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Research Article

# Characterization of *Hysterothylacium persicum* (Nematoda: Raphidascarididae) in *Scomberomorus commerson* from the Persian Gulf using morphological and molecular approaches

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#### **Keywords**

Scomberomorus commerson, Hysterothylacium persicum, Morphology, Molecular identification, Persian Gulf

#### Article info

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#### **Abstract**

Narrow-barred **Spanish** mackerel (Scomberomorus commerson) is one of the most important commercial fish species in the Persian Gulf and Oman Sea. Parasites of the genus Hysterothylacium (Ascaridoidea: Anisakidae) are common ascaridoid nematodes of fish-eating birds and sea The aim of this study is to identify mammals. morphological and molecular characteristics Hysterothylacium on S. commerson from April 2012 to December 2013. The morphological identification of Hysterothylacium specimens (n=10) was performed based on characteristic features, measurements and comparison with other publications. Larvae were counted, fixed in 70% ethanol and cleared in lactophenol for 48 h for identification. The molecular species description was based on a polymerase chain reaction (PCR) of sequences comprising ITS-1 and ITS-2. After the blast of nucleotide sequences, with the other recognized sequencing recorded in the GenBank, the most similarity was observed to Hysterothylacium persicum. Based on the morphological analysis and sequencing, the Hysterothylacium specimen was described as H. persicum. It is the first morphological and molecular identification of H. persicum in S. commerson from Iran. A combination of morphological description with a molecular technique seems to be the best practice for identifying anisakid larvae.

# Introduction

The Scombridae fish family contains 15 genera and about 50 species of epipelagic and generally migratory marine fish. It includes species of high commercial interest, such as mackerels, bonitos and tunas. Scombridae were distributed widely in the Persian Gulf and Oman Sea. Scomberomorus commerson (Teleostei. Perciformes, Scombridae) is a mackerel of the Scombridae family. This fish is a large and fast pelagic predator which is found in vast numbers in the tropical and subtropical waters (Kaymaram et al., 2010). Scomberomorus commerson is one of the most important and commercial species in the Persian Gulf and Oman Sea, which has many important roles in the foods of humans in the South of Iran.

Parasites play a crucial role in the biology and physiology of fish (Shamsi Barton, 2023). Adult anisakid nematodes of the genus Hysterothylacium are common parasites in the digestive tract of fishes in marine, brackish, freshwater environments. The Hysterothylacium includes more than 90 species, which are distributed in different continents (Pekmezci and Umur, 2015). Different species of Hysterothylacium have been reported from a wide variety of freshwater and saltwater fishes (Suthar and The occurrence Shamsi, 2021). Hysterothylacium sp. larvae was reported from Black sole fish, Brachirus orientalis (Bagherpour et al., 2011), greater lizardfish, (Saurida tumbil), Japanese thread fin bream (Nemipterus japonicus) (Dadar et al., 2016), Pick handle barracuda (Sphyraena jello) (Taheri Mirghaed et al.,

2016) and *Platycephalus indicus* (Azodi *et al.*, 2019) in Iran.

Fish-eating birds and marine mammals act as a final host, adult parasites live in the stomach or small intestines of these hosts. Cephalopods may act as a intermediate host (Roumbedakis et al., 2018) and various species of fish act as a second intermediate host. Adult worms. also, live in the intestines of marine mammals, dolphins whales and seals and the stomach of ducks, geese, swans, and a large number of aquatic birds. In humans, ingestion of raw or semi-raw fish, lightly salted or smoked fish leads to a severe disease known as anisakidosis (Mostafa et al., 2023). Symptoms of this disease depend on the location of the larval in organs. Third stage larvae in humans are likely to pierce the wall of the stomach or small intestine and symptoms such as vomiting, abdominal pain, fever, bloody diarrhea, eosinophilia, and emaciation were observed. In this disease, serologic particularly the enzyme-linked immunosorbent assay (ELISA) Western blot are very useful for clinical evaluation (Shamsi and Barton, 2023). No human anisakiasis has been yet reported from Iran, this issue could be attributed to the cuisine habit of fish in studied areas as well as other parts of the country.

The genus *Hysterothylacium* includes 89 accepted species, one taxon inquirendum, two nomina dubia, and ten unaccepted species (Hossen and Shamsi, 2019), but GBIF (2021) enlisted 97 species in this genus. Since morphological analysis of *Hysterothylacium* species is difficult and subjective, the developed molecular methods can be helpful and

useful. In the last decade, molecular biology techniques such as PCR and random amplified polymorphic have been used analysis identification of Hysterothylacium species (AlGabbani et al., 2021). PCR assays were used to differentiate among species of anisakid nematodes using the first and second internal transcribed spacers (ITS-1 and ITS-2) of ribosomal DNA (Simsek et al.. 2018). It is thought that the combination of morphological and molecular methods can be the best diagnostic strategy to identify the Hysterothylacium species. This study the identification presents of Hysterothylacium species S. commerson using both morphological and molecular methods.

### Materials and methods

One hundred S. commerson were caught from the Persian Gulf in Bandar Abbas in Hormozgan Province, south of Iran from April 2012 to December 2013. The fish were transported to the central laboratory of Shahrekord University for further analysis. The mean length and weight of  $40 \pm 19$ cm were  $5.0\pm0.2$ respectively. Then, the skin, abdominal cavity, stomach, sub-serous tissues, the contents of the stomach, intestine, livers, spleens, and gonads were inspected for parasites through naked eyes and under the standard investigation (Moravec, 2004; Buchmann, 2007).

The nematode larvae were primarily identified based on morphological characteristics (Shamsi *et al.*, 2013; Dadar *et al.*, 2016). Larvae were counted, fixed in 70% ethanol and cleared in lactophenol for

48 h for identification. Each parasite was photographed by a digital camera under a light microscope. Measurements were made under a microscope by means of eyepiece micrometer and the drawings were made. Photographs and drawings images of parasites were used for morphological analysis. Then, a middle piece of each nematode was fixed and stored in 70% ethanol for molecular analyses (Dadar *et al.*, 2016).

DNA was extracted using a DNA isolation kit (MBST, Iran) according to the manufacturer's instructions. Two pairs of primers were designed based on the sequence comprising ITS-1 and ITS-2 of *Hysterothylacium* sp. (Accession no: (5'-JN005755.1). Primer forward: GTAGGTGAACCTGCGGAAGGATC-3') was derived from nucleotides 1 to 23 of a sequence of 18S rRNA gene of Hysterothylacium spp. (Accession no. (5'-JN005755.1). Primer reverse TTAAATTCAGCGGGTAATCACG-3') was derived from nucleotide 903 to 924 of the sequence of the 28S rRNA gene of Hysterothylacium spp. (Accession no: JN005755.1).

The PCR was performed in a total reaction volume of 100  $\mu$ L containing: 10  $\mu$ L of 10X PCR buffer, 3  $\mu$ L MgCl<sub>2</sub> (50 mM), 2  $\mu$ L of dNTP (10 mM each), 0.5  $\mu$ L Taq DNA polymerase (5 U, Fermentas), 2  $\mu$ L of each primer (20  $\mu$ M), 74.5  $\mu$ L dH<sub>2</sub>O and 8 $\mu$ L of template DNA.

The reaction was repeated for 39 cycles under the following conditions: 5 min at 94°C (1 cycle), 45" at 94°C, 45" at 53°C, 45" at 72°C (37 cycles) and finally, PCR was completed with the additional extension step at 72°C for 10 min.

Distilled water was used as a negative control in each PCR reaction. PCR products were separated on 1.5% agarose gel in 0.5× Tris-borate-ethylene diaminetetraacetic acid (EDTA) buffer and stained with Ethidium Bromide and visualized by UV.

The PCR products were purified using a PCR purification kit (MBST, Iran) following the manufacturer's instructions. Purified fragments were sequenced from both sites of each PCR product using a method based on Sanger et al. (1977). Sequencing was carried out using the same primers as used for PCR amplification, by Kowsar Company. Sequences aligned using the computer program Clustal and then adjusted manually. The consensus data of all sequences were evaluated by the NCBI Basic Local Alignment Search Tool (Blast) to confirm matching rRNA gene. The ClustalW multiple alignment software was used to align the nearly complete rRNA genes of Iranian Hysterothylacium isolates with the different sequences present in the NCBI database. MEGA 6.0 was used to construct a phylogenetic tree using the Maximum Likelihood (ML) approach (Tamura et al., 2013). The tree with the highest log likelihood (-1719.1878) is shown. Initial tree(s) for the heuristic search were automatically obtained by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach and then selecting the topology with superior log likelihood value. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. The analysis involved 21 nucleotide sequences.

#### Results

Hysterothylacium larval were identified in 4 out of 100 fish samples (Fig. 1). The range of infection to Hysterothylacium was between 1-3 parasites. These larvae are characterized by excretory pores at the level of the nerve ring. The ventriculus had a nearly oval shape, the intestinal caecum was extended anteriorly and ventricular appendix was projected posteriorly. Four rectal glands were present. The conical tail was tipped with the arranged spines in a circle. Average length and width of body 7 mm (6–8) and 0.10 mm (0.07-0.13), respectively; lacking boring tooth; excretory pore opening just below nerve ring; nerve ring located from the 0.24mm (0.19-0.26)from the anterior end: muscular esophagus was 0.54 mm (0.44 – 0.61) long followed by a glandular ventriculus 0.05 mm long (0.04-0.06) and 0.03 mm (0.02-0.04) width; intestinal caecum 0.08 mm long (0.07-0.12); anus 0.12 mm (0.11-0.14) from the posterior end and tail conically shaped with a single spine. The survey revealed larval type consistent with Hysterothylacium type C in S. commerson (Fig. 2).

In this study, the amplification of nucleotide sequences of ITS-1 and ITS-2 were obtained from the individual *Hysterothylacium* parasite. After PCR assay, DNAs extracted gave the expected 889 bp PCR fragment, which is specific for Hysterothylacium parasite. After purification and sequencing of the PCR product, the nucleotide sequences revealed 889 bp lengths that were compared with

other *Hysterothylacium* sequences available in public databases (i.e. NCBI GenBank), By a BLAST search. Results showed that they are the most similar to *H. persicum*. The sequence of *H. persicum* isolated from *S. commerson* in the present study was deposited in Gen Bank under ID

number 2736622. The evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model (Tamura and Nei, 1993). The tree with the highest log likelihood (-1719.1878) is shown (Fig. 3).



Figure 1: Macroscopic Hysterothylacium larvae isolated from the intestine of Scomberomorus commerson.

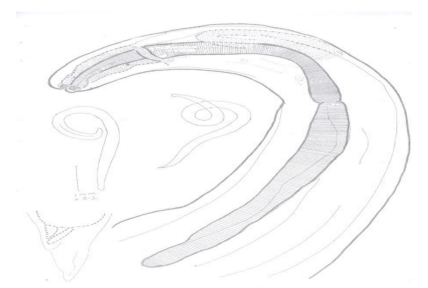


Figure 2: Drawing a picture of Hysterothylacium larvae type C from Scomberomorus commerson.

Initial tree (s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach and then selecting the topology with superior log likelihood

value. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. The analysis involved 21 nucleotide sequences. Codon positions included were 1st+2nd+3rd+Noncoding. All positions containing gaps and missing data were

eliminated. There was a total of 766 positions in the final dataset. Evolutionary analyses were conducted in MEGA6.

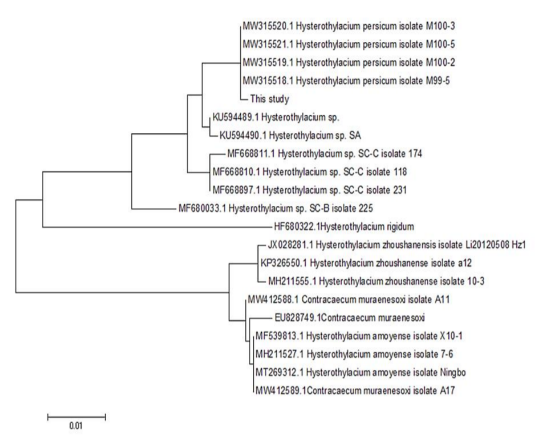


Figure 3: Molecular phylogenetic analysis by maximum likelihood method.

# **Discussion**

The Hysterothylacium genus has numerous species, the taxonomic status of many being unclear. Hence, species identification and distinguishing between some of them are not feasible (AlGabbani et al., 2021). Therefore, an identification of these species using improved approaches such as scanning electron microscopy and molecular techniques is essential. In Iran, only one systematic study of this group has been conducted (Shamsi et al., 2016).

One problem facing researchers when studying species of *Hysterothylacium* in Iran is that most of these species were poorly described. The present study supported the distinction among species of *Hysterothylacium* based on morphological data and was useful in confirming the taxonomic status of individual species. In the present study, using morphological as well as molecular methods, the systematic and taxonomic status of one species of *Hysterothylacium* in Iran has been reexamined.

In this study, from 100 pieces of *S. Commerson*, 4 pieces were infected with *H. persicum*. The range of contamination to *Hysterothylacium* was between 1-3 parasites, which were identified by morphological and molecular methods.

There have been other studies Hysterothylacium species in the Persian Gulf, such as the investigation by Taheri Mirghaed et al. (2016) which found a prevalence of 12% and a mean intensity of 4.8 in S. jello. Dadar et al. (2016) also collected anisakid larvae from two species of fish in the Persian Gulf, which were similar to Hysterothylacium type C. These studies highlight the presence Hysterothylacium species in the Persian Gulf and the need for further research on their taxonomy and identification.

It is important to highlight significance of phylogenetic analysis in this study, which confirmed the presence of H. persicum in the Persian Gulf and its similarity to other H. persicum in other regions. This analysis can provide insights into the evolutionary relationships and divergence of different species and can aid in the development of a comprehensive taxonomy for this group of nematodes. In our survey, molecular identification of Hysterothylacium was carried out by previously designed specific primers using PCR techniques. After the blast of nucleotide sequences, with the other recognized sequencing recorded in the Gene Bank, most similarity was observed to H. persicum. H. persicum was first described in the Persian Gulf by Shamsi et (2016). In parallel study, phylogenetic analyses of the current study showed that the highest similarity to H. persicum isolated from selected edible fish from the Persian Gulf (Shahmsi et al., 2016) and those previously described from Australasian (Shamsi et al., 2016) and Iraqi waters (Ghadam et al., 2018; Bannai, 2018). The result of the current study

agrees with the suggested of Mattiucci and Nascetti (2008) and Zhao et al. (2017) of "sibling identified the existence within species" the ascaridoids, morphologically very similar but genetically different in some specimens. Molecular identification using **PCR** techniques and specific primers can be a useful tool in identifying Hysterothylacium species. The blast of nucleotide sequences with those recorded in the Gene Bank can provide insights into the similarity of different species. For example, investigations in the Gene Bank revealed that species of the same genus of Hysterothylacium in studying gene locus were similar to 99%. However, further extensive studies are needed to provide more informative and useful data.

conclusion, the present provides the first report of H. persicum from S. Commerson in Iran using both morphological and molecular methods. The combination of these techniques can identifying Hysterothylacium species and confirming their taxonomic status. Considering the limited number of samples, it is evident that conducting molecular studies on a larger number of nematodes and investigating seasonal parasite contamination is essential in future studies. Further research on the and identification taxonomy of Hysterothylacium species is necessary, as many species remain poorly described and their taxonomic status unclear. The use of improved approaches such as scanning electron microscopy and molecular techniques can aid in species identification and differentiation. Studies such as the which present one, utilized both

morphological and molecular methods, can be useful in confirming the taxonomic status of individual species.

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## **Conflicts of interest**

The authors declare that they do not have any conflict of interest.

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