inside the metacercarian cyst. In our case, in M. terengganuensis and M. tasikkenyirensis infections, only disseminated spores inside trematode cysts and cysts with mature spores associated to metacercarians were found. We suggest that metacercariae invading the muscles of fish prefer tissue portions where plasmodia disintegrate their host cells and spores are released into the intercellular space. Obviously most of the released spores leave this location due to compression of the musculature but a certain number of spores remain on the spot and become surrounded by granulation tissue, as was found in the case of M. terengganuensis sp. nov.

Some of the fish species of Tasik Kenyir studied by us are commercially important fishes regularly seined or to be cultured in fish farms or cage systems in the future, so their myxosporean infection could become an economic problem. Among the fishes examined during the present survey, we found moderately infected and severely infected specimens. During the short term of the examinations, no disease symptoms were recorded and at present we cannot give data on
the pathogenicity of the myxozoan infections of Tasik Kenyir fishes. The large number of myxozoan species, the intensive infection of some fish specimens and the local changes around the parasites, however, suggest the possibility that in intensive culture systems, more severe symptoms and losses might also appear.

Of the species studied in the present survey, neither Myxobolus nor Henneguya spp. showed strict relationships in their DNA sequence to species in GenBank, because the sequences of myxobolid species deposited in Genbank are mostly from Europe, North America or China. Nevertheless some interesting observations can be made. M. terrangauensis was on the phylogenetic tree next to the M. cyprini group, which it resembles morphologically. M. tasikkenyirensis is also a muscle-infecting parasite, but its phylogenetic position is far away from M. terrangauensis. Its closest relative is M. csabai, which develops in the host’s renal interstitium. There is only a low similarity between M. dykovaevae and the European M. dujardini, M. obesus and M. hungaricus, but these species infect gills, as does M. dyakovaevae. Henneguya sharini does not have any close relative among the sequenced species.

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