Molecular and morphological evidence reveals a new genus of the subfamily Heteropterinae (Lepidoptera, Hesperiidae) from China

Yongxiang Hou¹, Hideyuki Chiba², Lijuan Zhu¹, Zhou Chang³, Lijun Ma⁴, Siyao Huang¹, Min Wang¹, Xiaoling Fan¹

¹ Department of Entomology, College of Plant Protection, South China Agricultural University, Guangzhou, Guangdong 510642, China ² B. P. Bishop Museum, 1525 Bernice Street, Honolulu, Hawaii, 96817-0916, USA ³ State Key Laboratory of Genetic Resources and Evolution, Kunming Institute of Zoology, Chinese Academy of Sciences, Kunming, Yunnan 650201, China ⁴ Institute of Plant and Environmental Protection, Beijing Academy of Agriculture and Forestry Sciences, Beijing 100097, China

Corresponding authors: Min Wang (minwang@scau.edu.cn); Xiaoling Fan (fanxiaol66@scau.edu.cn)

Academic editor: Martin Wiemers  |  Received 14 May 2021  |  Accepted 6 July 2021  |  Published 5 August 2021

http://zoobank.org/3C184FA2-423E-43F5-B77A-5D06C5639245

Citation: Hou Y, Chiba H, Zhu L, Chang Z, Ma L, Huang S, Wang M, Fan X (2021) Molecular and morphological evidence reveals a new genus of the subfamily Heteropterinae (Lepidoptera, Hesperiidae) from China. ZooKeys 1055: 55–67. https://doi.org/10.3897/zookeys.1055.68640

Abstract

Molecular phylogenetic analysis indicates that the genus Carterocephalus is not monophyletic. Based on combined molecular and morphological evidence, we propose a new genus, Pulchroptera Hou, Fan & Chiba, gen. nov., for Pamphila pulchra Leech, 1891. The adult, wing venation, and male genitalia of Pulchroptera pulchra comb. nov., Carterocephalus palaemon, and related genera are illustrated.

Keywords

Carterocephalus, Pulchroptera, new combination

Introduction

In recent years, the molecular phylogeny of the family Hesperiidae has attracted the attention of an increasing number of researchers (Warren et al. 2008, 2009; Sahoo et al. 2016; Toussaint et al. 2018; Cong et al. 2019; Li et al. 2019; Liu et al. 2020). At the
subfamily level, however, the phylogeny of the family Hesperiidae has yet to be established, and multiple new subfamilies (Zhang et al. 2019, 2020) and genera (Fan et al. 2016; Huang et al. 2016, 2019; Cong et al. 2019; Li et al. 2019) have been proposed in recent years.

Although Heteropterinae, including 13 genera from Africa, was established already by Aurivillius (1925), subsequent authors did not recognise the subfamily, presumably because it was a mixture of genera and species assigned to the subfamilies Hesperiinae as well as Heteropterinae in the current taxonomy. Evans, somehow, proposed three different genus group names for taxa of these skippers, namely the Astictopterus group for the African taxa (Evans 1937), the Heteropterus group for the European and Asian taxa (Evans 1949), and the Carterocephalus group for the American taxa (Evans 1955), all of which he considered a part of the subfamily Hesperiinae. This arrangement was accepted in subsequent taxonomic works until Higgins (1975) and Scott and Wright (1990) restored the subfamily Heteropterinae. Recent molecular studies strongly support the monophyly of the subfamily (Warren et al. 2008; Sahoo et al. 2016, 2017; Toussaint et al. 2018, 2020; Cong et al. 2019; Li et al. 2019; Zhang et al. 2019; Liu et al. 2020). Currently, Heteropterinae includes the following 13 genera: *Heteropterus* Dumeril, 1806; *Carterocephalus* Lederer, 1852; *Butleria* Kirby, 1871; *Argopteron*, 1893; *Dalla* Mabille, 1904; *Leptalina* Mabille, 1904; *Metisella* Hemming, 1934; *Dardarina* Evans, 1937; *Hovala* Evans, 1937; *Piruna* Evans, 1955; *Freemaniana* Warren, 2001; *Ladda* Grishin, 2019; and *Willema* Grishin, 2019. Moreover, the most recent studies indicate that all these genera are monophyletic (Cong et al. 2019; Toussaint et al. 2020).

The genus *Carterocephalus* includes more than 20 species distributed in the Holartic and Oriental regions. However, a cursory inspection of the male genitalia indicates that *C. pulchra* (Leech, 1891) is not a congener of the type species *Papilio palaemon* Pallas, 1771. Indeed, the findings of our morphological and molecular phylogenetic studies have revealed closer relationships with species in the genera *Heteropterus* and *Leptalina*. Accordingly, we consider that *Carterocephalus pulchra* should be placed in a new genus.

In the present study, we sought to assess the monophyly of the genus *Carterocephalus* and its relationship with other genera of Heteropterinae. On the basis of the evidence obtained, we describe a new genus.

**Materials and methods**

**Morphological examination**

For the morphological study, we followed the methods described by Fan et al. (2010). To examine wing venation, wings were removed from the thorax and cleaned with a 1:1 mixture of bleaching liquid (Blue Moon, Guangzhou, China) and water for approximately 3 to 4 min. Photographs of the wing venation and male genitalia were taken using a Keyence VHX-5000 digital microscope (Keyence, Osaka, Japan).
Taxon sampling

We sampled specimens from all genera listed in the subfamily Heteropterinae (Warren et al. 2008, 2009; Cong et al. 2019; Toussaint et al. 2020), including as many species as possible. We used a total of 44 specimens of 38 species in 13 genera as ingroup taxa, along with 12 species from other subfamilies (Coeliadinae, Pyrginae, Eudaminae, Euschemoninae, Barcinae, Trapezitinae, and Hesperiinae) as outgroup taxa. Among these specimens, 31 were newly sequenced in this study, with the remaining sequences being obtained from the GenBank database along with supplementary data presented by Sahoo et al. (2016) and Toussaint et al. (2020). The respective voucher specimens and additional information are listed in Suppl. material 1: Table S1. Vouchers bearing codes beginning with the abbreviation SCAU have been deposited in the collection of South China Agricultural University (SCAU), Guangzhou, China, and the specimens (JU19), (Dalla), and (SZSMETI) are retained in the private collections of J. Uehara, H. Chiba, and S. Sáfián, respectively.

Laboratory protocols

DNA was extracted from two or three legs of dried adult specimens using a TIANamp Genomic DNA Kit (Tiangen, Guangzhou, China) following the manufacturer's instructions. We amplified a single mitochondrial gene (658 bp of COI) and three nuclear genes (1066 bp of EF-1α, 610 bp of RPS5, and 403 bp of Wingless), for a total of 2737 bp. The primers used to amplify each gene were synthesised by Sangon Biotech (Shanghai, China) and are shown in Suppl. material 2: Table S2. DNA amplification was performed in 20-µL reaction volumes containing 1 µL of template DNA, 0.8 µL of each primer (10 µM), 10 µL of 2× EasyTaq PCR superMix (+dye) (Transgen, Beijing, China), and 7.4 µL of ddH₂O. The amplification protocol adopted is the one described by Huang et al. (2019). Sequencing of the amplicons thus obtained was performed by Sangon Biotech (Shanghai, China) and Tsingke Biological Technology (Beijing, China), and new sequences have been deposited in GenBank (Suppl. material 1: Table S1).

Phylogenetic analyses

Sequences were aligned using Clustal W (Thompson et al. 1997) and edited manually using MEGA 7.0 (Kumar et al. 2016). Gene data from Cong et al. (2019) were extracted from the genomic assembly in IDBA-UD (Peng et al. 2012). PartitionFinder v2.1.1 (Lanfear et al. 2012, 2016; Guindon et al. 2010) was used to select the optimal codon partitioning scheme under Akaike information criterion correction (AICc) (Suppl. material 3: Table S3). We inferred the phylogenetic trees using two methods, namely maximum likelihood (ML) and Bayesian inference (BI), for which we used the partition scheme produced by PartitionFinder. ML analyses were performed using IQ-TREE (Nguyen et al. 2015) as implemented in the IQ-TREE
web online server (iqtree.cibiv.univie.ac.at, Trifinopoulos et al. 2016), with branch support values evaluated based on 1000 replicates for ultrafast bootstrap (UFBoot) (Minh et al. 2013) and SH-aLRT (Guindon et al. 2010). BI analyses were performed using the CIPRES Science Gateway (https://www.phylo.org/) (Miller et al. 2010) with Markov Chain Monte Carlo (MCMC) randomisation in MrBayes using XSEDE 3.2.6 (Ronquist et al. 2012). Reversible-jump MCMC was used to facilitate sampling across the entire subduction rate model. We conducted two independent MCMC runs, with four Markov chains ($5 \times 10^6$ generations) for each analysis, of which the initial 25% of samples were discarded as burn-in. Bayesian posterior probabilities (PP) were used to evaluate branch support, and trees were visualised using FigTree v1.4.0.

Results and discussion

Phylogenetic relationships

The topological structures of the concatenated dataset inferred by ML and BI analyses were found to be generally consistent and strongly supported at most nodes (PP $\geq 0.98$, SH-aLRT $\geq 95$, UFBoot $\geq 98$) (Fig. 1). Moreover, the two analyses provided strong support for the monophyly of Heteropterinae (PP = 1, SH-aLRT = 99.9, UFBoot = 100), which excludes the genera Apostictopterus, Barca, Lepella, and Ttitana originally assigned to this subfamily, and is consistent with the findings of the most recent studies (Toussaint et al. 2018, 2020; Cong et al. 2019; Zhang et al. 2019). Within the subfamily Heteropterinae, four major clades were differentiated, with 14 well-supported monophyletic subclades, corresponding to the 13 currently recognised genera and the Carterocephalus pulchra clade. Certain results were consistent with those of previous studies (Cong et al. 2019; Toussaint et al. 2020): (1) of the 13 genera, 12 genera, excluding Carterocephalus, were monophyletic; (2) Argopteron and Butleria formed a strongly supported monophyletic group (PP = 1, SH-aLRT = 99.1, UFBoot = 100) that is sister to all other genera in Heteropterinae (PP = 1, SH-aLRT = 98.3, UFBoot = 99); (3) Carterocephalus, excluding the species C. pulchra, was sister to the clade containing Metisella, Hovala, and Willema with strong support (PP = 1, SH-aLRT = 99.5, UFBoot = 100); and (4) Piruna, Dardarina, Freemaniana, Ladda and Dalla formed a strongly supported monophyletic clade (PP = 1, SH-aLRT = 98.8, UFBoot = 99). Two findings, however, are inconsistent with those reported previously. Firstly, Piruna is sister to Dardarina (PP = 0.76, SH-aLRT = 87.2, UFBoot = 94), as opposed to sister to the four genera Dardarina, Freemaniana, Ladda, and Dalla. Based on the morphology of the male genitalia (Evans, 1955), Piruna shows a relatively close similarity to Dardarina, whereas species of Dalla show extensive variation. However, previous molecular phylogenetic studies, as well as our own, sampled only some representatives of Dalla. Accordingly, the monophyly of Dalla as well as the relationships among these five genera should be subjected to further studies.
A new genus of Heteropterinae from China

Secondly, we found that *Carterocephalus* is not a monophyletic group, given that the 11 species analysed in the present study were recovered in two distinct clades, with *C. pulchra* clustering with *Leptalina* and *Heteropterus* with strong support (PP = 1,

**Figure 1.** Maximum likelihood phylogenetic tree of the subfamily Heteropterinae. Values at nodes represent the posterior probabilities (PP) of BI analyses values, SH-aLRT values (SH-aLRT), and Ultrafast bootstrap support values (UFBoot) of the maximum likelihood analysis. * indicates that one of the values at a node exceeds the standard (PP ≥ 0.98, SH-aLRT ≥ 95, UFBoot ≥ 98). When the three node values all reach the standard, only one * is displayed. – indicates that the node was not recovered in the ML or BI tree.
The other ten species, including the type species *C. palaemon*, were recovered as a strongly supported monophyletic clade.

Although in this study we focused on relationships among the genera of Heteropterinae, it is worth mentioning that certain intra-generic relationships, namely, those between *C. abax* Oberthür, 1886 and *C. patra* Evans, 1939, *C. avanti* (de Nicéville, 1886) and *C. argyro stigma* (Eversmann, 1851), *C. longimaculatus* Hou, Fan & Chiba, 2021 and *C. alcina* Evans, 1939, *C. palaemon* (Pallas, 1771) and *C. silvicola* (Meigen, 1828) are strongly supported. As described by Toussaint et al. (2020), despite the lack of strong support (PP = 0.73, SH-aLRT = 85.3, UFBoot = 69), *C. houangty* and *C. dieckmanni* were clustered in a clade comprising *C. palaemon*, *C. silvicola*, *C. longimaculatus*, and *C. alcina*. In our previous study (Hou et al. 2021), we established that *C. dieckmanni* is sister to *C. abax* and *C. patra*. However, owing to an oversight, the names *C. dieckmanni* and *C. argyro stigma* were confused, which explains the discrepancy compared with the results reported herein. Accordingly, to determine relationships more comprehensively in the genus *Carterocephalus*, we ideally need to undertake additional and more extensive sampling.

Morphologically, although *C. pulchra* is similar to the type species of *Carterocephalus* with respect to wing shape and pattern (Fig. 2), the origin of vein R₄ on the hindwing is located nearly midway between the termen and the base in *C. pulchra*, *Heteropterus*, and *Leptalina*, whereas in other species of *Carterocephalus* the origin of vein R₄ is closer to the termen than to the base (Fig. 3). With regards to the male genitalia, the uncus in *C. pulchra*, *Heteropterus*, and *Leptalina* is deeply bifurcated, with arms distant from each other, whereas in the type species of *Carterocephalus* the uncus bifurcates with arms closely aligned (Fig. 4). These morphological similarities would accordingly appear to indicate that *C. pulchra* is more closely related to *Heteropterus* and *Leptalina* than to other species of *Carterocephalus*. Of these related genera, *C. pulchra* is autapomorphic with respect to its male genitalia. Notably, the gnathos is weakly sclerotized, membranous, and rounded at the tip, the valvae are asymmetrical, and the juxta is a heart-shaped ring with a narrow and long latero-central process. In summary, we propose a new genus, *Pulchroptera* Hou, Fan & Chiba gen. nov., for the *Carterocephalus pulchra* clade based on its autapomorphies and molecular evidence.

**Pulchroptera** Hou, Fan & Chiba, gen. nov. http://zoobank.org/3C184FA2-423E-43F5-B77A-5D06C5639245 Figures 2–4

**Type species.** *Pamphila pulchra* Leech, 1891

**Description.** Forewing length 11–12 mm. Antennae approximately half the length of forewing; nudum 8 on apiculus, dark brown. Palpi on second segment long and erect, yellow with long black hairs; on third segment black, thick, short, and porrect. Wing venation (Fig. 3): forewing: length of discoidal cell almost equal to 2/3 forewing length, Sc ends at 1/2 forewing length; origin of vein R₄ before vein R₃;
Figure 2. Male adults of the two skippers. Above: Pulchroptera pulchra (Leech, 1891) comb. nov. from Kunming, Yunnan, China; below: Carterocephalus palaemon (Pallas, 1771) from Moscow, Russia.

Figure 3. Wing venation of four genera of Heteropterinae A Pulchroptera pulchra (Leech, 1891) comb. nov. B Carterocephalus palaemon (Pallas, 1771) C Heteropterus morpheus (Pallas, 1771) D Leptalina unicolor (Bremer & Grey, 1852).

origin of vein $M_2$ in middle of veins $M_1$ and $M_3$; veins $CuA_1$, $CuA_2$, and 1A+2A almost parallel to each other; origin of vein $CuA_2$ nearly midway between vein $CuA_1$ and base. Hindwing: costa longer than dorsum; length of discoidal cell almost equal to 3/5 hindwing; origin of vein $Rs$ midway between base and termen; origin of vein $M_2$ slightly nearer $M_1$ than $M_3$. Wing ground colour and wing patterns: upper side dark
Figure 4. Male genitalia of four genera of Heteropterinae A *Pulchroptera pulchra* (Leech, 1891) comb. nov. B *Carterocephalus palaemon* (Pallas, 1771) C *Heteropterus morpheus* (Pallas, 1771) D *Leptalina unicolor* (Bremer & Grey, 1852).

brown with small yellow spots in central and submarginal areas; underside light brown, forewing patterns similar to upper side, hindwing with small silvery spots in spaces Rs, M$_3$, CuA$_1$, and CuA$_2$, and with a silvery longitudinal central streak. Mid and hind tibiae each with pair of spurs. Male genitalia: Tegument small and narrow, constricted at middle in dorsal view; uncus deeply bifurcated, V-shaped dorsally; gnathos long and wide, longer than tegument, membranous, undivided from basal 1/3; saccus long; valvae asymmetrical, bifid, distal end of left valva more sclerotized than right valva; aedeagus long, subzonal sheath shorter than suprazonal sheath, ratio of subzonal sheath to suprazonal sheath approximately 1:2, vesica with cornuti; juxta a heart-shaped ring with membranous extensions dorsally.

Remarks. The new genus superficially resembles *Carterocephalus* Lederer, 1852, although it is distinguishable from the latter with regards to the following characters: hindwing undersides with silver spots, a deeply bifurcated V-shaped uncus, juxta a heart-shaped ring, and valvae asymmetrical.

The new genus contains only the type species *Pulchroptera pulchra* (Leech, 1891) comb. nov., with the nominotypical subspecies and a further subspecies, *Pulchroptera pulchra ops* (Grum-Grushimaïlo, 1891) comb. nov. According to the description of Evans (1949), in *Pulchroptera pulchra pulchra* comb. nov. the upper side of the hindwing has a cell spot and the submarginal markings are notably more conspicuous, whereas in *Pulchroptera pulchra ops* comb. nov. the upper side of the hindwing lacks a cell spot
and has conspicuous submarginal markings. Whether the subspecies status of the latter is valid is subject to further verification.

**Etymology.** The name of the genus is taken from the specific epithet of the type species ‘pulchr-’, meaning beautiful, and ‘optera’, meaning wing. The gender is feminine.

**Distribution.**

*Pulchoptera pulchra pulchra* comb. nov.: China (Sichuan, Yunnan)

*Pulchoptera pulchra ops* comb. nov.: China (Gansu, Qinghai, Xizang)

**Acknowledgments**

This work was supported by the National Natural Science Foundation of China (no. 31872264, 31471984). We are grateful to Messes Jiro Uehara (Japan) and Szabolcs Sáfián (UK) for providing samples and the Cave Biology Laboratory of South China Agricultural University for taking the wing venation and genitalia photographs. We would like to thank Editage (www.editage.cn) for English language editing.

**References**

Abouheif E, Wray GA (2002) Evolution of the gene network underlying wing polyphenism in ants. Science 297: 249–252. [https://doi.org/10.1126/science.1071468](https://doi.org/10.1126/science.1071468)

Aurivillius C (1925) Die afrikanischen Tagfalter. In Seitz, Gross-schmett Erde 13: 11–613. [80 pls.]

Chou I (1994) Monographia Rhopalocerorum Sinensium. Henan Scientific and Technological Publishing House, Zhengzhou, 852 pp.

Cong Q, Zhang J, Shen JH, Grishin NV (2019) Fifty new genera of Hesperiidae (Lepidoptera). Insecta Mundi 0731: 1–56. [https://journals.fsvc.org/mundi/article/view/117773](https://journals.fsvc.org/mundi/article/view/117773)

Evans WH (1937) A Catalogue of the African Hesperiidae indicating the classification and nomenclature adopted in the British Museum (Natural History). British Museum (Natural History), London, 212 pp.

Evans WH (1949) A Catalogue of the Hesperiidae from Europe, Asia & Australia in the British Museum (Natural History). British Museum (Natural History). London, 502 pp. [https://www.biodiversitylibrary.org/page/48223970](https://www.biodiversitylibrary.org/page/48223970)

Evans WH (1955) A Catalogue of the American Hesperiidae Indicating the Classification and Nomenclature Adopted in the British Museum (Natural History). Part IV. Hesperiinae and Megathyminae. British Museum (Natural History), London, 499 pp.

Fan XL, Chiba H, Huang ZF, Fei W, Wang M, Sáfián S (2016) Clarification of the Phylogenetic Framework of the Tribe Baorini (Lepidoptera: Hesperiidae: Hesperiinae) Inferred from Multiple Gene Sequences. PLoS ONE 11(7): e0156861. [https://doi.org/10.1371/journal.pone.0156861](https://doi.org/10.1371/journal.pone.0156861)

Fan XL, Chiba H, Wang M (2010) The genus *Scobura* Elwes & Edwards, 1897 from China, with descriptions of two new species (Lepidoptera: Hesperiidae). Zootaxa 2490(1): 1–15. [https://doi.org/10.11646/zootaxa.2490.1.1](https://doi.org/10.11646/zootaxa.2490.1.1)
Folmer O, Black M, Hoeh W, Lutzet R, Vrijenhoek RC (1994) DNA primers for amplification of mitochondrial cytochrome oxidase subunit I from diverse metazoan invertebrates. Molecular marine biology and biotechnology 3(5): 294–299.

Grum-Grshimailo (1891) Lepidoptera nova in Asia centrali novissime lecta et descripta. Horae societatis entomologicae rossicae 25(3–4): e460. https://archive.org/details/trudyrusskagoent2526189092russ/page/n505/mode/1up?view=theater

Guindon S, Dufayard JF, Lefort V, Anisimova M, Hordijk W, Gascuel O (2010) New algorithms and methods to estimate maximum-likelihood phylogenies: assessing the performance of PhyML 3.0. Systematic biology 59(3): 307–321. https://doi.org/10.1093/sysbio/syq010

Häuser CL (1993) Die inneren weiblichen Genitalorgane der Tagfalter (Rhopalocera): Vergleichende Morphologie und phylogenetische Interpretation (Insecta, Lepidoptera). Zoologische Jahrbücher. Abteilung für Systematik, Ökologie und Geographie der Tiere 120: 389–439.

Higgins LG (1975) The Classification of European Butterflies. Collins, London, 320 pp.

Hou YX, Li YF, Wang M, Chiba H, Fan XL (2021) The relationships of *Carterocephalus alcinna* Evans, 1939 and relatives (Lepidoptera: Hesperiidae: Heteropterinae). Journal of South China Agricultural University 42(1): 101–108. http://doi.org/10.7671/j.issn.1001-411X.202009046

Huang ZF, Chiba H, Jin J, Kizhakke AG, Wang M, Kunte K, Fan XL (2019) A multilocus phylogenetic framework of the tribe Aeromachini (Lepidoptera: Hesperiidae: Hesperiinae), with implications for taxonomy and biogeography. Systematic Entomology 44: 163–178. https://doi.org/10.1111/syen.12322

Huang ZF, Chiba H, Wang M, Fan XL (2016) Establishment of a new genus for *Abraximorpha heringi* and *A. pieridoides* (Lepidoptera: Hesperiidae: Pyrginae: Tagiadiini). Zootaxa 4173(2): 147–154. https://doi.org/10.11646/zootaxa.4173.2.5

Kumar S, Stecher G, Tamura K (2016) MEGA7: Molecular Evolutionary Genetics Analysis version 7.0 for bigger datasets. Molecular Biology and Evolution 33: 1870–1874. https://doi.org/10.1093/molbev/msw054

Lanfear R, Calcott B, Ho SY, Guindon S (2012) PartitionFinder: combined selection of partitioning schemes and substitution models for phylogenetic analyses. Molecular biology and evolution 29(6): 1695–1701. https://doi.org/10.1093/molbev/msw020

Lanfear R, Frandsen PB, Wright AM, Senfeld T, Calcott B (2016) PartitionFinder 2: new methods for selecting partitioned models of evolution formolecular and morphological phylogenetic analyses. Molecular biology and evolution. https://doi.org/10.1093/molbev/msw260

Leech JH (1891) New Species of Rhopalocera from Western China. Entomologist 24: 57–61. https://archive.org/details/entomologist189195brit/page/59/mode/1up?view=theater

Li WL, Cong Q, Shen JH, Zhang J, Hallwachs W, Janzen DH, Grishin NV (2019) Genomes of skipper butterflies reveal extensive convergence of wing patterns. Proceedings of the National Academy of Sciences of the United States of America 116: 6232–6237. https://doi.org/10.1073/pnas.1821304116
Liu GC, Chang Z, Chen L, He JW, Dong ZW, Yang J, Lu SH, Zhao RP, Wan WT, Ma GL, Li J, Zhang R, Wang W, Li XY (2020) Genome size variation in butterflies (Insecta, Lepidoptera, Papilionoidea): a thorough phylogenetic comparison. Systematic Entomology 45(3): 571–582. https://doi.org/10.1111/syen.12417

Miller MA, Pfeiffer W, Schwartz T (2010) Creating the CIPRES Science Gateway for Inference of Large Phylogenetic Trees. Proceedings of the Gateway Computing Environments Workshop 14: 1–8. https://doi.org/10.1109/GCE.2010.5676129

Minh BQ, Nguyen MAT, Haeseler VA (2013) Ultrafast approximation for phylogenetic bootstrap. Molecular Biology and Evolution 30: 1188–1195. https://doi.org/10.1093/molbev/mst024

Monteriro A, Pierce NE (2001) Phylogeny of Bicyclus (Lepidoptera: Nymphalidae) inferred from COI, COII and EF-1α gene sequences. Molecular Phylogenetics And Evolution 18(2): 264–281. https://doi.org/10.1006/mpev.2000.0872

Nguyen LT, Schmidt HA, von Haeseler A, Minh BQ (2015) IQ-TREE: A fast and effective stochastic algorithm for estimating maximum-likelihood phylogenies. Molecular Biology and Evolution 32(1): 268–274. https://doi.org/10.1093/molbev/msu300

Peng Y, Leung HCM, Yiu SM, Chin FYL (2012) IDBA-UD: a de novo assembler for single-cell and metagenomic sequencing data with highly uneven depth. Bioinformatics 28(11): 1420–1428. https://doi.org/10.1093/bioinformatics/bts174

Ronquist F, Teslenko M, Mark PVD, Ayres DL, Darling A, Höhna S, Larget B, Liu L, Suchard MA, Huelsenbeck JP, Notes A (2012) MrBayes 3.2: Efficient Bayesian Phylogenetic Inference and Model Choice Across a Large Model Space. Systematic Biology 61(3): 539–542. https://doi.org/10.1093/sysbio/sys029

Sahoo RK, Warren AD, Wahlberg N, Brower AV, Lukhtanov VA, Kodandaramaiah U (2016) Ten genes and two topologies: An exploration of higher relationships in skipper butterflies (Hesperiidae). PeerJ 4: e2653. https://doi.org/10.7717/peerj.2653

Sahoo RK, Warren AD, Collins SC, Kodandaramaiah U (2017) Hostplant change and paleoclimatic events explain diversification shifts in skipper butterflies (Family: Hesperiidae). BMC Evolutionary Biology 17: e174. https://doi.org/10.1186/s12862-017-1016-x

Scott JA, Wright DM (1990) Butterfly phylogeny and fossils. In: Kudrna O (Ed.) Butterflies of Europe 2, Aula Verlag, Wiesbaden, 152–208.

Thompson J, Gibson TJ, Plewniak F, Jeanmougin F, Higgins DGG (1997) The CLUSTAL_X Windows Interface: Flexible Strategies for Multiple Sequence Alignment Aided by Quality Analysis Tools. Nucleic Acids Research 25(24): 4876–4882. https://doi.org/10.1093/nar/25.24.4876

Toussaint EFA, Breinholt JW, Earl C, Warren AD, Brower AVZ, Yago M, Dexter KM, Espeland M, Pierce NE, Lohman DJ, Kawahara AY (2018) Anchored phylogenomics illuminates the skipper butterfly tree of life. BMC Evolutionary Biology 18(1): e101. https://doi.org/10.1186/s12862-018-1216-z

Toussaint EFA, Ellis EA, Gott RJ, Warren AD, Dexter KM, Storer C, Lohman DJ, Kawahara AY (2020) Historical biogeography of Heteropterinae skippers via Beringian and post-Tethyan corridors. Zoologica Scripta 00: 1–12. https://doi.org/10.1111/zsc.12457
Trifinopoulos J, Nguyen LT, Haeseler AV, Minh BQ (2016) W-IQ-TREE: a fast online phylogenetic tool for maximum likelihood analysis. Nucleic Acids Research 44. gkw256. https://doi.org/10.1093/nar/gkw256

Wahlberg N, Wheat CW (2008) Genomic outposts serve the phylogenomic pioneers: Designing novel nuclear markers for genomic DNA extractions of Lepidoptera. Systematic Biology 57(2): 231–242. https://doi.org/10.1080/10635150802033006

Ward PS, Downie DA (2005) The ant subfamily Pseudomyrmecinae (Hymenoptera: Formicidae): phylogeny and evolution of big-eyed arboreal ants. Systematic Entomology 30: 310–335. https://doi.org/10.1111/j.1365-3113.2004.00281.x

Warren AD, Ogawa JR, Brower AV (2008) Phylogenetic relationships of subfamilies and circumscription of tribes in the family Hesperiidae (Lepidoptera: Hesperioidea). Cladistics 24(5): 642–676. https://doi.org/10.1111/j.1096-0031.2008.00218.x

Warren AD, Ogawa JR, Brower AV (2009) Revised classification of the family Hesperiidae (Lepidoptera: Hesperioidea) based on combined molecular and morphological data. Systematic Entomology 34(3): 467–523. https://doi.org/10.1111/j.1365-3113.2008.00463.x

Wiemers M, Chazot N, Wheat CW, Schweiger O, Wahlberg N (2020) A complete time-calibrated multi-gene phylogeny of the European butterflies. ZooKeys 938: 97–124. https://doi.org/10.3897/zookeys.938.50878

Wu CS, Hsu YF (2017) Butterflies of China. The Straits Publishing and Distributing Group, Fuzhou, 2036 pp.

Yuan F, Yuan XQ, Xue GX (2015) Fauna Sinica (Insecta: Lepidoptera: Hesperiidae). Science Press, Beijing, 769 pp.

Zhang J, Cong Q, Shen JH, Brockmann E, Grishin NV (2019) Three new subfamilies of skipper butterflies (Lepidoptera, Hesperiidae). ZooKeys 861: 91–105. https://doi.org/10.3897/zookeys.861.34686

Zhang J, Lees DC, Shen JH, Cong Q, Huertas B, Martin G, Grishin NV (2020) The mitogenome of a Malagasy butterfly *Malaza fastuosus* (Mabille, 1884) recovered from the holotype collected over 140 years ago adds support for a new subfamily of Hesperiidae (Lepidoptera). Genome 63: 195–202. https://doi.org/10.1139/gen-2019-0189

**Supplementary material I**

**Table S1**

Author: Yongxiang Hou
Data type: table
Explanation note: Label information and GenBank accession numbers of the specimens used in this study.
Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.
Link: https://doi.org/10.3897/zookeys.1055.68640.suppl1
Supplementary material 2

Table S2
Author: Yongxiang Hou
Data type: table
Explanation note: Primers used in this study.
Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.
Link: https://doi.org/10.3897/zookeys.1055.68640.suppl2

Supplementary material 3

Table S3
Author: Yongxiang Hou
Data type: table
Explanation note: The best-fit partition schemes and model calculated by PartitionFinder v2.1.1.
Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.
Link: https://doi.org/10.3897/zookeys.1055.68640.suppl3