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Controversy in the taxonomic evaluation of the Asian tree frog *Polypedates leucomystax* complex presents the challenging task of gaining insight into its biogeographical distribution and diversification. Here, we describe the dispersion and genetic relationship of these species in Thailand where we connect the population of the *P. leucomystax* complex of the Sunda Islands to the mainland population based on the mitochondrial cytochrome c oxidase subunit I (COI) gene, derived from 266 samples. Our maternal genealogy implies that there are four well-supported lineages in Thailand, consisting of Northern A (clade A: *Polypedates* sp.), Nan (clade B: *P. cf. impresus*), Southern (clade C: *P. cf. leucomystax*) and Northern B (clade D: *P. cf. megacephalus*), with Bayesian posterior probability >0.9. Phylogeny and haplotype networks indicate that clades A, B and D are sympatric. In contrast, clade C (*P. cf. leucomystax*) and clade D (*P. cf. megacephalus*) are genetically divergent due to the geographical barrier of the Isthmus of Kra, resulting in allopatric distribution. Climatic conditions, in particular rainfall, that differ on each side of the Isthmus of Kra may play an important role in limiting the immigration of both clades. For the within-populations of either clades C or D, there was no significant correlation between geographic and genetic distance by the isolation-by-distance test, indicating intraspecific gene flow of each clade. Population expansion occurred in clade C, whereas clade D showed a constant population. Taken together, the *P. leucomystax* complex in Southeast Asia may be diversified by climatic oscillation, leading to allopatric and/or sympatric speciation.
Phylogenetic relationships and genetic diversity of the *Polypedates leucomystax* complex in Thailand

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**Keywords** Amphibian, Allopatric, Demographic expansion, Evolution
INTRODUCTION

Southeast Asia is a hotspot of substantial genetic diversity of amphibians. Recent molecular phylogenetic analyses have disclosed many anuran lineages that contain cryptic species. Historically, complex changes in the region’s geology and climate (e.g., Pleistocene climatic oscillations) altered the topology and environmental conditions, resulting in an initial fragmentation of habitat. These mechanisms generated high species richness in the current period (Hall, 1998; Woodruff, 2010). Of interest to our research were the numerous frog species in Southeast Asia whose taxonomy is still controversial, such as Microhyla fissipes (Yuan et al., 2016), Staurois tuberilinguis (Matsui et al., 2007), Microhyla ornata (Matsui et al., 2005) and Polypedates leucomystax (Kuraishi et al., 2013; Rujirawan, Stuart & Aowphol, 2013). The clarification of ambiguous species is essential to better understand their speciation and diversification and their biogeography for conservation purposes.

The Asian tree frog, the Polypedates leucomystax (Gravenhorst, 1829) complex, is an Asian Rhacophoridae frog. These species are widely distributed in Southeast Asia, South China and India. In addition, this species has phenotypic plasticity and high adaptation to local environment, leading to its existence in diverse habitats such as forests and even buildings. These high levels of phenotypic plasticity present a great challenge for classification. Phylogenetic and taxonomic relationships of the P. leucomystax complex throughout Southeast Asia exhibit adaptive radiation (Kuraishi et al., 2013; Pan et al., 2013; Rujirawan, Stuart & Aowphol, 2013). At least six valid species, including P. braueri, P. leucomystax, P. macrotis, P. megacephalus, P. mutus and P. impresus, have been delimited from the P. leucomystax complex based on their morphology, advertisement calls and molecular data (Matsui, Seto & Utsunomiya, 1986; Brown
et al., 2010; Kuraishi et al., 2011; Kuraishi et al., 2013; Pan et al., 2013). Five species, *P. leucomystax*, *P. mutus*, *P. macroditis*, *P. colletti* and *P. megacephalus*, can be found in Thailand (Taylor, 1962; Heyer, 1971; Frost, 2013; Kuraishi et al., 2013; Pan et al., 2013; Rujirawan, Stuart & Aowphol, 2013). A study by Brown et al. (2010) indicated that much of the genetic divergence of the *P. leucomystax* complex was discovered in mainland rather than in insular populations distributed throughout thousands of islands of the Malay Archipelago, presumably resulting from range expansion mediated by transportation of agricultural products. Recently a new species, *P. discantus*, belonging to the *P. leucomystax* species complex from southern Thailand was discovered using data on morphological characteristics, advertisement calls and molecular evidence, which were dominantly dissimilar to those of *P. leucomystax* and *P. megacephalus* (Rujirawan, Stuart & Aowphol, 2013). Several studies have confirmed the presence of highly cryptic species of the *Polypedates leucomystax* complex (Matsui, Seto & Utsunomiya, 1986; Kuraishi et al., 2011; Blair et al., 2013; Kuraishi et al., 2013; Pan et al., 2013). We believe that the *P. leucomystax* complex in Thailand remains a highly cryptic species, as is the case elsewhere.

Understanding the phylogenetic relationships among species can give insight into how lineages diverged and how new species arose. The process of speciation can be organized based on the geographic overlap of emerging species during divergence. In this study, we investigated the genetic variation, phylogenetic relationships and other relevant factors that limit the dispersal of *P. leucomystax* complex in Thailand. The present results illustrate the range of distribution of *P. leucomystax* and *P. megacephalus*, which is influenced by climatic conditions, and the possible existence of *P. impresus* in sympatry with *P. megacephalus* in the north of Thailand.
MATERIALS AND METHODS

Sample collection, DNA extraction and sequencing

In this study, a total of 266 adult *Polypedates leucomystax* complex individuals were collected from 15 different localities in Thailand (Table 1). All samples were dissected to obtain the liver, which was then stored in absolute ethanol. Collecting and enthustication was approved by Center For Animal Research Naresuan University under project number NU-AE591028. Genomic DNA was extracted from liver tissue using a DNA extraction kit (RBC Bioscience, Singapore) and kept at −20 °C for further use. Individual DNA was used as a template for PCR amplification of the mitochondrial COI gene using Taq DNA polymerase in a total volume of 25 µL under the following condition: an initial denaturation at 94 °C for 5 min, followed by 35–40 cycles at 94 °C for 30 s, 50 °C for 30 s and 72 °C for 1 min, and a final extension step at 72 °C for 7 min. PCR products were visualized on 1.5% agarose gel under UV light. The expected size of a partial mitochondrial COI gene sequence was 688 bp. Subsequently, all PCR products were purified using a QIAquick PCR Purification Kit (Qiagen, Germany) and then sequenced (Macrogen, South Korea).

Phylogeny

Phylogenetic reconstructions were executed using Bayesian inference (BI) and maximum likelihood (ML) independently. The best-fit model of DNA sequence evolution for this locus was identified with the Akaike information criterion (AIC) implemented in MrModeltest v2.3 (Nylander, 2004). The GTR+I+G model was selected as the best model and used in the following
A Bayesian tree was generated using MrBayes 3.1.2 (Ronquist & Huelsenbeck, 2003). For BI analysis, two independent searches with random starting trees were run for 5 million generations while sampling over 1,000 generations and compared using four Markov chain Monte Carlo (MCMC) chains (temp = 0.2). Convergence was assessed by plotting the log-likelihood scores in Tracer v1.5 (Rambaut et al., 2013), and the first 25% of the generations from each run were discarded before building a consensus tree. Maximum likelihood analysis was performed using RAxML 7.0.4 (Stamatakis, Hoover & Rougemont, 2008). The same model of nucleotide substitution used for BI analysis was employed for the ML tree search with 1,000 bootstrap replicates. COI gene sequences of the *Polypedates leucomystax* complex were retrieved from GenBank, as follows: *P. impresus*: KP996822 (China), KP996846 (China), KP087862-70 (Laos); *P. leucomystax*: KR087871-2 (Thailand); *P. megacephalus*: KR087879, KR087881 (Thailand).

**Population genetics and structure**

A total of 266 sequences of mitochondrial *COI* were aligned using ClustalW (implemented in MEGA 6.0 with default parameters). The number of polymorphic sites, the parsimony-informative sites, singleton sites, the number of haplotypes, haplotype diversity (*Hd*), and nucleotide diversity for each clade were calculated using DnaSP 5.0 (Librado & Rozas, 2009). Genetic distances among taxa were calculated using the *p*-distance model in MEGA 6.0 (Tamura et al., 2011). Furthermore, we detected a boundary line in the genetic landscape between *P. megacephalus* and *P. leucomystax* using Barrier 2.2 (Manni, Guérard & Heyer, 2004). The minimum spanning network was constructed using PopART (Population Analysis with
Reticulate Trees) population genetics software to define the relationships among haplotypes and
the distribution of haplotypes in each locality (Bandelt, Forster & Röhl, 1999). To evaluate the
effect of geographic distance on the genetic divergence among populations of *P. megacephalus*
and among populations of *P. leucomystax*, a Mantel test with 1,000 permutations was carried out
using GenAlEx 6.5 (Peakall & Smouse, 2012).

**Demographic history**

To investigate the demographic history of *P. megacephalus* and *P. leucomystax* populations in
Thailand, multiple approaches were explored. Neutrality tests of Tajima’s *D* and Fu’s *Fs* for the
two species were calculated using DnaSP 5.0. A significantly positive value indicates a process
of subdivision or recent population bottleneck, whereas a population expansion shows a
significantly negative value. Pairwise mismatch distribution was used to assume a constant
population size using DnaSP 5.0. Multiple mismatch distribution implies stability of the
population, while unimodal mismatch distribution reflects an expanding population. In addition
to these methods, the raggedness index (*r*<sub>g</sub>) of the observed distribution was calculated using
DnaSP 5.0. A small *r*<sub>g</sub> indicates a demographic expansion.

**RESULTS**

**Sequence characteristics**
A total of 266 samples of the *P. leucomystax* complex yielded 688 bp fragments of the mitochondrial COI gene. All new sequences in this study were deposited in the GenBank database (xxxx-xxxx) [available upon manuscript acceptance]. After multiple alignment of all COI sequences, the sequences were trimmed to the same length, given as 437 bp before downstream analysis. We observed 82 polymorphic sites, which are 82 parsimony-informative sites without a singleton site, acquiring 15 haplotypes (Table 2). Overall nucleotide and haplotype diversity were 0.0664 and 0.9000, respectively (Table 2).

Phylogenetic analyses and haplotype distribution

Based on 266 mitochondrial COI sequences of the *P. leucomystax* complex, matrilineal genealogy was analyzed through a Bayesian analysis model with MrModeltest v2.3. Our results indicated that the *P. leucomystax* complex in Thailand could consist of four clades, including: clade A (Northern A), *Polypedates* sp.; clade B (Nan), *P. cf. impresus*; clade C (Southern), *P. cf. leucomystax*; and clade D (Northern B), *P. cf. megacephalus* (Fig. 1). With respect to phylogenetic interference, *Polypedates* sp. was treated as a sister group of *P. cf. impresus*, which was found in Nan. *Polypedates* sp., however, can be seen in genetic samples obtained from Kanchanaburi (KCB), Mae Hong Son (MHS) and Phetchaburi (PCB) provinces, and shared a habitat with clade D, which was recognized as *P. cf. megacephalus*; its distribution range was in the far north of the Isthmus of Kra as Chiang Mai (CM), MHS, KCB, PCB, Saraburi (SRB), Loei (LPR), Nakhon Ratchasima (NRS) and Prachuap Khiri Khan (PKK), whereas the dispersal areas of the clade C population as *P. cf. leucomystax*, including Chumphon (CP), Nakhon Si Thammarat (NST), Phuket (PK) and Ranong (RN), were south of the Isthmus of Kra (Fig. 2). To
determine a barrier for immigration between *P. megacephalus* and *P. leucomystax* based on the dataset of genetic distance (Kimura’s two-parameter model), Barrier 2.2 was ed. Likely, the Isthmus of Kra could be a significant area to restrict their immigration.

The minimum spanning network among the mitochondrial haplotypes was also constructed as a result of four groups having similar results of phylogenetic inference (Fig. 3). The haplotypes of each group showed unique features, and each group had a different number of haplotypes. *P. megacephalus* exhibited the highest number of haplotypes at seven (*Hd* = 0.746), followed by *P. leucomystax* with five haplotypes (*Hd* = 0.7526), as shown in Table 2.

Haplotypes A and B, seen in *Polypedates* sp., and haplotype C found in *P. cf. impresus* were unique haplotypes. Haplotypes D–I were noted in *P. cf. megacephalus*, while populations of northern, western and upper southern Thailand (KCB, PCB and PKK, respectively) shared haplotype J. Besides, we noted that the haplotypes G and F of *P. cf. megacephalus* in NRS had relatively high divergence from the congeners. *P. cf. leucomystax* had high haplotype diversity, and contained three unique haplotypes (M, N and O) and two shared haplotypes (K and L).

In addition, there was no significant isolation-by-distance effect among populations of *P. cf. megacephalus* and *P. cf. leucomystax* based on analyses of Mantel tests between the genetic distance of the mitochondrial COI gene sequence and the geographical distance (Fig. 4).

Demographic history

When we defined a significant barrier around the Isthmus of Kra leading to the genetic divergence between *P. megacephalus* and *P. leucomystax*, neutrality tests (Tajima’s *D* and Fu’s
Fs) of both species were not significantly positive, whereas Fu’s Fs of *P. leucomystax* was significantly positive (Table 3). Furthermore, the mismatch distribution was tested as a result of left-skewed multimodal mismatch distribution for *P. megacephalus* with moderate \( r_g \) (0.2031) but unimodal mismatch distribution for *P. leucomystax* with low \( r_g \) (0.0569) (Fig. 5). Overall, these results suggested a constant population size of *P. megacephalus* and population expansion of *P. leucomystax*.

**DISCUSSION**

The taxonomy of the Asian tree frog of the *Polypedates leucomystax* complex, one of the most notoriously challenging, is contentious owing to the species’ widespread distribution and their similar morphology. To solve the taxonomic status of the *P. leucomystax* complex in Thailand, COI mitochondrial gene sequences of these species were analyzed. Unfortunately, several studies on the genealogy of the *P. leucomystax* complex used mitochondrial 12S rRNA, tRNA valine, and the 16S rRNA gene (Brown et al., 2010; Kuraishi et al., 2013; Pan et al., 2013). We, therefore, could not exploit these sequences for our analyses. Our matrilineal genealogy implied four well-supported lineages, consisting of *P. cf. megacephalus* (Northern B clade), *P. cf. leucomystax* (Southern clade), *P. cf. impresus* (found only in Nan) and *Polypedates* sp. (Northern A clade). That *Polypedates* sp. occurs in the same geographic areas as *P. cf. megacephalus* was considered as sympatric distribution; it, however, is a sister lineage with *P. cf. impresus*. Nevertheless, to clarify the taxonomy of *Polypedates* spp., morphological observations, call advertisement and ecological habitat would need to be investigated in future work. Recently, Pan et al. (2013) validated *P. impresus* molecularly, based on approximately 2 kb of mitochondrial
12S rRNA, tRNA valine and the 16S rRNA gene, as a valid species found in southern China and northern Laos. This is the first report to verify the existence of *P. impresus* in Nan, Thailand, which is located at the border with northern Laos, indicating its dispersal range. The main characteristics of *P. impresus* originally described by Yang (2008) are: the top of the head obviously concave, upper lip margin white, dorsal body light brown, and with no zebra-like stripes. The putative *P. impresus* individuals in our study are consistent with these characteristics.

Bayesian inference strongly supported that the northern and southern clades were considered as *P. megacephalus* and *P. leucomystax*, respectively, indicating geographically distinct species. Furthermore, Monmonier’s algorithm suggested that the Isthmus of Kra seemed to be a significant region separating them due to a large genetic divergence between the populations north and south of the isthmus. This result was in accordance with a previous study by Kuraishi et al. (2013). In addition, we noted that the majority of the populations of the *P. leucomystax* complex in Thailand are *P. megacephalus* and *P. leucomystax*. They are widespread species, with the former having a dispersal range from the south of China (Guangxi and Yunnan) to the north of Thailand, Laos and Vietnam (Kuraishi et al., 2013; Pan et al., 2013), and the latter having a distribution range from the south of Thailand to the Malay Archipelago (Brown et al., 2010; Kuraishi et al., 2013). Considering that the Isthmus of Kra shapes the spatial distribution of the genetic lineage between the northern and southern clades, it seems remarkable that it lacks substantial topological features, e.g. a mountain range or river, which usually block migration routes of animals. Yet the Isthmus of Kra serves as a geographic barrier between the two clades, with *P. leucomystax* absent from areas north of the isthmus and, likewise, *P. megacephalus* not distributed south of the isthmus. Moreover, the matrilineal haplotype network was strong
evidence of the suppression of migration of the northern and southern clades. In contrast, a previous study revealed the range expansion of insular populations of *P. leucomystax* with small genetic distance among islands of the Malay Archipelago, as they can migrate from one island to another mediated by anthropogenic effects, in particular transportation of agricultural products (Brown et al., 2010). We therefore postulated that climatic conditions might be the key barrier for restricting the distribution range of the modern populations, because the Isthmus of Kra is a climatic joint between a drier climate to the north and a more humid climate to the south of the isthmus. Data on rainfall in Thailand by the Thai Meteorological Department indicates a difference in the amount of rainfall between areas to the north and south of the isthmus (Fig. 6). Todd et al. (2011) observed a shift in reproductive timing in ten amphibian species at a wetland in South Carolina, USA, over 30 years as a result of climate change. Furthermore, the variation of breeding season length of the *P. leucomystax* complex of Thailand (putative *P. megacephalus*; 6 months a year for breeding) and Singapore (putative *P. leucomystax*; every month) led to adaptation to shortened breeding season length by increasing clutch size (Sheridan, 2009). It is clear that climate can influence the alteration in phenotypes of frogs (Sheridan, 2009; Todd et al., 2011). The different climates of the northern and southern clades’ habitats likely led to the alteration of biological features such as behavior, reproductive timing or specific niche, contributing to the restricted distribution of both species. However, an initial cladogenesis of the *P. leucomystax* complex in Indochina, which includes the two species, was caused by climatic oscillations during the Miocene and the subsequent Plio–Pleistocene, resulting in increased aridity and a monsoonal weather system, sea fluctuation and habitat fragmentation which in turn promoted species diversification (Blair et al., 2013). Kuraishi et al. (2013) inferred that the most recent divergence time between *P. megacephalus* and *P. leucomystax* was in the late Pliocene or
early Pleistocene (1.4–4.0 MYBP). The speciation process was caused by a short disjunction of
the Malay Peninsula and Sunda Islands by the South China Sea in the early Pliocene (about 5
MYBP) when the sea level rose (Hall, 1998), resulting in genetic divergence into the two valid
species.

Within the population of *P. megacephalus*, maternal genealogy demonstrated that the
genetic samples from Nakhon Ratchasima province (NRS) seemed to be a naturally occurring
divergence because of the emergence of endemic haplotypes; however, it was a low-supported
lineage, with 0.7 Bayesian posterior probability (BPP). When we considered the topography of
this region, the population of NRS as clade D1 is partitioned from the other populations within
clade D by the two mountain ranges, Dong Phaya Yen and Sankamphaeng. This may be a
possible barrier to gene flow among the modern populations of *P. megacephalus* between
western and eastern Thailand. Unfortunately, only a limited number of populations from the east
of Thailand were investigated in this study. Further work for validating if Dong Phaya Yen and
Sankamphaeng are a great barrier for gene flow in *P. megacephalus* is required. According to the
demographic history, the population of *P. megacephalus* in Thailand was a stable population but
the population of *P. leucomystax* in southern Thailand showed a relatively similar unimodal
distribution with small raggedness index, possibly indicating a population expansion. This result
was similar to that observed in the population of *P. leucomystax* in the northern Philippines
(Brown et al., 2010). This scenario implied a genetically homogenous population, especially in
the population of Phuket Island which shared a haplotype with NST. We believe that these
events might be mediated by human activities like agricultural transportation (Brown et al.,
2010). Although the population of *P. leucomystax* expanded, it was limited to localities south of
the Isthmus of Kra.
CONCLUSIONS

Our matrilineal genealogy of the *Polypedates leucomystax* complex in Thailand suggested four lineages, i.e. Nan (putative *P. impresus*), Northern B (putative *P. megacephalus*), Southern (putative *P. leucomystax*) and Northern A (*Polypedates* sp.) clades. We noted that the populations of the Northern B clade, Nan and *Polypedates* sp. are in sympatry while their distributions are allopatric to the southern clade (*P. leucomystax*), separated by the Isthmus of Kra. Climatic conditions may be a major contributor to limited migration of the current populations of both clades but climatic oscillation in the Pliocene and Pleistocene is a highly possible scenario that drove a speciation mechanism for diversification of the *P. leucomystax* complex in Southeast Asia and China, the divergence of the southern and northern clades in Thailand included.

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Author Contributions
K.B. designed the experiments, analyzed the statistical data and wrote the manuscript. C.S. conducted the experiments on *P. leucomystax* complex samples. Both authors read and approved the final manuscript.

REFERENCES

Bandelt H-J, Forster P, Röhl A. 1999. Median-joining networks for inferring intraspecific phylogenies. *Molecular Biology and Evolution* **16**:37–48.

Blair C, Davy CM, Ngo A, Orlov NL, Shi HT, Lu SQ, Gao L, Rao DQ, Murphy RW. 2013. Genealogy and demographic history of a widespread amphibian throughout Indochina. *Journal of Heredity* **104**:72–85.

Brown RM, Linkem CW, Siler CD, Sukumaran J, Esselstyn JA, Diesmos AC, Iskandar DT, Bickford D, Evans BJ, McGuire JA, Grismer L, Supriatna J, Andayani N. 2010. Phylogeography and historical demography of *Polypedates leucomystax* in the islands of Indonesia and the Philippines: evidence for recent human-mediated range expansion? *Molecular Phylogenetics and Evolution* **57**:598–619.

Frost DR. 2013. Amphibian species of the world: an online reference. Version 6.0. New York: American Museum of Natural History. [electronic database] Available at http://research.amnh.org/herpetology/amphibia/index.html (accessed 22 June 2017).

Hall R. 1998. The plate tectonics of Cenozoic SE Asia and the distribution of land and sea. In: Hall R, Holloway JD, eds. *Biogeography and geological evolution of SE Asia*. Leiden, The Netherlands: Backhuys Publishers, 99–131.
Heyer WR. 1971. Mating calls of some frogs from Thailand. Chicago: Field Museum of Natural History.

Kuraishi N, Matsui M, Hamidy A, Belabut DM, Ahmad N, Panha S, Sudin A, Yong HS, Jiang JP, Ota H, Thong HT, Nishikawa K. 2013. Phylogenetic and taxonomic relationships of the Polypedates leucomystax complex (Amphibia). Zoologica Scripta 42:54–70.

Kuraishi N, Matsui M, Ota H, Chen S-L. 2011. Specific separation of Polypedates braueri (Vogt, 1911) from P. megacephalus (Hallowell, 1861) (Amphibia: Anura: Rhacophoridae). Zootaxa 2744:53–61.

Librado P, Rozas J. 2009. DnaSP v5: a software for comprehensive analysis of DNA polymorphism data. Bioinformatics 25:1451–1452.

Manni F, Guérard E, Heyer E. 2004. Geographic patterns of (genetic, morphologic, linguistic) variation: how barriers can be detected by using Monmonier’s algorithm. Human Biology 76:173–190.

Matsui M, Ito H, Shimada T, Ota H, Saidapur SK, Khonsue W, Tanaka-Ueno T, Wu GF. 2005. Taxonomic relationships within the pan-oriental narrow-mouth toad Microhyla ornata as revealed by mtDNA analysis (Amphibia, Anura, Microhylidae). Zoological Science 22:489–495.

Matsui M, Mohamed M, Shimada T, Sudin A. 2007. Resurrection of Staurois parvus from S. tuberilinguis from Borneo (Amphibia, Ranidae). Zoological Science 24:101–106.

Matsui M, Seto T, Utsunomiya T. 1986. Acoustic and karyotypic evidence for specific separation of Polypedates megacephalus from P. leucomystax. Journal of Herpetology 20:483–489.
Nylander JA. 2004. MrModeltest v2.3. [program distributed by the author] Uppsala, Sweden: Evolutionary Biology Centre, Uppsala University.

Pan S, Dang N, Wang J, Zheng Y, Rao D, Li J. 2013. Molecular phylogeny supports the validity of *Polypedates impresus* Yang 2008. *Asian Herpetological Research* 4:124–133.

Peakall R, Smouse PE. 2012. GenAlEx 6.5: genetic analysis in Excel. Population genetic software for teaching and research—an update. *Bioinformatics* 28:2537–2539.

Rambaut A, Suchard MA, Xie D, Drummond AJ. 2013. Tracer v1.5. Available at http://beast.bio.ed.ac.uk/Tracer (accessed 22 June 2017).

Ronquist F, Huelsenbeck JP. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* 19:1572–1574.

Rujirawan A, Stuart BL, Aowphol A. 2013. A new tree frog in the genus *Polypedates* (Anura: Rhacophoridae) from southern Thailand. *Zootaxa* 3702:545–565.

Sheridan JA. 2009. Reproductive variation corresponding to breeding season length in three tropical frog species. *Journal of Tropical Ecology* 25:583–592.

Stamatakis A, Hoover P, Rougemont J. 2008. A rapid bootstrap algorithm for the RAxML web servers. *Systematic Biology* 57:758–771.

Tamura K, Stecher G, Peterson D, Filipski A, Kumar S. 2013. MEGA6: Molecular Evolutionary Genetics Analysis version 6.0. *Molecular Biology and Evolution* 30:2725–2729.

Taylor EH. 1962. The amphibian fauna of Thailand. *University of Kansas Science Bulletin* 43:265–599.
Todd BD, Scott DE, Pechmann JH, Gibbons JW. 2011. Climate change correlates with rapid
delays and advancements in reproductive timing in an amphibian community.
*Proceedings of the Royal Society B: Biological Sciences* **278**:2191–2197.

Woodruff DS. 2010. Biogeography and conservation in Southeast Asia: how 2.7 million years
of repeated environmental fluctuations affect today’s patterns and the future of the
remaining refugial-phase biodiversity. *Biodiversity and Conservation* **19**:919–941.

Yuan Z-Y, Suwannapoom C, Yan F, Poyarkov NA Jr, Nguyen SN, Chen H-m, Chomdej S,
Murphy RW, Che J. 2016. Red River barrier and Pleistocene climatic fluctuations
shaped the genetic structure of *Microhyla fissipes* complex (Anura: Microhylidae) in
southern China and Indochina. *Current Zoology* **62**:531–543.
Phylogeographic relationships of the *Polypedates leucomystax* complex among populations in Thailand

It consists of clade A as *Polypedates* sp., clade B as *P. cf. impresus*, clade C as *P. cf. leucomystax*, and clade D as *P. cf. megacephalus*, as well as outgroups (KR087858, KP996762 = *P. braueri*) inferred from Bayesian analysis of mitochondrial *COI* gene sequences. Bayesian posterior probability values are expressed above internodes. The asterisks above branches represent bootstrap support for Bayesian posterior probabilities and maximum likelihood (>95%). Scale bar represents 0.5 nucleotide substitutions per site.
Clade A  Northern A; *Polypedates* sp.

Clade B  Nan; *P. cf. impresus*

Clade C  Southern part; *P. cf. leucomystax*

*P. leucomystax* complex

Clade D  Northern B; *P. cf. megacephalus*
Figure 2 (on next page)

Haplotype distribution of the *Polypedates leucomystax* complex throughout Thailand.

The abbreviations for each locality are given in Table 1. Different colors represent the different haplotypes.
The minimum spanning network of all haplotypes found in the *Polypedates leucomystax* complex in Thailand.

The mutation points between haplotypes are expressed by hatch marks. The different colors indicate the localities where the samples were collected.
The correlation of genetic distance and linear geographic distance (km) for *Polypedates megacephalus* (A) and *P. leucomystax* (B).
A

y = 1E-05x + 0.0037

R² = 0.0855, p = 0.11

B

y = 1E-05x + 0.0047

R² = 0.0618, p = 0.05

y = 1E-05x + 0.0037

R² = 0.0855, p = 0.11
Mismatch distribution of the mitochondrial COI gene in *Polypedates megacephalus* (A) and *P. leucomystax* (B).

The raggedness \( (r_g) \) index is calculated to evaluate the population expansion of each species. Ramos-Onsins and Rozas’s \( R^2 \) statistic represents the population growth.
A

$r_g$ index = 0.2031
$R^2$ statistic = 0.1063

B

$r_g$ index = 0.0569
$R^2$ statistic = 0.149
Figure 6 (on next page)

The relationship between climatic condition and genetic differentiation across *Polypedates megacephalus* and *Polypedates leucomystax*.

(A) annual rainfall (in mm) in Thailand for 2015 (Image credit: Thai Meteorological Department); and (B) a significant barrier to partition the distribution of the Northern B (*P. megacephalus*) and Southern (*P. leucomystax*) clades, by Barrier version 2.2.
Rainfall

Millimeter (mm)

A

Isthmus of Kra

B

Dry winter

Short dry winter

Rainfall

Millimeter (mm)

Isthmus of Kra

Dry winter

Short dry winter
**Table 1** (on next page)

Localities of sample collection for *Polypedates leucomystax* complex in Thailand.
Table 1 Localities of sample collection for *Polypedates leucomystax* complex in Thailand.

| Locality                          | Abbreviation | Number | Altitude (m above sea level) | Longitude  | Latitude   |
|-----------------------------------|--------------|--------|-----------------------------|------------|------------|
| Nan province                      | NAN          | 12     | 665                         | 18.980974  | 101.182594 |
| Kanchanaburi province             | KCB          | 20     | 917                         | 14.69329   | 98.40535   |
| Loei province: Phu Ruea           | LPR          | 11     | 939                         | 17.48193   | 101.34982  |
| Nakhon Ratchasima province        | NRS          | 14     | 865                         | 14.49336   | 101.87364  |
| Chiang Mai province: Mae Wang     | CM           | 7      | 678                         | 18.657305  | 98.681831  |
| Chiang Mai province: Doi Saket    | CM           | 13     | 402                         | 18.98777   | 99.11455   |
| Chiang Mai province: Omkoi        | CM           | 13     | 460                         | 17.47137   | 98.45785   |
| Mae Hong Son province             | MHS          | 44     | 396                         | 19.24797   | 97.99542   |
| Saraburi province                 | SRB          | 12     | 105                         | 14.70993   | 100.81819  |
| Phetchaburi province              | PCB          | 22     | 329                         | 14.70993   | 100.81819  |
| Prachuap Khiri Khan province      | PKK          | 15     | 23                          | 11.43678   | 99.56011   |
| Ranong province                   | RN           | 14     | 18                          | 9.6052     | 98.4669    |
| Nakhon Si Thammarat province      | NST          | 37     | 97                          | 8.76902    | 99.80349   |
| Phuket province: Thalang          | PK           | 17     | 31                          | 7.96804    | 98.33589   |
| Chumphon province                 | CP           | 15     | 103                         | 10.110278  | 99.082778  |
Table 2

Summary of the *P. leucomyrstax* complex in Thailand

major lineages clades, putative scientific name, number of individuals (*N*), number of mtDNA haplotypes (*n*), number of polymorphic sites (*P*), parsimony-informative sites (*PI*) and singleton sites (*S*), haplotype diversity (*Hd*) and nucleotide diversity (*π*).
Table 2 Summary of the *P. leucomystax* complex in Thailand: major lineages clades, putative scientific name, number of individuals (*N*), number of mtDNA haplotypes (*n*), number of polymorphic sites (P), parsimony-informative sites (PI) and singleton sites (S), haplotype diversity (*H_d*) and nucleotide diversity (*π*).

| Clade | Scientific name     | *N* | *n* | *π*   | *H_d* | P  | S  | PI |
|-------|---------------------|-----|-----|-------|-------|----|----|----|
| A     | Polypedates sp.     | 40  | 2   | 0.0037| 0.4089| 4  | 0  | 4  |
| B     | *P. impresus*       | 12  | 1   | 0     | 0     | 0  | 0  | 0  |
| C     | *P. megacephalus*   | 131 | 7   | 0.0048| 0.746 | 15 | 1  | 14 |
| D     | *P. leucomystax*    | 83  | 5   | 0.0073| 0.7526| 7  | 0  | 7  |
| Total |                     | 266 | 15  | 0.0664| 0.9   | 82 | 0  | 82 |
Table 3 (on next page)

Summary of statistics used to compute the demographic history of populations of *P. megacephalus* and *P. leucomystax*. 
Table 3 Summary of statistics used to compute the demographic history of populations of *P. megacephalus* and *P. leucomystax*.

| Species             | Tajima’s $D$ | Fu’s $F_s$ | $P$ value |
|---------------------|--------------|------------|-----------|
| *P. megacephalus*   | 0.439        | >0.1       | 3.213     | 0.045     |
| *P. leucomystax*    | 1.176        | >0.1       | 3.031     | 0.071     |