**Paragonimus paishuihoensis** Metacercariae in Freshwater Crabs, *Potamon lipkei*, in Vientiane Province, Lao PDR

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**Abstract:** Among *Paragonimus* species, *P. paishuihoensis* is one of the most mysterious and poorly understood species. Metacercariae are characterized by having a unique dendritically branched excretory bladder. However, the morphology of the adult worm remains unknown. To date, metacercariae of this species have been reported only in China and Thailand. In this study, we first found *P. paishuihoensis* metacercariae in freshwater crabs, *Potamon lipkei*, in Hinheub District, Vientiane, Lao PDR, with a prevalence of 77.7% and the average intensity of 10.3 (range 1-28) metacercariae per crab. The molecular data based on ITS2 and CO1 markers indicated that *P. paishuihoensis* from Laos and Thailand were almost completely identical and were close to members of the *Paragonimus bangkokensis/Paragonimus harinasutai* complex. Attempts to infect experimental animals (cats, dogs, and rats) with *P. paishuihoensis* were unsuccessful, suggesting that these animals might be unsuitable definitive hosts for the species. Further studies are necessary to elucidate the taxonomic status and life cycle of *P. paishuihoensis*.

**Key words:** *Paragonimus paishuihoensis*, metacercaria, freshwater crab, *Potamon lipkei*, Lao PDR

**INTRODUCTION**

Among about 50 nominal species of the genus *Paragonimus*, *Paragonimus paishuihoensis* Tsao and Chung 1965 is one of the most mysterious and poorly understood species because no adult has ever been obtained [1]. The metacercariae of this species, originally found in Szechuan (Sichuan) Province, China, are unique in the genus *Paragonimus* because they possess a highly branched excretory bladder [2]. After experimental infection in cats, only 1 juvenile worm was obtained [2]. Subsequently, Wang et al. [3,4] found *P. paishuihoensis* metacercariae in freshwater crabs in Yunnan Province, China. At almost the same time, Cui et al. [5] determined nuclear ribosomal second internal transcribed spacer region (ITS2) and a partial mitochondrial cytochrome c oxidase subunit 1 gene (CO1) sequenc-
es of *P. paishuihoensis* and another 4 *Paragonimus* species in China and revealed that *P. paishuihoensis* was genetically close to *Paragonimus bangkokensis*, *Paragonimus menglaensis* and *Paragonimus xiangshanensis*. In addition to those reports from China, Ranşinijü et al. [6] discovered *P. paishuihoensis* metacercariae in Chanthaburi Province, Thailand, and reported genetic similarities of this species to *P. bangkokensis* and *Paragonimus harinasutai* based on analyses of ITS2 and CO1 sequences. Here, we report morphological and molecular features of *P. paishuihoensis* metacercariae first found in freshwater crabs, *Potamon lipkei*, in Lao PDR.

**MATERIALS AND METHODS**

Collection of metacercariae

*Paragonimus* metacercariae were collected from freshwater crabs, *Potamon lipkei*, in Hinheub district, Vientiane Province, Lao PDR, by the methods described previously [7]. Nine crabs were examined individually in detail to determine the prevalence of each *Paragonimus* species and their distribution in crab tissues (Table 1). *P. paishuihoensis* metacercariae collected from...
Additionally, crabs used for experimental infection were not included in Table 1. Metacercariae were identified based on morphological characteristics. Metacercariae that have a dendritic excretory bladder were regarded as *P. paishuihoensis* and were used for experimental infection and for morphological and molecular analyses.

Experimental infection to laboratory animals

Fresh metacercariae having a dendritic excretory bladder were given to 5 cats, 1 dog, and 2 rats either orally or intraperitoneally (Table 2). Fecal examination was carried out 1 month after infection and weekly thereafter. Since none of those animals became *Paragonimus* egg-positive after several months, they were autopsied at the designated time (Table 2) to examine the lungs. In case of rats, the whole body was examined using pepsin digestion.

Molecular analyses

From 3 metacercariae of *P. paishuihoensis* (codes: MC32-MC34), we amplified and sequenced the ITS2 region using the primer pair 3S and A28 [8] and a partial CO1 gene using the primer pair of CO1-1-forward and CO1-2-reverse [9]. The methods for molecular study followed those described previously [10]. The obtained sequences were deposited in the GenBank with accession no. AB679285-AB679290. In addition, we downloaded and used further sequences of *Paragonimus* spp. from GenBank, including the available sequences (ITS2: AB471886 and CO1: AB471887) of *P. paishuihoensis* from Thailand and 1 ITS2 sequence (AB370191), which was originally registered as *Paragonimus* sp. from Laos and is now re-registered as *P. paishuihoensis* by us. Two sequence data sets, ITS2 and CO1, were used to reconstruct the phylogenetic trees. Each set was aligned using Clustal-W [11] with default options. Evolutionary analyses were conducted in MEGA5 [12]. Maximum likelihood trees were reconstructed using the best models: the Kimura 2-parameter with gamma rate heterogeneity (+G) for ITS2 tree and Hasegawa-Kishino-Yano model with invariable sites (+I) for CO1 tree.

RESULTS

**Morphology of *P. paishuihoensis* metacercariae**

When we examined *Paragonimus* metacercariae in the freshwater crabs, *P. lipkei*, caught in Hinheub District, Vientiane, Lao PDR, we found metacercariae having an excretory bladder of dendritic appearance (Fig. 1) in addition to 3 previously reported species; *P. harinasutai, Paragonimus heterotremus,* and *P. bangkokensis* [13,14]. By light microscopic observations, these metacercariae were spherical in shape with 2 cyst walls (Fig. 1A,B). The outer cyst (n = 8) was 702.0 ± 46.0 × 670.5 ± 49.1 µm (range 577-725 × 631-755) in diameter, while the inner cyst (n = 17) was 620.9 ± 30.9 × 573.9 ± 28.8 µm (range 518-622 × 580-681). A distinct space was present between the 2 walls. The thickness of outer and inner cyst wall was 4.9 ± 0.3 and 5.7 ± 0.7 µm, respectively. Two metacercariae were artificially excysted (Fig. 1C) and measured. They measured 1,502 × 737 and 1,436 × 533 µm respectively; oral suckers measured 105 × 118 and 117 × 112 µm, and ventral suckers 186 × 211 and 158 × 192 µm, respectively. A branched excretory bladder runs the
length of the body. All these morphological and morphometric data of encysted and excysted metacercariae were compatible with those of *P. paishuihoensis* reported previously [2,3,6].

Prevalence of *Paragonimus* metacercariae in crabs

In this study, 9 crabs were examined individually in detail to determine the prevalence of each *Paragonimus* species and their distributions in crab host tissues. Among 4 co-existing *Paragonimus* metacercariae in crabs, *P. harinasutai* was the most common species which was found in 8 (88.8%) crabs and accounted for 75.5% of total metacercariae found, followed by *P. heterotremus*, *P. paishuihoensis*, and *P. bangkokensis* (Table 1). Individual crab harbored metacercariae of 1-4 *Paragonimus* species. Regardless of the *Paragonimus* species, the majority of metacercariae were distributed in the muscles and hepatopancreas of the crab, and only a few were found in the gills or other tissues (Table 1).

Experimental infection

None of the 8 experimental animals (5 cats, 1 dog, and 2 rats), which were infected with *P. paishuihoensis* metacercariae, became egg-positive after infection (range of times until autopsy; 2.0-16.5 months). At autopsy, all of them were negative for *Paragonimus* worms (Table 2). No lesion suggestive of *Paragonimus* infection was found in the pleural and abdominal cavity of any experimental animal.

**Fig. 1.** Encysted form (A, B) and excysted form (C) of *P. paishuihoensis* metacercariae from crabs (*P. lipkei*) collected in Lao PDR.

**Fig. 2.** The Maximum Likelihood tree constructed from ITS2 (A) and CO1 (B) sequences. Bootstrap scores (percentages of 1,000 replications) are presented for each node. The sequences obtained from the samples in this study are represented in bold, while others from DNA database are shown with the GenBank accession no., species name, and country code.
Molecular analyses

For each gene region, the sequences of *P. paishuihoensis* isolates from Lao PDR were almost completely identical with each other, only 1 and 2 nucleotide differences were observed in the ITS2 (363 bp) and CO1 (354 bp) alignments, respectively. Similarities between the Lao sequences and those from the Thai sample were 99.9% in ITS2 (in spite of an AG indel) and 96.2% in CO1 sequences. Consequently, Lao and Thai samples formed a reasonably well-supported group in trees inferred from either gene region (Fig. 2). *P. paishuihoensis* was shown to be close to the *P. bangkokensis/P. harinasutai* complex.

**DISCUSSION**

The present study clearly showed that *P. paishuihoensis* is present in Vientiane Province, Lao PDR as the third country after China [2,3,13] and Thailand [6]. *P. paishuihoensis* metacercariae were often found together with 1-3 other Paragonimus species: *P. harinasutai*, *P. heterotremus*, and *P. bangkokensis* in a same crab, indicating that competition among these species of metacercariae may not so strict in *P. lipkei*. Consistent with a previous report [13], *P. harinasutai* was by far the commonest species in this study area (Table 1).

Until now, fully mature adult worms of *P. paishuihoensis* have never been found in wildlife, nor in experimentally infected animals except for 1 juvenile obtained by experimental infection in a cat in China [2]. We infected cats, a dog, and rats, which are known as susceptible definitive hosts for almost all *Paragonimus* spp. [1], with *P. paishuihoensis* metacercariae, but no worms were established themselves in these hosts. This indicates that dogs, cats, and rats are not suitable definitive hosts for *P. paishuihoensis*; it might have narrow host specificity to an unknown mammalian host, or require a paratenic host. If a paratenic host is required, our results suggest that it is not the rat; no juvenile worms were found when the body of each rat was completely digested. Further studies are necessary to elucidate the life cycle of *P. paishuihoensis*.

In molecular studies, Cui et al. [5] reported that *P. paishuihoensis* obtained in Yunnan Province, China, was genetically similar to *P. xiangshanensis* and *P. bangkokensis* from China. Unfortunately, those sequence data are not registered in GenBank. To date, *P. paishuihoensis* from Thailand is represented in GenBank by only a single ITS2 and a single CO1 sequence [6]. In the present study, molecular phylogenetic analyses revealed that *P. paishuihoensis* isolates from Lao PDR and Thailand were genetically almost completely identical with each other, thus forming a moderately to well-supported group in ITS2 and CO1 trees. Consistent with the previous reports [14,15], *P. paishuihoensis* sequences were clustered with *P. harinasutai* of China with a moderate bootstrap value of 76% to make a group separated from the rest of the *P. bangkokensis/P. harinasutai* complex. However, grouping with *P. paishuihoensis* and *P. bangkokensis/P. harinasutai* complex was supported by very low bootstrap values (less than 50%), especially in CO1 tree. For further investigation on taxonomic status in this group, more sampling and sequence data accumulation is necessary. Since only 1 each of ITS2 and CO1 sequence of *P. harinasutai* and none of other related species (*P. xiangshanensis* and *P. bangkokensis*) from China are available in the GenBank database, more sequence data of those species from China and also from other geographical locations would be useful for better understanding of their taxonomic status and phylogenetic relationship.

In conclusion, we demonstrated the presence of *P. paishuihoensis* metacercariae in Lao PDR, suggesting a wider distribution of this species in Indochina than previously suspected. Since the definitive hosts of *P. paishuihoensis* have not yet been identified, further studies are required to clarify the taxonomic status and life cycle of this species.

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