Phylogeographic genetic variation of \textit{Indoplanorbis exustus} (Deshayes, 1834) (Gastropoda: Planorbidae) in South and Southeast Asia

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\textbf{A B S T R A C T}

The freshwater snail \textit{Indoplanorbis exustus} play an important role as the sole intermediate host of several medically- and economically-important trematodes, especially zoonotic schistosomes and echinostomes, which can infect and cause diseases in livestock and people. This study aims to explore the mitochondrial cytochrome c oxidase subunit I sequence variation of \textit{I. exustus} collected from new geographical areas; 459 specimens of \textit{I. exustus} were collected from 43 localities in South and Southeast Asia. The 42 haplotypes (Ie1 – Ie42) we detected were classified into haplogroups I – V. Phylogenetic analyses revealed five major clades, A – E, in concordance with all previous studies. Clade E contained two subclades, E1 (haplogroup I) and E2 (haplogroup II). The most widespread genetic group was subclade E1. Clade A, clade B (haplogroup V), and clade C (haplogroup IV) were found only in South Asia, whereas clade D (haplogroup III) was specifically found in Southeast Asia. In Thailand, \textit{I. exustus} showed high genetic divergence with 21 haplotypes. Several isolates showed significant genetic differences from others with unique haplotype(s). Hence, we confidently conclude our findings support all previous studies that \textit{I. exustus} is a species complex with at least four major lineages and five haplogroups. Our additional analyses of 35 samples from Sri Lanka showed these were indeed an independent genetic group as previously found, but they can now be classified as a unique group forming subclade E2 (haplogroup II) of \textit{I. exustus} sensu lato.

\textbf{ARTICLE INFO}

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1. Introduction

The freshwater snails, \textit{Indoplanorbis exustus} (Deshayes, 1834) are intermediate hosts for several medically-important parasites, such as those responsible for cattle schistosomiasis, especially the \textit{Schistosoma indicum} species group. They are also intermediate hosts for the zoonotic intestinal trematodes \textit{Artyfechinostomum sufrartyfex} and \textit{Echinostoma sp}. [26,29]. In addition, \textit{I. exustus} has been implicated in human cercarial dermatitis outbreaks in India, Lao PDR, Malaysia and Thailand [6,13,21,24]. Cercarial dermatitis is caused by a cutaneous allergic reaction in people who are exposed to larval schistosomes (cercariae) which are shed by infected snails into freshwater bodies (e.g. ponds, lakes and rice paddy fields). Cercarial infections cause popular pruritic eruptions: severe secondary infections may result from cercarian attempts at infecting non-permissive definitive hosts and subsequently dying in the skin [20].

In addition, \textit{I. exustus} is a hermaphroditic, invasive snail species with high fecundity and self-fertilization ability. It has the ability to colonize habitats containing other well-established pulmonates and prosobranch snails within a year [25]. In optimal conditions, e.g. 30 °C, each snail can
lay up to 968 eggs every 23 days [27]. The snails commonly attach to aquatic plants in small ponds, pools, tanks, lakes including stagnant pools of water in rivers and reservoirs, and semi-permanent pools flooding fields, where they can survive buried in mud during the dry season. In the latter case, dispersal may occur in clumps of mud stuck to cattle and water buffalo or disperse via water in debris [17]. Currently, *I. exustus* is known to be a widely distributed species in tropical and subtropical zones throughout South and Southeast Asia, southern China and across Africa to the French West Indies [18,19]. In addition, sea flooding fields, where they can survive buried in mud during the dry pools of water in rivers and reservoirs, and semi-permanent pools aquatic plants in small ponds, pools, tanks, lakes including stagnant pools of water in debris [17], which may comprise more than one species or, in fact, be a species complex [5,10,17,19]. It was initially classified as clades existing in north India (Assam and Bangladesh) and Cratonic India (Sri Lanka), ubiquitous in Thailand, the Philippines, and Nepal, Sundaic in Indonesia, as well as Arabian and Malaysian clades [17]. Subsequently, these clades were classified and named as clade A to clade E corresponding to their geographical origins. For instance, clade A contained samples from Nepal, clade B from Myanmar and Nepal, clade C from Lao PDR, clade D from India and Nepal, and clade E from a wide distribution covering Sri Lanka, Oman, Nepal, Myanmar, Thailand, Indonesia, Malaysia, and the Philippines [5,10]. More recently, the five major clades were categorized into only four clades. The previously classified clade D and C by Gaubre-Autelin et al. [10] were merged as one clade, following examination of new localities from Africa and the French West Indies [19].

The mitochondrial cytochrome c oxidase subunit 1 (CO1) has been shown to be the best potential marker to elucidate phylogeographic genetic variations in this snail [10,19]. The majority of previous studies have used the CO1 sequence to examine the molecular systematics of *I. exustus*. Unfortunately, previous studies analyzed few geographical localities within the broad-scale distribution clades of *I. exustus* [5,10,17,19]. Thus, our study aims to use the CO1 sequence to examine phylogeographic genetic variations of *I. exustus* collected from novel localities in South and Southeast Asia and in particular, a detailed

### Table 1
Sample collection localities and haplotype frequencies examined in this study.

| Code | Province | District | Region | Country | Habitat | No. | Haplotype frequency |
|------|----------|----------|--------|---------|---------|-----|--------------------|
| CMI  | Chiang Mai | San Sai | North | Thailand | Canal | 10 | Ie1(8), Ie2(2) |
| NAM  | Nan | Na Noi | North | Thailand | Canal | 10 | Ie1(10) |
| CRI  | Chiang Rai | Wiang Pa Pao | North | Thailand | Paddy field | 10 | Ie1(9), Ie7(1) |
| LPG  | Lampang | Muang | North | Thailand | Canal | 10 | Ie1(5), Ie2(2), Ie9(3) |
| PRE  | Prat | Kong Kwang | North | Thailand | Swamp | 10 | Ie6(1), Ie4(3), Ie15(1) |
| UTT  | Uttaradith | Muang | North | Thailand | Canal | 10 | Ie7(1), Ie11(3) |
| PNB  | Petchaboon | Losmak | Central | Thailand | Swamp | 10 | Ie10(1) |
| PLK  | Pitsanulok | Wang Thong | Central | Thailand | Paddy field | 10 | Ie1(10) |
| PCT  | Pichit | Sak Lek | Central | Thailand | Canal | 10 | Ie6(10) |
| STI  | Sukhothai | Muang | Central | Thailand | Canal | 10 | Ie1(10) |
| PTE  | Pathumthani | Klong Luang | Central | Thailand | Canal | 10 | Ie8(1), Ie4(1), Ie5(1) |
| LRI  | Lopburi | Muang | Central | Thailand | Canal | 10 | Ie10(1) |
| AYA  | Ayuthaya | Wang Noi | Central | Thailand | Swamp | 10 | Ie3(1), Ie7(5), Ie19(2) |
| ATG  | Arngthong | Wiset Chai Chan | Central | Thailand | Paddy field | 10 | Ie1(10) |
| SPR  | Supan Buri | Muang | Central | Thailand | Swamp | 10 | Ie6(10) |
| NYK  | Nakhon Nayok | Ongkhark | Central | Thailand | Canal | 10 | Ie8(1), Ie4(2) |
| KRI  | Kanchanaburi | Muang | West | Thailand | Canal | 10 | Ie5(1), Ie20(5) |
| TAK  | Tak | Mae Sot | West | Thailand | Canal | 10 | Ie1(10) |
| RBR  | Ratchaburi | Suan Phuang | West | Thailand | Swamp | 10 | Ie9(1), Ie7(1) |
| CBI  | Chonburi | Ban Bueng | East | Thailand | Swamp | 10 | Ie4(10) |
| UBN  | Ubon Ratchathani | Khong Chiam | Northeast | Thailand | Paddy field | 10 | Ie1(10) |
| YST  | Yasothon | Muang | Northeast | Thailand | Canal | 10 | Ie10(1) |
| KKN  | Khon Kaen | Muang | Northeast | Thailand | Canal | 10 | Ie7(1), Ie8(3) |
| NKI  | Nong Khai | Muang | Northeast | Thailand | Swamp | 10 | Ie5(1), Ie4(5) |
| UDN  | Udonthani | Muang | Northeast | Thailand | Swamp | 10 | Ie10(1) |
| ACR  | Amnatchareon | Muang | Northeast | Thailand | Canal | 10 | Ie16(1), Ie18(4) |
| NBP  | Nong Boua Lamphu | Non Sang | Northeast | Thailand | Swamp | 10 | Ie10(1) |
| MKM  | Maha Sarakham | Kantharakwachai | Northeast | Thailand | Paddy field | 10 | Ie3(1), Ie10(7) |
| RET  | Roi Et | Muang | Northeast | Thailand | Swamp | 10 | Ie1(10) |
| BRM  | Buri Ram | Kranang | Northeast | Thailand | Paddy field | 10 | Ie5(1), Ie4(5) |
| NRT  | Nakho Rat Chasima | Choe Chai | Northeast | Thailand | Paddy field | 10 | Ie6(1), Ie12(3), Ie13(1) |
| STN  | Satoon | Muang | South | Thailand | Swamp | 10 | Ie10(1) |
| SNI  | Surathani | Hat Yai | South | Thailand | Canal | 10 | Ie21(9), Ie22(1) |
| LVV  | Viantiane | Vang Vieng | -- | Lao PDR | Song River | 10 | Ie8(1), Ie6(2) |
| LVF  | Viantiane | Vang Vieng | -- | Lao PDR | Canal | 10 | Ie1(10) |
| LSV  | Savannakhet | Kaisone Phomvihan | -- | Lao PDR | Canal | 10 | Ie10(1) |
| CSR  | Siem Reab | Siem Reab | -- | Cambodia | Canal | 10 | Ie10(1) |
| CPS  | Pursat | Pursat | -- | Cambodia | Swamp | 10 | Ie26(6), Ie27(4) |
| CPF  | Phnom Penh | Phnom Penh | -- | Cambodia | Swamp | 10 | Ie23(10) |
| CKC  | Kampong Cham | Kampong Cham | -- | Cambodia | Swamp | 10 | Ie24(4), Ie25(6) |
| PHP  | Kampot | Mografi | -- | Philippines | Canal | 13 | Ie13(1), Ie42(5) |
| BAN  | Mymensingh | Mymensingh | -- | Bangladesh | Canal | 11 | Ie28(7), Ie29(9), Ie30(1), Ie31(1), Ie32(1) |
| SRI  | Central | Barigama | -- | Sri Lanka | Canal | 13 | Ie33(2), Ie34(4), Ie35(3), Ie36(2), Ie37(1), Ie38(1) |
| NRT  | North Central | Sekirawa | -- | Sri Lanka | Canal | 9 | Ie35(3), Ie36(2), Ie40(2) |
| NRT  | North Central | Polonnaruwa | -- | Sri Lanka | Canal | 8 | Ie35(6), Ie41(2) |

\(^a\) Regions of Thailand.

\(^b\) Number in the bracket after haplotype name indicates the observed number of a particular haplotype in each locality.
analysis of samples collected on a fine geographical scale at localities throughout Thailand, including CO1 sequences generated by all previous studies.

2. Materials and methods

2.1. Sample collection

The 459 specimens of I. exustus were collected from various natural habitats, e.g. swamps, rivers, canals, and rice paddy fields. Samples were collected from 33 geographical localities in Thailand, three localities in Lao PDR, four localities in Cambodia, four localities in Sri Lanka, one locality in the Philippines, and one locality in Bangladesh (Table 1). After collection, snails were preserved in 80% alcohol before transport to the laboratory. Each snail species was morphologically identified as described by the Danish Bilharziasis Laboratory [4]. The ethics of animal use and biosafety for this research were approved by The Animal Care and Use Committee and the Institute Biosafety Committee of Thammasat University, respectively.

2.2. Molecular analyses

Snails were individually used for total genomic DNA extraction using the E.Z.N.A.® Mollusc DNA Kit (Omega Bio-Tek, USA), following manufacturer’s instructions. The mitochondrial CO1 gene was amplified using primers LCO1490 (5’-GGT CAA CAA ATC ATA AAG ATA TTG G-3’) and HCO2198 (5’-TAA ACT TCA GGG TGA CCA AAA AAT CA-3’) [9]. PCR consisted of a 25 µl final reaction volume containing 1× Ex buffer (Takara, Japan) with 0.2 mM each of dNTP, 0.2 µM of each forward primer, 0.625 U of Ex Taq DNA Polymerase (Takara, Japan), and 1 µl (−10–50 ng) of the DNA sample. PCR conditions used for both gene amplifications were: 94°C for 4 min followed by 35 cycles of 1 min each of 94°C, 50°C, and 72°C, final extension at 72°C for 8 min. Amplified products were analyzed using 1.0% agarose gel electrophoresis, which was then cut and purified using an E.Z.N.A.® Gel Purification Kit (Omega Bio-Tek, USA). Purified PCR products were cycle-sequenced at Eurofins Genomics Company, Japan.

2.3. DNA sequence and haplotype analyses

Sequences were manually checked and edited using ABI Sequence Scanner Software v1.0 and BioEdit v7.2.6 [11]. Multiple sequence alignment was performed using ClustalW [15]. Genetic diversity indices, i.e. the segregating sites, haplotype number, haplotype diversity, and nucleotide diversity, were computed and generated by DnaSp v5 [16]. Pairwise values of genetic differentiation (ΦST) based on the Kimura two-parameter model [12] and genetic structure (Analysis of Molecular Variance: AMOVA) were calculated in Arlequin v.3.5.2.2 [8]. The minimum spanning haplotype network of CO1 and rrnL sequences was constructed with Network v.5.0.1.1 (http://www.fluxus-engineering.com) based on the median-joining algorithm [1]. Based on haplotype network analysis, the haplogroups of CO1 sequences were defined based on mutational steps (ms) ≥ 10.

2.4. Phylogenetic analyses

Time-reversible variation data among sites which were generated from a gamma distribution model (GTR + G + I) [22] were selected for analyses by maximum likelihood (ML) and Bayesian inference (BI) methods as determined by the MrModelTest v.2.3 [23]. The ML tree was constructed using MEGA X [14]. The BI was run using MrBayes v.3.2 [28] with 15,000,000 generations using four chains. Trees were sampled every 100th generation, resulting in 150,000 trees. The first 110,000 trees were excluded as burn-in because the average standard deviation of split frequencies (ASDF) was >0.01. Furthermore, convergence of the run was assessed by checking the potential scale reduction factor (PSRF) values of each parameter in MrBayes as well as the effective sample size (ESS) values of each parameter in Tracer 1.6 (http://beast.bio.ed.ac.uk/Tracer). A value of PSRF ~1.00 and of ESS > 200 were considered suggestive of convergence. The remaining 40,000 trees were used to calculate Bayesian posterior probabilities of individual clades. The 98 CO1 sequences of I. exustus deposited in GenBank from previous reports, namely HM104223, GU451738 – GU451750 [17], KR811332 – KR811361 [5], KY024420 – KY024472 [10], and MH037074 – MH037075 [19] were included, whereas a CO1 sequence of Balinus tropicus (AY282583) was used as an out-group. The Poisson tree processes model (PTP) for species delimitation and a Bayesian implementation of the PTP (bPTP) were used to infer putative species boundaries on a given phylogenetic input tree [31]. The following parameters were included in PTP and bPTP analyses, namely, MCMC, 500000 generations; thinning, 100; burn-in, 0.1; and seed, 123.

3. Results

3.1. Sequence analyses

The CO1 sequences of 459 individual I. exustus from 43 different localities in South and Southeast Asia were determined and deposited in GenBank under accession numbers MT274329 – MT274370. Genetic divergence was detected within all populations, revealed by nucleotide diversity ranging between 0.001 ± 0.001 to 0.043 ± 0.023, with an average of 0.012 ± 0.001. Haplotype diversity varied from 0.129 ± 0.079 to 0.831 ± 0.024, with an average of 0.671 ± 0.087 (Table 2). Population comparisons defined by different countries showed genetic differentiation (ΦST) was between 0.048 and 0.962, and there were significant differences (p < 0.05) between all populations, the exceptions being Thailand vs. Lao PDR, Cambodia vs. Philippines, Lao PDR vs. Philippines (Table 3). Comparisons between 33 different micro-geographical localities (isolates) in Thailand revealed genetic differences (ΦST) between 0.000 and 1.000. Comparisons between the 33 Thai isolates found that the isolates of Pitchit (PCT), Ayuthaya (AYA), Supanburi (SPB), Chonburi (CBI), Maha Sarakham (MKM), and Sukhothai (STI) provinces were significantly genetically different (ΦST) when compared to all other isolates (Supplementary Table 1).

3.2. Haplotype analyses

Based on nucleotide variation of the CO1 sequence, the 459 samples were classified into 42 haplotypes (Ie1 – Ie42) in our study. The most common haplotype detected in Southeast Asia was Ie1. Another common haplotype was Ie4, found in several geographical localities in Thailand, and haplotype Ie7 found in isolates from Chiang Rai (CRI) and Ayutthaya (AYA) provinces (Table 1, Fig. 1). All remaining haplotypes were found to be unique in a particular isolate, such as haplotype Ie3 being found only in the Pitchit (PCT) isolate, Ie21 and Ie22 unique to the Satoon (STN) isolate, Ie24 was detected only in Kampong Cham, Cambodia, and Ie42 was present only in Manila, and the Philippine isolate. The South Asian isolates clearly differed from Southeast Asian ones with haplotypes Ie28 – Ie32 seen only in Bangladesh, whereas Ie33

| Population | n | S | H | \( π \pm SD \) | \( Hd \pm SD \) |
|------------|---|---|---|----------------|----------------|
| Thailand   | 330 | 28 | 21 | 0.002 ± 0.001 | 0.507 ± 0.034 |
| Lao PDR    | 30  | 8  | 2  | 0.001 ± 0.001 | 0.129 ± 0.079 |
| Cambodia   | 40  | 62 | 6  | 0.021 ± 0.011 | 0.831 ± 0.024 |
| Philippines| 13  | 1  | 2  | 0.001 ± 0.001 | 0.513 ± 0.082 |
| Bangladesh | 11  | 67 | 5  | 0.043 ± 0.023 | 0.618 ± 0.164 |
| Sri Lanka  | 35  | 15 | 9  | 0.004 ± 0.002 | 0.717 ± 0.075 |
| All populations | 459 | 126 | 42 | 0.012 ± 0.001 | 0.671 ± 0.087 |

n = number, S = number of polymorphic sites, H = number of haplotypes, \( π \) = Unique haplotypes, \( π \) = nucleotide diversity, \( Hd \) = haplotype diversity.
Ie41 were detected only in Sri Lanka (Table 1, Fig. 1). The haplotype network analysis of the 459 CO1 sequences (42 haplotypes) were placed into five haplogroups, I–V based on mutation steps (ms) ≥ 10. Haplogroup I contained 27 haplotypes (Ie1 – Ie23, Ie25 – Ie27 and Ie42) from all isolates in Southeast Asia, haplogroup II contained nine haplotypes (Ie33 – Ie41) from Sri Lanka, haplogroup III contained haplotype Ie24 from four samples in Kampong Cham, Cambodia, haplogroup IV consisted of three haplotypes (Ie29, Ie31 and Ie32) from Bangladesh, and haplogroup V had haplotypes Ie28 and Ie30 from Bangladesh (Fig. 1). All haplotypes generated in this study combined with all retrieved sequences from GenBank were also used to construct a haplotype network, in which the genetic grouping was similar as indicated in Fig. 1 (Figure Supplementary Fig. 1).

### 3.3. Phylogenetic analyses

Based on previously studies, five clades (A – E) were classified and sequences were deposited in GenBank, i.e. clade A consisted of sequences from Nepal, clade B from Myanmar and Nepal, clade C from Lao PDR, clade D from India and Nepal, and clade E from wide range covering Sri Lanka, Oman, Nepal, Myanmar, Thailand, Indonesia, Malaysia, and the Philippines. Current phylogenetic tree analyses, combining our 42 haplotypes with 98 sequences from previous studies retrieved from GenBank, demonstrated there were five major lineages, clades A – E, supported by high bootstrap values and BI posterior probabilities (Fig. 2). Importantly, the PTP/bPTP analyses also supported the presence of five separate cryptic genetic groups corresponding to the five clades (A – E) classification in the phylogenetic tree (Supplementary Fig. 1). This was based on an acceptance rate of 0.61454; merge rate of 250,058; and split rate of 249,942. The estimated number of species was between 4 and 104 with a mean of 53.0 species. We subdivided clade E into two subclades, namely subclades E1 and E2 (Fig. 2). Of these subclades, E1 was the most common and widespread genetic group worldwide, whereas subclade E2 was specifically detected in Sri Lanka and more distantly in the Caribbean (Fig. 3). Our study revealed for the first time that several samples from Bangladesh belonged to clade C together with the sequences from Nepal and India. Almost all of the samples from Bangladesh in our study were found to belong to clade B together with the sequences from Myanmar and Nepal. Clade D was rarely found; only three deposited sequences from Lao PDR and four sequences recently examined from Cambodia clustered in this genetic group. We found that none of the sequences generated in our study belonged to clade A (Fig. 3).

### 4. Discussion

Our current study was conducted to examine the phylogenetic relationships of I. exustus that had been proposed by previous studies [5,10,17,19]. The phylogenetic analyses herein were based on more samples than previous studies, which also covered novel localities with a...
specific focus in South and Southeast Asia, especially Thailand. Genetic groups classified in our study were in concordance to all previous studies [10,17,19]. Clades A – E were initially classified corresponding to their geographical origin by Gauffre-Autelin et al. [10], clade A was found specifically in Nepal, clade B occurred in Myanmar and Nepal, clade C was specifically found in Lao PDR, clade D was found in India and Nepal, clade E occurred throughout a wide geographical range covering Sri Lanka, Oman, Nepal, Myanmar, Thailand, Indonesia, Malaysia, and the Philippines. Subsequent analyses proposed that the five major phylogenetic lineages should be classified into only four major lineages, namely clades A – D by the merged of clade C and D as a monophyletic group [19].

However, our study was based on haplotype networks and phylogenetic analyses, and it showed that *I. exustus* examined in our study can be classified into five haplogroups, belonging to four major evolutionary lineages being clade B – E. We found no samples belonged to clade A, which is congruent with previous report by Gauffre-Autelin et al. [10]. Furthermore, our data showed that clade E were subdivided into two subclades, namely E1 and E2, corresponding to a report by Gauffre-Autelin et al. [10]. All 35 samples from Sri Lanka examined in this study were grouped into subclade E2 (haplogroup II) together with a sequence from Sri Lanka [17] and two sequences from the French West Indies [19]. Our results showed that subclade E2 can confidently be classified as a novel cryptic genetic group, whereas, subclade E1 is the most common genetic group with a worldwide distribution. Furthermore, we discovered two and three novel haplotypes from Bangladesh

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**Fig. 2.** Phylogenetic tree analyses of CO1 sequences of 42 haplotypes of *Indoplanorbis exustus* sensu lato (sl), which were classified into five haplogroups (I – V) generated in this study, including 98 CO1 sequences of *I. exustus* deposited in GenBank under accession number HM104223, GU451738 – GU451750, KR811332 – KR811361, KY024420 – KY024472, and MH037074 – MH037075 based on maximum likelihood (ML) and Bayesian inference (BI). The tree of *I. exustus* were classified into five major lineages, i.e. clade A – E, whereas clade E were separated into subclades E1 and E2. Nodal supports are sequential values of bootstrap value generated by ML, and Bayesian posterior probability generated by BI. A CO1 sequence of *Bulinus tropicus*, accession number AY282583, was used as an out-group.
grouped within clade B and C, respectively.

Previously samples from Lao PDR have been classified as an independent clade D [10]. This was, however, was based on a low sample size of three sequences from Lao PDR (rare genotypes) which may not reflect actual phylogenetic relationships. Subsequently, further analyses by Mouahid et al. [19] of samples from Bangladesh, India and Nepal provided evidence to merge Clade D from Lao PDR into Clade C. Our results now provide the basis to separate clade C [19] into two different clades, i.e. clades C and D corresponding to the previous classification of Gaufrère-Autelin et al. [10]. Clades C and D classification was also supported by the PTP species delimitation test. Furthermore, we discovered that a unique haplotype of four snails from a novel place in Cambodia clustered within clade D together with previously generated sequences from Lao PDR. This suggests that this genetic group may be specifically genetically adapted and restricted within these geographical areas. Additionally, I. exustus from Philippines were sampled from limited localities, hence, only genotypes that belong to a common genetic group were detected, for instance, subclade E1. Other cryptic genetic groups/genotypes may also exist. Therefore, it is important to examine the extent of genetic variation at micro-geographical scales, covering a far broader range between these areas than has been done to date. Therefore, it is important to examine the extent of genetic variation at micro-geographical scales, covering a far broader range between these areas than has been done previously.

Hence, we also examined the genetic variation of I. exustus on a micro-geographical scale covering 33 localities within Thailand, together with three localities in Lao PDR and four in Cambodia. In Thailand, we found that all 21 haplotypes belonged to subclade E1, the most common genetic group distributed worldwide. However, there were several unique haplotypes found in particular localities, leading to significant genetic differences between many localities. The high self-fertilization capacity of I. exustus [2,7] may have facilitated clonal production of new genetic types adapted to each environment, leading to the unique haplotypes found in several different areas evidenced in our study. The different catchments in Thailand may also influence genetic differences and genetic structure of I. exustus, as water flow is limited or not possible between different catchments. This would lead to limited migration and low gene flow between the I. exustus populations across these catchments. For instance, in northeastern Thailand, the sampling locality Nong Khai (NKI) is in the Lower Mekong catchment, Maha Sarakham (MKM) is located in the Chi catchment, and Buri Ram (BRI) is located in Mun catchment. Our results showed that there were significant genetic differences between isolates from these different catchments in northeast Thailand compared to other isolates examined. Similar patterns of genetic differences have previously reported in the other freshwater species of snails, e.g. Bithynia siamensis [30] and Hydrobioides nassa [3].

The differences and variation in seasonal rainfall patterns and different ecological conditions between different regions of Thailand may also influence the genetic differences we have detected in I. exustus. For instance, significant genetic differences were detected when isolates from the following geographical localities in Thailand were compared to all other isolates examined, namely, isolates from Pichit (PCT), Ayuthaya (AYA), and Supan Buri (SPB) located in the alluvial plain in the central region of Thailand, Lampang (LPG) from the large mountain ranges in northern Thailand, Surathani (SNI) in the peninsular of Thailand, Kanchanaburi (KRI) located in steep river valleys in western, Chonburi (CBI) nearby the Gulf of Thailand in the eastern region, and NKI, MKM and BRI are localities in the Khorat Plateau in northeast Thailand. Interestingly, we found several isolates from geographically distant localities within a defined catchment and isolates from different catchments which showed low or no genetic differences (i.e. a low Fst value). These results may have been due to human activities of trading aquatic plants on which I. exustus can be attached. This is one of main sources of migration and gene flow of I. exustus in Thailand and worldwide.

In conclusion, our mitochondrial CO1 sequence analyses of the phylogeographic genetic variation of I. exustus, which was based on a large sample size, validated the worldwide distribution determined by previous studies [10,17,19]. Currently, at least six genetic groups have been classified based on haplotype networks and phylogenetic analyses. Subclade E1, or haplogroup I, is the most common genetic group distributed globally. Subclade E2 or haplogroup II is a novel cryptic group identified and defined in our study, specifically found in Sri Lanka and recently shown to exist in the French West Indies [19].

Furthermore, we found that clade D occurs in southern Lao PDR and as well as Cambodia. We also found high genetic variation and genetic differences between I. exustus collected from different micro-geographical localities in Thailand. The uncovering of unique subclades and haplotypes confirms that I. exustus, found worldwide, is a species complex and should hence be referred to as ‘I. exustus sensu lato’.

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Declaration of Competing Interest
The authors declare that they have no competing interests.

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