Morphological and molecular studies on two myxosporean infections of cyprinid fishes: *Thelohanellus pyriformis* from tench and *Thelohanellus cf. fuhrmanni* from nase

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**ABSTRACT**

During surveys on myxosporean parasites of Lake Balaton and River Danube fishes, two *Thelohanellus* spp. were found on tench (*Tinca tinca*) and on common nase (*Chondrostoma nasus*). They were identified as *Thelohanellus pyriformis* and *Thelohanellus cf. fuhrmanni*, respectively. Myxospores of *T. pyriformis* from tench were collected from artery branchialis afferens of gill filaments. The mature myxospores of this species were pyriform in shape and 19 ± 0.6 (18–19.5) long, 8.2 ± 0.54 (7.5–9) wide, 7.3 ± 0.25 (7–7.5) thick containing polar tubules with 9–10 turns. The plasmodia of *Thelohanellus cf. fuhrmanni* were collected from under the skin of snout of the common nase. The myxospores were pyriform, 16.3 ± 0.39 (15.5–16.5) long, 6.5 ± 0.55 (6.3–7) wide, 6.3 ± 0.53 (5.8–7) thick containing polar tubules with 6 turns. Small subunit ribosomal DNA sequences of both *Thelohanellus* species differed from other known myxozoans. The myxospores morphology, histopathology and ssrDNA sequences supported a diagnosis of *T. pyriformis* from tench and *T. cf. fuhrmanni* from common nase.

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**1. Introduction**

Members of the genus *Thelohanellus* Kudo, 1933 are among the most common myxozoans usually infecting freshwater fishes. Zhang et al. (2013) in their synopsis reported on 108 nominal species, but this number has been increased recently with several newly described species. Most of the described *Thelohanellus* species infect cyprinid fishes but majority of species are originated from India and China (Basu et al., 2006; Zhang et al., 2013; Chen and Ma 1998). After a general revision, most probably the large number of these species will prove to be synonyms. Similar problems concern the host range and tissue specificity of members of this genus. Often several fish species are recorded as hosts for a given *Thelohanellus* sp. (Kudo, 1919; Shulman, 1966; Donec and Shulman, 1984), while others argue that *Thelohanellus* spp. are rather host specific parasites infecting mostly a single species (Akhmerov, 1955, 1960; Molnár, 1982; Shin et al., 2014). Moreover, recent studies prove that most species can be characterized by strict tissue and organ specificity (Akhmerov, 1955; Molnár and Eszterbauer, 2015; Liu et al., 2016; Zhang et al., 2017). The synopsis on *Thelohanellus* by Zhang et al. (2013) focused to designate only the host and organ from where the *Thelohanellus* species were first described. The *Thelohanellus* fauna of the common carp and the goldfish has been thoroughly studied, however little is known on species infecting other cyprinids. Donec and Shulman (1984) recorded *Thelohanellus* spp. in more than a dozen cyprinids fishes but they classified them to some already known species such as *T. pyriformis* (Thelohan, 1892), *T. fuhrmanni* (Auerbach, 1909), or *T. ocullilaeiscii* (Trojan, 1909). Type host of *T. pyriformis* is the tench while the type host of *T. fuhrmanni* is the roach (*Rutilus rutilus* L.). Little is known on the *Thelohanellus* infection of the nase. Only a single report was presented up to this time on its occurrence on this fish by Molnár (1979). More intensive studies were conducted by Dyková and Lom (1987) who studied the histology and the morphology of a case of *T. pyriformis* infection in tench by electron microscope. Data are scarce on the ssrDNA sequences of the majority of *Thelohanellus* species, which may answer the validity of these morphologically similar species.

In the present study, we document *Thelohanellus pyriformis* in the type host, tench, and report a *Thelohanellus cf. fuhrmanni* infection found in nase. Morphological details and histopathological analysis of
Polymerase chain reaction primers used for the amplification and sequencing of the ssrDNA gene.

| Primers | Sequences | References |
|---------|-----------|------------|
| ERBB1   | ACCCTGGTGATCCTGCCA | Barta et al. (1997) |
| ERBB1O  | CTTCCGAGGTCCTACACTGGG | Barta et al. (1997) |
| CR1R    | CTAGGGCTGGTATCTGAGCTTCG | Szekely et al. (2015) |
| CR1F    | CGGAGAGCTAGATAAGGCTTAG | Szekely et al. (2015) |
| MXGENF4F | GTCCCTAGAATACATCGAG | Diamant et al. (2004) |
| ACR1F   | TGGGTAATTTGCCGCTTGAGC | Hallett and Diamant (2001) |
| ACR1R   | ATTGGTGTTCATG | Rocha et al. (2014) |

these species are supported by ssrDNA sequences.

2. Materials and methods

During a long-term fish health survey of natural water in Lake Balaton and River Danube, Hungary, fishes were regularly examined for myxosporean infections. Large sized fish specimens were obtained from fishing companies and sport anglers, and others were selected from the trap of Lake Balaton Drainage System. Small fishes were collected by a 15 m long seine net. All caught fishes were carried to the institute alive and then some of the myxospores were placed in oxygenated water, kept in aquarium in aerated water, and dissected within 3–5 days after collection. Tench (Tinca tinca) is a widespread but rare fish in both Lake Balaton and River Danube, only seven specimens (8–31 cm in length) were examined in 2019. Common nase (Chondrostoma nasus) is one of the most common fish in River Danube. In the case of common nase, parasitological dissections were conducted on 6 specimens of 31 cm in length caught in Lake Balaton at Balatonszemes (46°17′35″N 17°45′55.9″E). When inspecting hemibranchia under dissecting microscope for myxosporeans. The intestines were opened and pieces of muscle were compressed between two glass plates. In the case of a single cyst, small portion of myxospores was absorbed by a syringe for morphological and molecular studies and the cyst and the surrounding tissues were saved for histology. Myxospores from the cysts were studied native, and then some of the myxospores were placed in glycerine-jelly under a coverslip on a slide and preserved as a reference preparation. Myxospores were also collected into Eppendorf tubes in 95% ethanol for subsequent molecular studies. Tissue samples from infected organs had been fixed in Bouin’s solution, embedded in paraffin wax, sectioned at 4 μm and stained with hematoxylin and eosin. The vitality of the spores was checked by 0.4% solution of urea; spores of a given plasmodium were regarded as mature when at least 90% of them extruded polar filaments in that solution. Unfixed spores were studied using differential interference contrast optics on an Olympus BH2 microscope. Fresh spores were photographed with an Olympus DP20 digital camera or recorded on videotapes; digitized images were obtained and measurements were taken from fresh spores and digitized photos. All measurements are given in micrometers (μm). In description, we use the term “polar tubule” instead of “polar filament” as Ben-David et al. (2016) suggested it.

3. Genomic DNA isolation and amplification

Preserved isolated plasmodia or myxospores in 95% ethanol were centrifuged at 8000 × g for 10 min then ethanol was removed by pipetting and evaporation. Genomic DNA was extracted from the pellet using the Genaid Tissue Genomic DNA Mini kit (Genaid Tawai City, Taiwan), following the manufacturer’s recommended protocol for animal tissue. The ssrDNA was amplified by semi-nested PCR: first round with universal primers ERBB1 and ERBB1O (Barta et al., 1997). This was followed by two semi-nested PCR reactions to generate over-lapping sequences, ERBB1 with CR1R (Szekely et al., 2015) and ERBB1O with CR1F (Szekely et al., 2015) were paired. Reaction mixtures and thermal conditions followed the protocol previously published by Goswami et al. (2021). The primers used for PCR amplification and sequencing are listed in Table 1. PCR products were analysed in 1% agarose gel and the specific bands were excised from the gel.

PCR products cut off from the gel were purified with the Gel/PCR DNA-Fragments Extraction Kit (Geneaid Tawai City, Taiwan) and directly sequenced with sequencing primers (Table 1) in both directions using the BigDye Terminator v3.1 Cycle Sequencing Kit (Life Technologies, Waltham, MA USA) with an ABI PRISM 3100 Genetic Analyser (Life Technologies, Waltham, MA USA), using the amplification and inner primers.

4. Phylogenetic analysis

Assembly of the sequences was conducted in MEGA X (Kumar et al., 2018) and were corrected by manual adjustments. Assembled ssrDNA sequences were verified as myxozoan by GenBank BLAST search and related myxozoan sequences were added to the alignment. The selected sequences were restricted to only those of at least 1000 base pairs in length and 89% similarity. Gblocks 0.91b (Castresana, 2000) was used to remove ambiguous regions and highly variable sections. The sequences were aligned by using Clustal W (Thompson et al., 1994) implemented in the MEGA X. Pairwise distances were computed by using the p-distance model matrix in MEGA X. Evolutionary model for Maximum Likelihood analysis was determined by model testing of the dataset for the nucleotide substitution model of judging by the Akaike information criterion (AIC), which suggested the GTR + G + I 1000 bootstrap replicates were performed to assess node support. Chloromyxum cristatum (AY604198) was chosen as an out-group in the final alignment.

5. Results

5.1. Infection in tench (Tinca tinca)

Thelohanellus infection in tench was found in June 2019 in a fish of 29 cm in length caught in Lake Balaton at Balatonszemes (46°48′36.4″N 17°45′55.9″E). When inspecting hemibranchia under dissecting microscope, small dark nodules were found in the arteria afferens, which proved to be plasmodia filled with spores of a Thelohanellus sp. When studying the Gill filament under a compound microscope round or oval plasmodia were located in the lumen of the arteria afferens (Fig. 1). Round cysts measured 25 × 25 and elongate ones 25 × 160. They contained from 500 to 3000 spores. Each of the eight arches was infected with 2–8 plasmodia. Based on the shape and size of the spores and the specific location, this species was identified as T. pyriformis (Thelohan, 1892).

5.2. Description of spores

Mature spores (Fig. 2a and b and 2c) were pyriform both in frontal and sutureal view, slightly tapering toward the anterior end and round at posterior end, 19.0 ± 0.61 (18.0–19.5) long (n = 50), 8.2 ± 0.54 (7.5–9.0) wide (n = 50) and 7.3 ± 0.25 (7.0–7.5) thick (n = 25). In some spores the anterior end slightly bent. Spore wall formed by two shell valves of equal-size separated by sutural ridge, and light microscopically showing an about 0.55 in thickness. The wall at the posterior pole of the spore thickened showing some nodules on its surface, regarded as cultural notches by Dyková and Lom (1987). Single pyriform polar capsule presents close to apex of spore, 8.4 ± 0.055 (7.5–9.0) (n = 50) long, 4.6 ± 0.3 (4.0–5.0) (n = 50) wide. Polar tubules closely coiled with 9, or 10 turns, arranged perpendicular to the longitudinal axis of capsule. At one side of the polar capsule close to the blunt end a bright nucleus of the
capsulogenic cell, about 1 in diameter is present. Sporoplasm located at the posterior pole of spore, contains two round nuclei. No iodinophilous vacuole had been seen. At the anterior end of the spore besides the polar capsule two bright round globules (probably the nuclei of the valvogenic cells), about 1 in diameter are located. Some of the spores are surrounded by an average 15.8 ± 22.7 mucous envelope (Fig. 2c arrows).

5.2.1. Taxonomic summary
Host: Tench, Tinca tinca L.
Locality: Lake Balaton, Balatonszemes, Hungary (46°48’36.4”N 17°45’55.9”E).
Site of tissue development: Arteria branchialis afferens.
Material: Photo-types and histological preparations were deposited in the parasitological collection of the Zoological Department, Hungarian Natural History Museum, Budapest, Coll. No. HNHM-PAR-72082. The ssrDNA sequence (1973 bp, long) of *T. pyriformis* was deposited in the GenBank under accession numbers OM764632.

Prevalence: 1 specimen from 7 fish.
Molecular data: The phylogenetic analysis revealed the closest relative of this species to *T. cf. fuhrmanni* (present study data) with 96.4% nucleotide sequence identity. 95.1% similarity was found with *T. magnacysta* (MN540268) and 91.6% similarity was observed to a *Thelohanellus* species-2018 (MK053786) from intestinal epithelium of the Cobitis paludica in Portugal.

5.3. Infection in common nase (Chondrostoma nasus)
During our long-term examinations on myxosporan infections of the common nase, *Thelohanellus* infection was detected only in two cases. Firstly, in 2004 in a 34.4 cm long nase collected in the Danube close to the city Győr (46°48’36.4”N 17°45’55.9”E). Secondly, in 2019 when *Thelohanellus* infection was found in one of the six nase of 31–33 cm long caught from the Danube at Nagymaros (47° 47’ 16.9368” N 18° 57’ 14.9256” E). In both cases, a single large, flat, 4 × 4 mm size cyst was found on the snout of the specimens. The cysts were visible to the naked eye. Inside the cysts, 5–6 global shape plasmodia were located filled by about 10,000 myxosporas.

5.4. Description of spores
Mature myxosporas were pyriform both in frontal and sutural view, slightly tapering toward the conical anterior end and rounded at the posterior end 16.3 ± 0.39 (15.5–16.5) long (n = 50), 6.5 ± 0.55 (6.3–7.0) wide (n = 50) and 6.3 ± 0.53 (5.8–7.0) thick (n = 25) (Fig. 3a, b, c, d). Myxospore wall formed by two shell valves of equal-size separated by a thick and fairly well marked sutural ridge. Light microscopically in sutural plane, the spore wall was about 0.5–0.6 in thickness but at the posterior end the myxospore thickened and bearing some nodules on its surface. Single flask-shaped polar capsule presents close to apex of spore, 6.5 ± 0.8 (5.5–7.0) long, 3.0 ± 0.1 (3.0–3.2) thick. Polar tubule coiled with 6 turns arranged obliquely to the longitudinal axis of capsule. At one side close to the blunted end of the polar capsule a bright nucleus of the capsulogenic cell, about 1 in diameter is present. Sporoplasm located close to posterior pole of myxospore, containing two round nuclei. At the anterior end of the myxospore besides the thickened polar capsule one or two bright round globules (probably the nuclei of the valvogenic cells) are located. Sutural ridge thickened and fairly well marked. Most of the myxosporas surrounded by a pale, oval shape on average 13.5 × 9.0 mucous envelope (Fig. 3 c, d arrows) attaching to the wall of the myxospore at the anterior pole. The spores of the species resembled to *T. fuhrmanni* described by Auerbach (1909).

5.5. Histology
Plasmodia of this species located inside the loose connective tissue of the snout. Five to six round shape plasmodia with a diameter of 1–1.3 mm developed in close vicinity within an about 4.3–4.5 mm long and 3–3.2 mm wide cyst formed by the host (Fig. 4a). The thickness of the cyst was not measured. Plasmodia were surrounded by a relatively thin dense connective tissue wall of the host (Fig. 4). This layer wrapped plasmodia from outside and separated plasmodia from each other also in the interplasmodial space. Plasmodia were separated from the host capsule by their own very thin eosinophil staining ectoplasm (Fig. 4b and c). Dark staining early developmental stages located in a single layer at the periphery (Fig. 4c). Centrally, it was followed by a layer of sporoblasts and forming myxosporas (Fig. 4d), while the centre of the plasmodium was filled by young and matured myxosporas.

Remarks: Nodules at the thickened wall at the posterior end found also at *T. pyriformis* are more emphasized in this species.

5.5.1. Taxonomic summary
Host: Common nase, Chondrostoma nasus L.
Locality: River Danube, Nagymaros, Hungary (47° 47’ 16.9368” N 18° 57’ 14.9256” E).
Site of tissue development: Plasmodium under the skin of the snout.
Material: Photo-types were deposited in the parasitological collection of the Zoological Department, Hungarian Natural History Museum, Budapest, Coll. No. HNHM-PAR-72083. The ssrDNA sequence (1547 bp, long) was deposited in the GenBank under accession number OM793063.

Prevalence: 2 specimens of 6 nases of 3–4 years old.
Molecular data: This species showed 97.6% similarity with *Thelohanellus magnacysta* (MN540267.1) described from the somatic muscles of *Cyprinella venusta* Girard by Ksepka et al. (2020), and 96.4% similarity was found to *Thelohanellus pyriformis* sequenced by us in this study from the gill of Tinca tinca. 93.4% similarity was observed with a *Thelohanellus* sp. (SR-2018 (MK053786), obtained from the intestinal epithelium of Cobitis paludica (Fig. 5)). This clade of Thelohanellus species was branched together with several Myxobolus spp. from cyprinid fishes apart from other notable Thelohanellus species, like *T. nikolitsii* and *T. horvokai*.

6. Discussion
The genus *Thelohanellus* Kudo, 1933 is one of the richest myxosporan genera in species. Since then Zhang et al. (2013) in their synopsis reported on 108 *Thelohanellus* species, at least 14 more *Thelohanellus*
spp. have been described from China, India, Brazil, Angola, Egypt and the USA. Most of them are plasmodia-forming parasites infecting various tissues of freshwater fishes, and some are pathogenic, causing economical losses in aquaculture (Zhang et al., 2013). Identification of several *Thelohanellus* spp. is difficult because descriptions of the old species are often relatively poor, lacking several important characteristics; moreover, many species were originally described as *Myxobolus* sp. with one polar capsule (Kudo, 1933).

Recently, publications from Asia (Yokoyama et al., 2012; Zhang et al., 2013; Singh and Kaur, 2012) described many new species or re-described old ones, but there is still a lot of questions regarding their host-, tissue-, and organ specificity. In our present study, *T. pyriformis* in the gills of tench was identified based on the correspondence of morphometrical characteristics of the spores, the location of the plasmodia

Fig. 2. Spores of *Thelohanellus pyriformis* a) Schematic drawing of a spore in frontal view, b) Schematic drawing of a spore in sutural view. c) Microscopic photos of spores.
and the identity of the type host. Morphometrically differing myxospores are classified to certain species described long ago despite their evolutionary distinct hosts, like *T. pyriformis* (Thelohan, 1892), *T. oculileucisci* (Trojan, 1909) or *T. fuhrmanni* (Auerbach, 1909). Many of the host fishes are probably infected by currently undescribed *Thelohanellus* species which are assumed to be one of those above-mentioned species. Akhmerov (1955, 1960) realized first the large morphological diversity and species richness of *Thelohanellus* spp. in common carp and in other cyprinids, though Shulman (1966) furtherly regarded most of his species’ synonyms. Therefore, the redescriptions of *T. pyriformis* and *T. fuhrmanni* are necessary. The myxospores of *T. pyriformis* were found in the same species as in the original description and spores corresponded to the original data. However, the myxospores of *T. cf. fuhrmanni* were found in common nase (*Chondrostoma nasus*) altering from the type host (*Rutilus rutilus*) in many respects but belonging to the same subfamily Leuciscinae. Regarding *T. pyriformis*, it was originally reported from the gills, kidneys, and spleen of tench. Later, however several new hosts were reported, like Molnár (1979) described this parasite from the gills and skin of *T. tinca* and *C. nasus* from Hungary, however the myxospores found in the skin probably corresponded to *T. fuhrmanni*. Donec and Shulman (1984) reported *T. pyriformis* from several additional fish hosts and from various organs. Dyková and Lom (1987) described plasmodia of *T. pyriformis* from the vessels of gills of the type host tench and concluded that the infection could lead to hypertrophy of endothelial cells in gills. Large plasmodia of *T. cf. fuhrmanni* found together in a cyst caused well observable tumors in the nostrils of
the nase. Similar nodules with pathogenic characters were found in the skin of *Cyprinus carpio* infected by *T. kitauei* (Egusa and Nakajima, 1981; Zhai et al., 2016; Liu et al., 2019) and in *Cyprinella venusta* infected by *T. magnacysta* (Ksepka et al., 2020). Molnár (1994) suggested that host species and infection site are important factors for myxosporean identification. Host specificity of myxosporeans was mostly studied on *Myxobolus* spp. Several *Myxobolus* species are known to have a strict host specificity causing infection of a single host or some closely related species (Molnár et al., 2011; Cech et al., 2012). In our assumption, host specificity of *Thelohanellus* spp. might be similar to *Myxobolus* spp. Tench as a single member of Tincinae, and nase as a member of Leuciscinae are relatively distinct inside the family Cyprinidae, therefore it is unlikely that *T. pyriformis* could infect other cyprinids than tench. Recent publications (Shin et al., 2014; Lewisch et al., 2015; Liu et al., 2014, 2019) using molecular data suggest the relatively strict host specificity on *Thelohanellus* spp. of the common carp and goldfish (*Carassius auratus* L.). The common carp and its closest relative, the goldfish e.g., have no documented *Thelohanellus* species infecting the both fish species (Molnár and Kovács-Gayer, 1982; Zhang et al., 2013, 2018; Shin et al., 2014). Therefore, the problem of the occurrence of *T. cf. fuhrmani* in common nase requires more caution. The species from nase identified as *T. cf. fuhrmani* has similar looking but somewhat different sized spores compared to the original description. Interestingly, the site of infection in the roach documented by Auerbach (1909) as “connective tissue under the mucous membrane of the mouth” fairly corresponds to our data, therefore we designated this species as *T. cf. fuhrmani* until further evidence emerges. Both common nase and roach belong to the Leuciscinae subfamily of Cyprinidae but they differ greatly regarding their body structure and natural freshwater habitats. Phylogenetic studies have demonstrated that the origin and radiation of myxozoans reflect the evolution of their hosts (Carriero et al., 2013; Kodádková et al., 2015; Holzer et al., 2018; Patra et al., 2018). *Thelohanellus cf. fuhrmani* showed the close similarity with *T. magnacysta* (Ksepka et al., 2020) described from the muscles of *Cyprinella venusta* Girard, 1856 and 96.4% similarity was found to *Thelohanellus pyriformis* (present study) from the gill of *T. tinca*. However, there are no previously published sequences of *Thelohanellus* species from tench or common nase, therefore further evolutionary comparisons cannot be made beside the fact that these two newly sequenced species in this study showed a close relationship to each other. The genus *Thelohanellus* is polyphyletic, as its members are always scattered around within the *Myxobolus* cluster which is consistent to the previous studies (Liu et al., 2014; Székely et al., 2009). Both the species of *Thelohanellus* also showed closer similarity with *Myxobolus intimus* Zaika, 1965 isolated from the capillary network of gill lamellae of *Aspius aspius* L., as well with many *Myxobolus* species from cyprinid fishes just like the other *Thelohanellus* species. Molecular data clearly indicate a close genetic relation existing among genera *Myxobolus, Thelohanellus,* and *Henneguya*
Fig. 5. Phylogenetic tree generated by maximum likelihood analysis of ssrDNA sequences of *Thelohanellus pyriformis* and *Thelohanellus furhmanni* and other closely related myxozoan species identified by BLAST; GenBank accession numbers. Numbers at nodes indicate the bootstrap confidence values (ML). *Chloromyxum cristatum* was used as an outgroup.
(Zhang et al., 2019). The difference is in the number of polar capsules between *Thelohanellus* species (one polar capsule) and *Myxobolus* species (two polar capsules), however one of the two polar capsules in some *Myxobolus* species is too small to be recognized (Yokoyama and Ogawa, 2015; Griffin and Goodwin, 2011). Measurements given by different authors vary highly (Tables 2 and 3), therefore there is a possibility that they identified both species incorrectly, and parasites regarded as *T. pyriformis* or *T. fuhrmanni* designated by them represent different but closely related species. Further molecular data of new isolates might answer these uncertain species designations.

Unfortunately, our data rely on two cases, although we agree with Zhai et al. (2016) that measurements of spores should come from several different plasmodia to increase the reliability of the morphological data. Intraspécific morphometric variation of myxosporean species commonly occurs, therefore we are reluctant to describe *Thelohanellus* sp. found in nase as a new species. However, the great differences in morphology and size given for the two species (Table 2 and 3), and the large number of genetically far distinct hosts suggest that identification of the two *Thelohanellus* species by different authors was not correct and by more thorough examinations several morphologically similar but host specific species can be separated.

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