INTRODUCTION

Several kinds of mammals, i.e., humans, rats, cats, and dogs, as well as chicks, have been reported as definitive hosts of this fluke. This trematode is known to be the possible cause of clinical problems in some Asian countries [1,2]. In addition, metacercariae (= the infective stage) are mainly found in brackish water or marine fish, which serve as the second intermediate host, especially the group of mullets known as Mugil spp. and Liza spp. [3,4]. However, several previous studies have reported that S. falcatus metacercariae can also be found in freshwater fish, such as the half-beak (Dermogenys pusillus) [5] and climbing perch (Anabas testudineus) [6], for which a high infection rate has been reported, and the rate of infection has been found to be high in the northern region of Thailand. Even though the symptomatic features of human S. falcatus infection have not been clearly determined, the first clinical reports have revealed that S. falcatus can be found at autopsy and many embryonated eggs remained in the blood vessels of the cardiac muscle of the host [7]. Hence, it should be reminded that S. falcatus has the potential to retain its clinical importance as a great foodborne zoonotic trematode.

Nowadays, molecular approaches are the most efficient and accurate tools used to understand the relationships that exist in an organism and for screening the genetic variations that occur among many populations of parasites. The high annealing temperature-randomly amplified polymorphic DNA (HAT-RAPD) PCR technique had been applied to study the phylogenetic relationships of heterophyid trematodes [8,9]. However, currently, this technique is not commonly used because the HAT-RAPD PCR technique does not involve a DNA profile and exhibits a lower level of stability than other meth-
ods. The internal transcribed spacer 2 (ITS2) of nuclear rDNA is far more variable in the sequence, and because of the relatively rapid rate at which new mutants are fixed, these regions may reveal closely distinguished related species that otherwise would show little genetic divergence [10]. Therefore, phylogenetic analyses of ITS2 rDNA sequences have successfully been used to resolve evolutionary relationships among closely related species. Differing sequences of a given region are often assumed to be homogenized within a population of the same species by concerting evolution [11].

PCR based-methods targeting ITS2 have been applied for phylogenetic studies of various trematodes, such as Haplorchis taichui and H. pumilio [12,13], Clonorchis sinensis [14], Fasciola hepatica [15,16], and trematodes in the family Paramphistomidae [17]. Recently, the phylogenetic analysis of this gene was performed by more appropriate methods than the HAT-RAPD procedure applied in previous reports [5,8,9,18,19]. The purpose of this study was to investigate the prevalence of S. falcatus infection in the half-beaks (D. pusillus) and the worm recovery in experimentally infected chicks. In addition, phylogenetic relationships of S. falcatus with other heterophyid trematodes were analyzed for additional information to the recent reports [8,9].

**MATERIALS AND METHODS**

**Investigation of metacercarial infections in the fish host**

Thirty individual specimens of 4 species of freshwater fish, including D. pusillus, Henicorhynchus siamensis, Puntius brevis, and Cyclocheilichthys armatus, were subjected to analyses by surveying the metacercarial infections of heterophyid trematodes. Each individual fish was digested using an acid pepsin solution (conc. HCl 1 ml: pepsin 1 g: 0.85% sodium chloride solution 99 ml) for 3 hr at 37°C. The digested material was then rinsed with 0.85% sodium chloride solution and examined for metacercariae under a light microscope. Some metacercarial specimens were excysted and fixed in 4% formalin under the cover slip. The specimens were then stained with hematoxylin, dehydrated in alcohol series, cleared with xylene, and mounted in permount®. The metacercariae were examined and observed under a stereomicroscope. The specimens were measured and drawn under a compound microscope for morphological studies.

**Parasite preparation for phylogenetic analysis**

The metacercariae of S. falcatus, H. taichui, and C. formosanus were collected from different fish hosts; D. pusillus, H. siamensis, and P. brevis, respectively. Meanwhile, the adult specimens of Haplorchoidea sp. were gathered from the intestines of the Yellow cat fish, Hemibragus filamentosus, and the adult stages of other trematodes, namely the giant liver fluke, Fasciola gigantica and the rumen fluke, Orthocoeilium streptocoeilium, were recovered from cows (Bos taurus).

**Total genomic DNA extraction**

The genomic DNA of all parasites was extracted and purified from the adult worms using a DNeasy Tissue Kit (Vivantis, Oceanside, California, USA) according to the instructions of the manufacturer. All extracted genomic DNA were diluted to a working concentration of 30 ng/µl and stored at -20°C until used.

**Amplification of ITS2 region**

The PCR amplification of partial ITS2 fragments was done using a pair of primers as has been described in the previous report [20]. This consisted of (ITS3) 5’ GCA TCG ATG AAG ACGGCA GC 3’ as a forward primer and (ITS4) 5’ TCC TCCTGCTATG TAT GC 3’ as a reverse primer. The reaction was carried out at a final volume of 20 µl containing 1 × PCR buffer, 2 mM MgCl₂, 10 μM of each dNTP, 1 μM of each primer, and 1 U of Vivantis Tag DNA polymerase. The analysis was then performed in a MyCycler™ Thermocycler (BioRAD, Her-
cules, California, USA). Recommended PCR protocols were as follows; 1 cycle of 95°C for 2 min, 35 cycles of 94°C for 45 sec, 50°C for 45 sec, 72°C for 1 min, and 1 cycle of final extension at 72°C for 7 min. Specific PCR products were separated on 1.4% TBE agarose gel electrophoresis, stained with ethidium bromide, and photographed using a Kodak digital camera Gel Logic 100 and direct sequenced by the dideoxy terminator method (First Base Company). The sequence was checked using the BLAST program in the NCBI (National Center for Biotechnology Information, Bethesda, Maryland, USA) database, to confirm the PCR target. The eletropherograms of each sequence were examined for sequence accuracy using a Sequence Scanner version 1.0 and Bioedit version 7.1. All sequences were aligned automatically using Clustal X version 2.0. The phylogenetic relationships of 11 trematodes were constructed using MEGA version 5.0. All molecular data were analyzed by UPGMA (Unweighted Pair Group Method with Arithmetic Mean) method (Table 1). The reliability of the internal branches was assessed using the bootstrap method per 1,000 replicates.

Table 1. List of materials and sequences of ITS2 used for phylogenetic analysis

| Species                    | Accession no./ references |
|----------------------------|----------------------------|
| Haplorchis taichui         | KJ630831                   |
| Haplorchoides sp.          | KJ630832                   |
| Stellantchasmus falcatus   | KJ630833                   |
| Orthococcoellum sterpiocoelem | KJ630834                 |
| Fasciola hepatica          | KJ630835                   |
| Centrocestus formosanus    | KJ630836                   |
| Haplorchis taichui         | EU826640*                  |
| Metagonimus yokogawai     | AB470525*                  |
| Centrocestus sp.           | KF658459*                  |
| Opisthorchis viverrini     | AY584735*                  |

*The sequence data were from GenBank database.

Table 2. Summary of fish species, prevalence, and intensities of heterophyid trematodes examined in each fish species

| Fish species                | No. of fish infected with each heterophyid species/No. of fish examined | Mean |
|-----------------------------|--------------------------------------------------------------------------|------|
| S. falcatus                 | H. taichui                                                               |      |
| Dermogenys pusillus         | 27/30                                                                    | 90.0%|
| Henicorhynchus siamensis    | 0/17/30                                                                  | 83.3%|
| Puntius brevis              | 0/0                                                                      | 33.3%|
| Cyclocheilichthys armatus   | 0/0                                                                      | 23.3%|

*The intensity values represent the mean no. of metacercariae per fish (range).
μm in width. The right testis was found to be 42.5-90.0 μm long and 20.0-42.5 μm wide, while the left testis was 37.5-70.0 μm long and 22.5-30.0 μm wide. On day 2 PI, mature worms were observed. The body measured 360.0-510.0 μm in length and 130.0-210.0 μm in width. The ovary enlarged to 30.0-62.5 μm long and 27.5-50.0 μm wide. The right testis was 55.0-87.5 μm long and 32.5-52.5 μm wide, while the left testis was 47.5-80.0 μm long and 30.0-52.5 μm wide. The expulsor was oval and relatively large, with a length of 32.5-82.5 μm and a width of 25.0-37.5 μm. The ventrogenital sucker was slightly submedian to the right side and contained a ventral sucker. The ventral sucker armed with 2 dense lateral groups of small spines on the lip. The ovary was submedian and located slightly to the right side between the ventral sucker and right testis and was 30.0-62.5 μm long and 27.5-50.0 μm wide. The vitellaria extended dorsally into the ovariotesticular zone (Fig. 1D).

**Fig. 1.** Morphologic changes of *Stellantchasmus falcatus* recovered from experimental chicks (A-C) and illustrations demonstrating the permanent slide capture and line-drawing (D).

**Fig. 2.** The rooted phylogeny from partial ITS2 sequences of heterophyid trematodes based on the UPGMA method. Bootstrap values were computed independently for 1,000 resembling.
Phylogenetic analysis

The partial sequence data of the ITS2 region were used to understand their phylogenetic relationships. The length of this fragment was 410-590 bp. The trematodes in the family Heterophyidae appeared to be in a monophyletic clade. The heterophyids were separated into 2 sister groups including S. falcatu's group and the other heterophyid group (Fig. 2).

DISCUSSION

In this study, we reported the metacercarial infection status of 4 heterophyid flukes, i.e., S. falcatu's, H. taichu, Haplorchoides sp., and C. formosanus, which were collected individually from their fish hosts to be 90.0%, 86.7%, 33.3%, and 23.0%, respectively. Our findings corresponded well with previous reports that have confirmed a high prevalence of infection in S. falcatu's and H. taichu, particularly in the northern part of Thailand [21-23]. Various reports have detected the presence of intestinal trematodes in the family Heterophyidae, such as, H. taichu, H. pumilio, S. falcatu's, C. formosanus, and Procerovum sp., which were found to be wildly distributed and had a high prevalence and intensity. Previous surveys have also revealed that the metacercariae, which were mostly found in Mae Taeng District of Chiang Mai Province, belonged to the family Heterophyidae [21-23]. Our findings corresponded well with those of Radomyos et al. [24] who reported that certain heterophyid species were widely distributed in certain regions in the north of Thailand. Epidemiological surveys from 1997-2013 have shown an increase in trematode infections in the northern and central parts of Thailand. The surveyed region has been reported as an endemic area of heterophyid trematodes with high prevalences in a variety of intermediate and definitive hosts. For instance, Wongsawad et al. [6] investigated on trematode infections in freshwater fish from Mae Sa stream, Chiang Mai Province, where 4 species of heterophyid trematodes were found, i.e., Haplorchis spp., Haplorchoides sp., C. formosanus, and S. falcatu's [6].

From the present results, it is strongly suggested that the half-beaks (D. pusillus) in northern Thailand typically serve as the highly compatible secondary host for infection with S. falcatu's. This result corresponded well with a previous report [25]. In contrast, the metacercarial stage of S. falcatu's has been detected also in marine or brackish water fish [3,4]. However, further studies must be done to investigate S. falcatu's metacercariae in fish species from a larger area in order to confirm that this fish species (D. pusillus) serves as the most popular second intermediate host of this parasite.

Based on the results of experimental infection, chicks seemed to be a susceptible host for S. falcatu's infection revealing much higher rates of infection and worm recovery than those observed in mice (Mus musculus) (13.6%) and rats (Rattus norvegicus) (20.2%) [26]. These results are supported by the far higher worm recovery that was recorded in mice (13.6%) and chicks (33.8%). In our study, parasites were observed to have invaded the jejunum during days 1-6 PI and later they were typically observed in the ileum. A similar pattern was observed in H. taichu and H. pumilio which were found to have invaded the upper part of the small intestines during the first period of infection, but when they matured the parasites were more common in the middle and lower parts of the intestinal tract because the flukes became gradually excreted into the lumen of the intestinal canal along with mucus and other exudates. The flukes then reattached themselves further down the tract [27]. However, host-parasite relationships in heterophyid infections have never been studied in detail. Since human infections with S. falcatu's and other heterophyid trematodes are expected to increase in areas where fresh and brackish water fish are popularly eaten raw, studies on host-parasite relationships are considered highly useful.

According to the phylogenetic analysis, the partial sequences of ITS2 of some heterophyid trematodes are declared to be known now. However, only a few studies have focused on the relationship that exists among the ITS2 of S. falcatu's in Thailand [28]. Our sequence data of ITS2 agreed well with the sequence data reported in other recent reports [28].

The UPGMA trees showed the monophyletic relationships of heterophyid trematodes. Heterophyid trematodes were separate in a same group except S. falcatu's which was separated as another group with a high bootstrap value. This agrees with the reports on phylogenetic relationships based on mitochondrial cytochrome c oxidase 1 (mtCOI) sequence data, which summarized that only S. falcatu's has a ventrogenital complex [29], whereas the results of a phylogenetic study of S. falcatu's using the HAT-RAPD method showed a group of S. falcatu's together with other heterophyid trematodes [8].

ITS2 has been used to study the systematic analysis of some heterophyid trematodes, i.e., Haplorchis [12,13] and Metagonium [30]. From our results, it is believed that the ITS2 locus could be used as a barcode for authenticating animal species, as well as a complementary locus to other barcoding genes for
the purpose of identifying all organisms, including heterophyid trematodes. Several regions that have become known for a relevant presence of nuclear ribosomal DNA (nu rDNA) were selected and used for identification of various stages as well as for a study of the life cycle of trematodes (cercarial, metacercarial, and adult stages) in freshwater fish and snails because of the high levels of accuracy, sensitivity, and rapidity of this method. ITS2 has been revealed as a sensitive marker at the species level for trematodes. The PCR-based method targeting ITS2 has been applied for use in the phylogenetic study of various trematodes, such as *H. taichui* and *H. pumilio* [12,13], *C. sinensis* [14], *F. hepatica* [15], and trematodes in the family Paranthistomidae [17]. Using the sequence data of the ITS2 region, the species-level identification could be achieved, and the ITS2 analysis could actually provide a phylogenetic study of these trematodes.

This study can conclude that *D. pusillus* fish serves as a second intermediate host for *S. falcatus* with the highest rate of prevalence and rapid development being found in the experimental host, chicks (*G. gallus domesticus*). Finally, the present study provided valuable and beneficial information that can be used to demonstrate the epidemiological situation, as well as to examine the development and phylogenetic relationships of *S. falcatus* with regard to the biological control and monitoring of this parasitic disease.

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**CONFLICT OF INTEREST**

We declare that we have no conflict of interest related to this study.

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