Molecular and phylogenetic characterization of *Thelohanellus qadrii* (Myxozoa, Myxosporea, Bivalvulida) infecting the secondary gill epithelium of Indian major carp, *Catla catla* (Hamilton, 1822)

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**ABSTRACT**

Myxosporean taxonomy which is traditionally based on the morphology of the myxospore stage, is in a state of flux given new insights provided by the expanding dataset of DNA sequences. To date, more than 40 species of *Thelohanellus* from India have been described according to morphometric characteristics. Nevertheless, molecular data on these histozoic myxosporean parasites of freshwater fish are scarce. In the present study, molecular characterizations of *Thelohanellus qadrii* infecting the secondary gill epithelium of Indian major carp *Catla catla* (Hamilton, 1822) and its phylogenetic relationship is reported. The sub-adult cultured catla were observed to have low to moderate gill myxosporean infections. The morphometry of mature spores was in compliance with original descriptions of *T. qadrii*. Based on the analysis of 18S rRNA gene, phylogenetic clusters which were established according to a consensus sequence, illustrated the taxonomic placement of a series of myxobolids. The DNA sequence homogeneity of *T. qadrii* (KF170928) with other *Thelohanellus* spp. ranged from 78% to 95% and formed a dichotomy with cyprinid gill lamellae infecting *T. toyamai* (HQ338729). Distance matrix results indicated a high genetic diversity among myxosporeans. The present report is the first on the molecular and phylogenetic characterizations of *T. qadrii*.

**Key words:** *Catla catla*; Myxosporean infection; *Thelohanellus qadrii*; Molecular characterization; Phylogenetic relationship

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INTRODUCTION

Over 62 genera of phylum Myxozoa have been described from fishes, primarily on the basis of myxosporean spore structure [1]. Among them, species of *Thelohanellus* Kudo, 1933 are typically histozoic (rarely coelozoic) bivalvulidan myxosporean parasites of freshwater fish with a high diversity of infection sites [2]. In recent years, myxosporeans have received much attention as emerging fish pathogens as they are continually threatening the development of aquaculture [3-6]. The increased importance of myxosporean diseases has initiated scientific research on life cycles, host-pathogen relationships and the development of diagnostic tests for myxosporean parasites. Molecular systematics using small subunit ribosomal DNA sequences and sensitive PCR tests are currently used to resolve specific relationships within the Myxozoa and to identify the parasites [3, 7-10]. *Thelohanellus* is the sixth most speciose myxozoan genus with 108 nominal species reported worldwide to date, of which 40 species are from India [11]. Molecular studies on Indian myxosporeans are rare. The authors have recently reported on the molecular characterization of *T. caudatus* infecting the caudal fin of *Labeo rohita* [12]. In this article, for the first time we present the molecular characterizations of *Thelohanellus qadrii* infecting the secondary gill epithelium of Indian major carp *Catla catla* (Hamilton, 1822) and its phylogenetic relationship.

MATERIALS AND METHODS

A total of 60 juvenile to sub-adult carp (*Catla catla*) from a polyculture pond in Garia (Lat. 22°27′59″N; Long. 88°24′18″E), Kolkata, West Bengal, India, were screened during the routine survey of carp myxosporean infections in 2013. Myxosporeans infecting the secondary gill epithelia of catla were collected and characterized by morphometric techniques. Myxosporean identification was performed according to Lom & Arthur [13]. In brief, a fresh plasmodium was first taken on a clean grease-free glass slide with a few drops of distilled water and then slightly ruptured. The spores released from the plasmodium were then spread onto clean grease free glass slides, covered with cover slips and sealed with Distrene, Plasticizer and Xylene (DPX) for examination under an oil immersion (100X) lens. Two fresh spore smears were treated with 2% KOH (w/v) for polar filament extrusion. The Indian ink method was employed to observe the mucous membrane around the spores. Smears of fresh spores were treated with Lugol’s iodine solution to observe iodonophilic vacuoles in the sporoplasm. For permanent slides, air dried smears were fixed with acetone free absolute methanol for about 8 min and stained with Giemsa solution for 40 min afterwards. The Giemsa solution was prepared by dissolving 0.5 g Giemsa powder in 33 mL glycerol at 50-60°C for 90 min in a water bath followed by the addition of 33 mL methanol. This solution was matured in the dark for 15 days and diluted with phosphate buffer (pH 7.2) with a 1:2 ratio prior to use. The slides containing myxosporean spores were observed under an oil immersion (100X) lens of a Motic BA400 microscope with...
an inbuilt digital camera. Morphometric measurements were done in μm by Motic Image Plus Version 2 software.

Morphologically identified myxosporeans was further characterized by molecular techniques as described previously [12]. After morphometric confirmation of the first plasmodium spores, the spores were suspended in a 500 μL lysis buffer (100 mM NaCl, 10 mM Tris, 10 mM EDTA, 0.2% SDS, 0.4 mg/mL Proteinase K) and incubated overnight at 55°C. Afterwards, 500 μL of phenol:chloroform (1:1) was added to the digested spores, mixed gently and centrifuged at 5200g for 10 min. The upper phase was later transferred to a new tube and mixed with a 1/10 volume of sodium acetate (3 M, pH 5.2) and 2 volumes of 96% ethanol (Amresco, USA). DNA was precipitated at –20°C overnight and pelleted by centrifugation at 10000g for 30 min. The pellet was washed once with 70% ethanol, air-dried for several minutes and resuspended in 30 μL of molecular biology grade water. The universal eukaryotic primers -ERIB1, 5´-ACC TGG TTG ATC CTG CCA G-3´ and ERIB10, 5´-CTT CCG CAG GTT CAC CTA CGG-3´ [14] were used for the amplification of 18S rDNA by Eppendorf Master cycler Pro S. The PCR was run using a mixture of 50 ng of genomic DNA, 10 μM of each primer and a 2X PCR TaqMixture. Amplification was done by initial denaturation at 95°C for 5 min, followed by 35 cycles of denaturation at 95°C for 30 sec, annealing of primers at 51°C for 30 sec and extension at 72°C for 60 sec. The final extension was at 72°C for 5 min. PCR product was analysed on a 1.5% agarose gel containing 0.5 μg/mL ethidium bromide in a 1X Tris-acetate-EDTA (TAE) buffer. Following the purification of the amplified PCR product by an EXO-SAP treatment, the DNA was quantified and subjected to automated DNA sequencing on an ABI 3730xl Genetic Analyzer (Applied Biosystems, USA). BigDye® Terminator v3.1 Cycle sequencing kit (Applied Biosystems, USA) was used for sequencing as per the manufacturers’ instructions. Electrophoresis and data analyses were carried out on the ABI 3730xl Genetic Analyzer.

Phylogenetic analysis was performed on a selection of 18S rRNA sequences that comprised the new sequence (KF170928) and 17 additional sequences from closely related species available in the NCBI GenBank database using the basic local alignment search tool (BLAST) and other representatives of the Myxobolidae clade (Table 1) as described by Fiala [9]. Buddenbrockia plumatellae (AY074915) of the class Malacosporea was used as an out-group.

The sequences were compared by standard nucleotide BLAST (www.ncbi.nlm.gov/BLAST/). Data analysis and multiple alignments were performed by ClustalX [15] and MEGA5 [16] software, respectively. Genetic distance analyses were conducted using the Kimura 2-parameter model [17]. Codon positions included were 1st + 2nd + 3rd + Noncoding. All positions containing gaps and missing data were eliminated. Evolutionary history was inferred using the maximum likelihood method.

A bootstrap consensus tree inferred from 1000 replicates was taken to represent the evolutionary history of the taxa analyzed. Branches corresponding to partitions reproduced in <50% bootstrap replicates were collapsed. The percentage of replicate trees in which the associated taxa were clustered together in the bootstrap test (1000 replicates) is shown next to the branches [18]. The nucleotide sequence generated in the
present study was deposited in the NCBI GenBank database under accession number KF170928.

RESULTS AND DISCUSSION

During the survey, 25 out of 60 cultured catla (41.66%) were observed to have low to moderate gill myxosporean infection. The plasmodia found on the secondary gill epithelia were very small (0.2-0.3 mm), whitish and oval shaped. Mature spores measured 14.80 ± 0.61 (14.0-15.9) × 5.52 ± 0.25 (5.0-5.9) μm. Spores were elongated pyriforms in valvular view with rounded posteriors and tapering anterior ends. In the sutural view, they appeared slightly lenticular with indistinct sutural lines. The polar capsule with a size of 8.51 ± 0.42 (7.6-8.9) × 3.22 ± 0.19 (3.0-3.7) μm was situated at the anterior portion and was similar to the shape of the spore. The flat anterior surface was situated at a short distance below the anterior end of the spore. The polar filament formed 14-16 coils within the capsule. The 93 μm long thread-like polar filament was wavy and tapering when completely extruded (Fig. 1). The sporoplasm contained two large nuclei and a large irregular shaped iodinophilous vacuole. Mucus envelopes around the spore and intercapsular processes (ICP) were absent. Based on morphometry, the myxosporean infecting the secondary gill epithelium was identified as Thelohanellus qadrii.

Figure 1: Wet mount preparation showing mature spores of Thelohanellus qadrii with extended polar filament at 200X magnification (bar = 1 μm)

Universal primers successfully amplified approximately 1708 bp fragments (Fig. 2) of the 18S rRNA gene of T. qadrii. The phylogenetic cluster was established based on the 1703 bp consensus sequence. The phylogenetic tree (Fig. 3) illustrated the taxonomic placement of a series of myxobolids based on the analysis of the 18S rRNA gene. The novel DNA sequence of T. qadrii clustered phylogenetically with other
Theelohanellus spp. with 78-95% homogeneity (Table 1). Evolutionary pair-wise distances among T. qadrii and other species of myxosporeans, measured by Kimura-2 parameter algorithm (Table 1), ranged from 0.00 (T. toyamai HQ338729) to 0.56 (Kudoa iwatai AY514038).

Figure 2: Agarose gel (1.5%) showing 18S rRNA gene amplification of Thelohanellus qadrii (1708bp) from Catla catla. 1: DNA ladder, 2: Thelohanellus qadrii

Figure 3: Phylogenetic tree generated by maximum likelihood of the 18S rRNA gene sequences of Thelohanellus qadrii (NCBI Accession number KF170928) and other Thelohanellus spp., and related taxa. Bootstrap confidence values are shown at nodes (1000 replications). Abbreviations: G-Gill, PF-Pelvic fin, CF-Caudal fin, I-Intestine, Vo-Various organs, S-Skin, Fi-Fin, Mu-Muscle, GB-Gall bladder, F-Freshwater clade, M-Marine clade, Cypr-Cypriniformes, Silu-Siluriformes, Perc-Perciformes, Salm-Salmoniformes. No details on infection site, host and habitat are available on Thelohanellus sinensis in NCBI GenBank database.
| Myxozoan species                                  | A* | 1   | 2   | 3   | 4   | 5   | 6   | 7   | 8   | 9   | 10  | 11  | 12  | 13  | 14  | 15  | 16  | 17  | 18  |
|-------------------------------------------------|----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| KF170928: *Thelepanulius quadrilobus*           | 100| 0.00|     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| KF476777: *Thelepanulius quadrispiculatus*       | 85 | 0.25| 0.00|     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| KF536776: *Thelepanulius quadrusomus*           | 85 | 0.25| 0.00| 0.00|     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| KF865807: *Thelepanulius caudatus*              | 78 | 0.43| 0.13| 0.13| 0.00|     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| IQ696557: *Thelepanulius katsurae*              | 87 | 0.22| 0.04| 0.04| 0.14| 0.00|     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| AY131401: *Thelepanulius horikoshi*             | 87 | 0.26| 0.07| 0.13| 0.04| 0.00|     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| HCJ381929: *Thelepanulius tojumai*              | 95 | 0.09| 0.25| 0.25| 0.43| 0.22| 0.25| 0.00|     |     |     |     |     |     |     |     |     |     |     |     |
| AY165116: *Thelepanulius wakamensis*            | 84 | 0.27| 0.10| 0.10| 0.22| 0.06| 0.07| 0.27| 0.00|     |     |     |     |     |     |     |     |     |     |     |
| DQ241156: *Thelepanulius ribicati*              | 85 | 0.27| 0.11| 0.12| 0.27| 0.10| 0.13| 0.26| 0.14| 0.00|     |     |     |     |     |     |     |     |     |     |
| EU036630: *Thelepanulius zakhrovi*              | 89 | 0.20| 0.20| 0.40| 0.20| 0.21| 0.19| 0.25| 0.23| 0.00|     |     |     |     |     |     |     |     |     |     |
| DQ502143: *Thelepanulius cinerascens*           | 87 | 0.23| 0.11| 0.12| 0.25| 0.06| 0.11| 0.22| 0.12| 0.18| 0.09| 0.00|     |     |     |     |     |     |     |
| KF995620: *Myxobolus carolinensis*              | 83 | 0.23| 0.11| 0.11| 0.21| 0.09| 0.09| 0.22| 0.12| 0.12| 0.18| 0.09| 0.00|     |     |     |     |     |     |
| AF380481: *Myxobolus neoarcticus*               | 87 | 0.21| 0.29| 0.29| 0.53| 0.30| 0.21| 0.20| 0.33| 0.20| 0.22| 0.29| 0.28| 0.00|     |     |     |     |     |
| AF021181: *Hemibapia arctica*                   | 85 | 0.26| 0.27| 0.43| 0.24| 0.26| 0.28| 0.30| 0.27| 0.25| 0.26| 0.32| 0.00|     |     |     |     |     |     |
| AY114028: *Kidae iwatai*                       | 82 | 0.56| 0.61| 0.62| 0.89| 0.60| 0.62| 0.55| 0.67| 0.57| 0.60| 0.61| 0.60| 0.63| 0.64| 0.00|     |     |     |
| AF001570: *Catenezaia zebra*                    | 81 | 0.61| 0.57| 0.57| 0.83| 0.57| 0.62| 0.50| 0.65| 0.64| 0.58| 0.60| 0.58| 0.55| 0.63| 0.33| 0.00|     |     |
| DQ577858: *Myxidium decoratum*                  | 81 | 0.63| 0.56| 0.56| 0.77| 0.52| 0.52| 0.57| 0.51| 0.57| 0.52| 0.53| 0.59| 0.53| 0.28| 0.37| 0.00|     |     |
| AY074916: *Buddistokichia plumatellus*          | 79 | 0.65| 0.73| 0.73| 1.00| 0.71| 0.76| 0.65| 0.73| 0.78| 0.72| 0.72| 0.71| 0.65| 0.70| 0.80| 0.73| 0.64| 0.00 |
Although morphological data on many Indian *Thelohanellus* spp. are available [2, 5, 11], molecular data are not. Prior to molecular characterization, we morphometrically characterized the myxosporean species isolated from the secondary gill epithelium of carp. The spore length (LS) to breadth (BS) ratio (1:0.37) and the length of polar capsules (LPC) to their breadth (BPC) ratio (1:0.38) were, more or less, in conformity with the original descriptions of *Thelohanellus qadrii* (LS:BS = 1:0.36; LPC:BPC = 1:0.46), a species described by Lalithakumari (1969) [19] from *Labeo potail* in Andhra Pradesh, India. Although the spore size slightly differed, it did not exceed the limits of natural variations typical of species' populations. Several other gill-infecting *Thelohanellus* spp. from Indian cyprinids have been described [11]. In the present study, a comparison of the morphometric data of *T. qadrii* with the representatives of *Thelohanellus* spp. infecting the gills of cyprinids [11] further revealed its morphometrical difference with the others. These observations, thus, confirmed that the *Thelohanellus* species found on the gills of cultured catla was *T. qadrii* in terms of morphology, host (carp) specificity and tissue (gill) tropism.

This is the first report regarding the molecular and phylogenetic characterization of *T. qadrii*. Earlier, we characterized *T. caudatus* (KC865607) infecting the caudal fin of *L. rohita* [12]. In this study, approximately 1708 bp fragments of the 18S rRNA gene of *T. qadrii* were successfully amplified. Our phylogenetic tree, established by the maximum likelihood method, defined topologies resembling those generated by Fiala [9]. According to Esztterbauer [8], site specificity is an important factor in myxozoan speciation. The evolutionary tree of this study further demonstrated that tissue tropism plays an important role in genetic relationships among myxozoan species. *Thelohanellus qadrii* clustered with cyprinid gill infecting myxosporeans with 89-95% homogeneity. The phylogenetic tree placed *T. qadrii* within the freshwater clade. It formed a dichotomy with the closely related *T. toyamai* (HQ338729) that infected the gill lamellae of common carp, *Cyprinus carpio* from the USA with maximum node support. In addition, *T. qadrii* exhibited maximum DNA sequence homogeneity (95%) with *T. toyamai* (HQ338729). Our previously characterized fin infecting *T. caudatus* (KC865607) exhibited 78% homogeneity with *T. qadrii*. Furthermore, freshwater and marine clades (*Myxidium incurvatum* (DQ377708) and *Kudoa iwatai* (AY514038)) were distinctly separated within the lineage Myxosporea. Other representatives of the Myxobolidae clade such as *Ceratomyxa, Henneguya, Kudoa, Myxidium* and *Myxobolus* were also distinctly different from the gill-infecting *Thelohanellus* spp. and clustered separately. The out-group *B. plumatellae* (AY074915) of the Malacosporea class was distinctly and phylogenetically clustered as a separate lineage with Myxosporea. The wide range observed in the evolutionary pair-wise distances between *T. qadrii* and other myxosporeans (0.00-0.56) including the gill infecting *Myxobolus carnaticus* (0.23) and the fin infecting *T. caudatus* (0.25-0.43) from India possibly indicated the high genetic diversity of myxosporeans.

Myxosporeans are best known for the diseases they cause in commercially important fish hosts. With the huge global expansion of freshwater aquaculture, several myxosporeans have been recognized or elevated in status as important pathogens [5, 6]. Since the gills of catla had low to moderate infections (with the parasitic frequency
index of 41.66%), negative impacts on their respiration and physiology can be generally assumed. Although no mortality was noticed, such infection may cause substantial production loss. Further studies are warranted to establish this pathogenicity and its role in production loss.

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Conflict of Interest: The authors declare that they have no competing interest.

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