A multilocus assessment reveals two new synonymies for East Asian Cyclommatus stag beetles (Coleoptera, Lucanidae)

Jiao Jiao Yuan\(^{1,2}\), Dan Chen\(^{1,2}\), Xia Wan\(^{1,2}\)

1 Department of Ecology, School of Resources and Engineering, Anhui University, 111 Jiulong Rd., Hefei 230601, China 2 Anhui Province Key Laboratory of Wetland Ecosystem Protection and Restoration, Hefei 230601, China

Corresponding author: Xia Wan (wanxia@ahu.edu.cn)

Abstract

Cyclommatus scutellaris Möllenkamp, 1912, Cyclommatus elsae Kriesche, 1921 and Cyclommatus tamdaoensis Fujita, 2010 are East Asian stag beetle species with long-debated taxonomic relationships due to high intraspecific morphological variability. In this study, we applied multilocus phylogenetic analyses to reassess their relationships. Two mitochondrial genes (16S rDNA, COI) and two nuclear genes (28S rDNA, Wingless) were used to reconstruct the phylogeny through the Bayesian inference (BI) and Maximum Likelihood (ML) methods. Both topologies supported two clades: the clade C. scutellaris was sister to the clade (C. elsae + C. tamdaoensis) with the subclade C. tamdaoensis embedded in the subclade C. elsae. The Kimura 2-parameter (K2P) genetic distance analysis yielded a low mean value (≤0.035) among the three taxa, which was well below the minimum mean value between other Cyclommatus species (≥0.122). We also compared the accuracy and efficiency of two approaches, GMYC and ABGD, in delimitating the three lineages. The result shows that ABGD is a better approach than GMYC. Our molecular data recognizes the three species as different populations of a single species, ranging from Taiwan Island to the continent. Therefore, we propose two new junior synonyms for C. scutellaris: C. tamdaoensis, syn. nov. and C. elsae syn. nov.

Keywords

East Asia, Lucanidae, molecular phylogeny, morphology, new synonym, species delimitation
Introduction

The genus *Cyclommatus* Parry, 1863 (Lucanidae), includes some of the most striking stag beetles with enormous male mandibles and metallic body colorations. Many species display strong sexual dimorphism and male polymorphism, causing substantial confusion in morphological-based species delimitation. Phylogenetic analysis using molecular markers such as mitochondrial and nuclear genes can clarify many morphology-based species/subspecies’ taxonomic positions. Yet, in *Cyclommatus*, only a few such studies have been performed for a limited number of species (Kim and Farrell 2015; Tsai and Yeh 2016) and only mitogenomic data of *Cyclommatus vitalisi* Pouillaude, 1913 from southwestern China have been reported (Liu et al. 2017). To our knowledge, no multilocus phylogenetic analysis has been applied to resolve taxonomic debates in this genus to date.

This study aims to resolve the long-debated taxonomic relationships among three *Cyclommatus* species, namely *C. scutellaris* Möllenkamp, 1912; *C. elsae* Kriesche, 1921 and *C. tamdaoensis* Fujita, 2010. *C. scutellaris* was first described by Möllenkamp (1912) and believed to be endemic to Taiwan Island. *Cyclommatus elsae* and *C. tamdaoensis* are distributed in southeastern China and the China-Vietnam border, respectively (Fig. 1). The three lineages’ taxonomic relationships were frequently revised by different studies, resulting in inconsistent conclusions (Möllenkamp 1912; Kurosawa 1974; Bomans 1979; Lacroix 1988; Maes 1992; Fujita 2010; Schenk and Nguyen 2015). Most of these revisions, however, lack sufficient data support. For instance, Fujita (2010) treated *C. scutellaris* and *C. elsae* as valid species and described the new taxon *C. tamdaoensis* from northern Vietnam. More recently, Huang and Chen (2017) considered them three subspecies based on morphological characters and considered *C. princeps* Schenk & Nguyen, 2015 from northern Vietnam as a junior synonym of *C. tamdaoensis*. Both studies used some key morphological diagnostic traits showing a considerable degree of intraspecific variabilities, such as the stripes on the pronotum and the male genitalia. The black lateral stripes in male individuals of *C. elsae* display two forms: the complete-form extending to the anterior angle of the pronotum and the partly patched form along the margins [see pl. 194, figs 8–2–2, 8–2–4 in Huang and Chen (2017)]. The black median stripe shows three forms: ranging from the broadly diamond shape to the reduced midline, then disappearing completely [see pl. 194, figs 8–2–2, 8–2–4 in Huang and Chen (2017)]. The lateral stripes of male *C. scutellaris* are patched and the middle one is absent, while those of *C. tamdaoensis* are the complete-form and diamond-shaped, respectively [see pl. 194, figs 8–1, 8–3 in Huang and Chen (2017)]. Moreover, characters of male genitalia, including the median lobes and the basal piece, which are very important in lucanid taxonomy (Holloway 2007), show variability within the samples of *C. elsae*, *C. scutellaris* and *C. tamdaoensis* [see pl. 196 in Huang and Chen (2017)]. The protibiae of males of all three taxa consistently show yellow setae along 3/5 of their inner margin [see pl. 194, figs 8–1–1, 8–2–1, 8–3–1 in Huang and Chen (2017)]. Finally, the females of all three taxa look uniform with broad lateral stripes and a
Two new synonyms of *Cyclommatus* stag beetles

Figure 1. Collection localities of *C. scutellaris*, *C. elsae* and *C. tamdaoensis*. The map is generated using ARCGIS 10.2 software based on the geospatial data from the National Geomatics Center of China.

The broadly distributed *C. elsae* displays substantial morphological variability within its range. Populations from Mt. Nanling in Guangdong are morphologically almost identical to *C. scutellaris*, whereas the Mt. Simianhshan (Chongqing) population are nearly indistinguishable from *C. tamdaoensis*. The three groups’ intraspecific variations and interlineage morphological convergence suggest that they are likely different populations of the same species. Yet, the lack of molecular data makes it difficult to clarify their conspecific status. In the current study, we applied the multilocus approach to recognize the three species’ relationship. Our goal is to resolve this long-debated issue in *Cyclommatus* species phylogeny while testing the multilocus molecular approach’s power and efficiency in resolving species delimitation problems for lucanid beetles.

**Materials and methods**

**Taxon sampling**

Thirty-three stag beetle specimens were examined, including 29 samples of *Cyclommatus* (15 *C. elsae*; 9 *C. scutellaris*; 5 *C. tamdaoensis*) as in groups, and 4 samples as outgroups...
(C. albersi Kraatz, 1894, C. mniszechi (Thomson, 1856); C. nagaii Fujita, 2010; C. vitalisi). Voucher specimens and their extracted genomic DNA are deposited in the research collection at the Museum of Anhui University, China (Suppl. material 1: Table S1).

**Laboratory protocols**

The specimens were stored in 100% ethanol at -20 °C for molecular analysis. A small muscle was extracted from the sample using the Blood and Tissue Kit for total DNA extraction (Qiagen, Germany). Extracted DNA was stored at -20 °C until needed for the polymerase chain reaction (PCR). Four gene regions were targeted using the PCR. These regions included mitochondrial genes (16S rDNA, COI) and two nuclear genes (28S rDNA, Wingless). COI is often used in DNA barcoding because of its strong species identification ability (Han et al. 2016; Thormann et al. 2016). 16S rDNA is a conserved gene in animal mitochondria, which has a slow evolution rate. This gene is suitable for the study of distant taxa (Simon et al. 1994). COI and 16SrDNA are the most commonly used mitochondrial genes in the Lucanidae phylogeny (Tsai and Yeh 2016). 28S rDNA is relatively conservative in the evolutionary process and contains a highly variable region in the conserved sequence. It is also an excellent molecular marker to solve the phylogenetic relationship from species to families. Because wing pattern is a vital criterion for insect identification, Wingless is a necessary candidate gene for studying insect phylogeny. Still, the gene is only used in Lepidoptera and Coleoptera systematics (Liu and Jiang 2005). So, all specimens were sequenced for 16S rDNA, COI, 28S rDNA, and Wingless.

The primer sets used to amplify 16S rDNA, COI, 28S rDNA, and Wingless are shown in Table 1. PCR amplification reaction was carried out in a volume of 25 μL, in which 10 μM of each primer (forward and reverse) 1 μL, 2 μL template DNA, 12.5 μL 2×EasyTaq SuperMix (dye) and 8.5 μL sterile double distilled water (ddH2O) constituted a final volume of 25μL. PCR conditions were as followed: initial denaturation at 94 °C for 2 minutes, followed by 35–37 cycles of denaturation at 94 °C for the 40 s and annealing at 52–58 °C for 50s, and elongation at 70 °C for 50s, and then a final extension step at 72 °C for 7 min. The length of the fragments determined the annealing temperature. For sequencing, we used the ABI PRISM BigDye terminator version 3.1.

**Table 1.** List of the primer pairs and their respective reference used during the present study.

| Gene     | Primer name | Sequence (5’–3’) | Reference                  |
|----------|-------------|------------------|---------------------------|
| COI      | COI-F1      | CAACATTTATTTTGTATTTTGG | Simon et al. (1994)       |
|          | COI-R1      | TCCAATGCACTAATCTGCCATATA | Simon et al. (1994)       |
| 16S rDNA | 16S-F1      | CCGTTTTGACTCGACATCGT | Hosoya et al. (2001)     |
|          | 16S-R1      | TAATTTATTGACCTTGTGTACAG | Hosoya et al. (2001)     |
| 28S rDNA | 28SDD       | GGACCCCTTTGAAACAC   | Monaghan et al. (2007)   |
|          | 28SSF       | TTACACACTCCTTACGGGAT | Monaghan et al. (2007)   |
| Wingless | Wg550F      | ATCCGTACGARTGAYAARTGCAYGGATGTC | Wild and Maddison (2008) |
|          | WgAbRZ      | CACTTNACTRCRCACCAARTG | Wild and Maddison (2008) |
|          | Wg578F      | TGCACNGTGARACTGCTTGATGT | Ward and Downie (2005)   |
|          | WgAbR       | ACYTCCGACGCACCGGAA  | Abouheif and Wray (2002) |
Two new synonyms of *Cyclommatus* stag beetles

Sequencing kit (Life Technologies, USA) and cycle sequencing reactions were performed on ABI PRISM 3730xl automated sequencers (Life Technologies, USA) at Sangon Bio-tech Company, China. The sequences were submitted to GenBank with the accession number (Suppl. material 1: Table S1).

**Phylogenetic analyses**

Sequences were assembled in GENEIOUS PRIME 2019.1.1. All sequences were aligned in MEGA 7 (Kumar et al. 2016). Divergences among taxa were analyzed using MEGA 7.0 based on the Kimura 2-parameter model. DNA COI sequences of *Cyclommatus* were assembled for genetic distance analyses. Bayesian inference (BI) and Maximum Likelihood (ML) analyses were conducted using MRBAYES 3.2.7a and IQ-TREE web server, respectively. The BI and ML analysis was conducted on the CIPRES Science Gateway (Miller et al. 2011). Two independent chains starting from a random tree were run for 20,000 generations, sampling the tree every 10 generations. The initial 25% trees of each Markov Chain Monte Carlo (MCMC) chain were discarded as burn-in. A consensus tree was computed from the remaining 1500 trees combined from two runs, and the two runs converged at a maxdiff of less than 0.1. For ML analyses, the “automatic” option was set under the optimal evolution model, and the phylogenetic trees were constructed using an ultrafast bootstrap approximation approach with 10,000 replicates. Phylogenetic trees were viewed and edited in FIGTREE 1.4.4.

**Divergence time analysis**

Divergence time was estimated with a relaxed clock Exponential model (Drummond and Rambaut 2007) in BEAST 2.6.0 (Bouckaert et al. 2014). The MCMC chain was run for $8.5 \times 10^8$ generations, with four independent runs. Substitution rates of 1.77% per lineage in a million years (Myr) for COI combining with 0.54%/lineage/Myr for 16S rDNA have been suggested optimally for beetles (Papadopoulou et al. 2010). The resultant BEAST log files were viewed using TRACER 1.7 (Rambaut et al. 2018) to analyze the output results of the Effective Sample Size (ESS) for the posterior distribution of estimated parameter values. With a 25% burn-in threshold, all post-burn-in trees from the four independent runs were combined using the software LOG COMBINER 2.6.0 (Bouckaert et al. 2014). TREE ANNOTATOR 2.5.1 (Bouckaert et al. 2014) was used to summarize information (i.e., Nodal posterior probabilities, posterior estimates and highest posterior density limits) from the individual post-burn-in trees onto a single maximum clad credibility (MCC) tree. The summarized information was visualized on the MCC tree using FIGTREE 1.4.4.

**Species delimitation**

We performed species delimitation using Automatic Barcode Gap Discovery (ABGD) and Generalized Mixed Yule Coalescent (GMYC) methods. The ABGD detects a gap
in divergence distribution, which corresponds to differences between intraspecific and interspecific distances. When a gap exists, the process works well for species delimitation (Puillandre et al. 2012a). The ABGD analyses were performed at the webserver (https://bioinfo.mnhn.fr/abi/public/abgd/abgdweb.html). The following setting was used: steps (20), distance Kimura (K80) TS/TV (2.0), other parameter values employed defaults. GMYC delimits distinct genetic clusters by optimizing the set of nodes defining the transitions between inter- and intraspecific processes (Pons et al. 2006). The analysis was conducted using BEAST 2.6.0 under a relaxed clock Exponential model. ESS values assessed convergence. A burn-in with 25% was set to obtain an optimal consensus tree. The resulting tree was then used to analyze the data under the GMYC species delimitation approach in the software R with the package ‘splits’ using the single-threshold method (Ezard et al. 2015).

Results

Phylogenetic relationships

The BI and ML analysis showed consistent topology with a highly supported backbone (Fig. 2; Suppl. material 2: Fig. S1). The clade *C. scutellaris* was sister to *C. elsae* + *C. tamdaoensis* (Bayesian posterior probability, BPP = 1, Maximum Likelihood Bootstrap, MLB = 100). And the subclade *C. tamdaoensis* embedded in *C. elsae* (BPP = 0.7, MLB = 76). *Cyclommatus scutellaris* was an early branch in the three taxa and the species was monophyletic.

Genetic distances

The genetic distances using the COI gene were calculated among the three taxa. The mean genetic distances between the *C. scutellaris*, *C. elsae* and *C. tamdaoensis* were no more than 0.035 (Table 2). The numbers were well below the minimum mean genetic distances of 0.122 among interspecific taxa of *Cyclommatus*.

Divergence time analysis

Based on the COI gene and 16S rDNA gene, calibration time was analyzed to describe these taxonomically controversial species’ possible differentiation history. The analysis under BI converged well as all parameters had ESS values above 200. The mean divergence age estimates and 95% High Posterior Density (HPD) for nodes of interest-based on the BEAST analysis are presented in Fig. 3, and Suppl. material 3: Fig. S2. Among the three taxa, *C. scutellaris* diverged during the Middle Pleistocene circa 1.23 million years ago (Mya) (95% HPD: 0.58–2.28 Mya). Within the clade *C. elsae* + *C. tamdaoensis*, the crown node’s age estimate was dated to be 0.62Mya (95% HPD: 0.32–1.06 Mya) in the Late Pleistocene. The age of *C. tamdaoensis* was estimated to be 0.49 Mya (95% HPD: 0.27–0.82 Mya) in the Late Pleistocene.
Two new synonyms of *Cyclommatus* stag beetles

Figure 2. Bayesian topology showing the relationships within *C. scutellaris*, *C. elsae*, *C. tamdaoensis* and outgroups. Values next to each node represent Bayesian posterior probabilities (first number) and maximum likelihood bootstrap support (second number). The phylogenetic tree is based on Bayesian inference analysis of concatenated DNA sequence data from 16S rDNA, 28S rDNA, COI and Wingless. The columns on the right show numbers of entities identified by the ABGD, GMYC.

Species delimitation

Two species delimitation methods were employed to evaluate which one was most suitable for multilocus phylogenetic analyses. ABGD analyses suggested the three taxa were one molecular operational taxonomic unit (MOTU), whereas GMYC divided *C. scutellaris*, *C. elsae* and *C. tamdaoensis* into two MOTUs (Fig. 2).
Table 2. The mean genetic distances among all the taxa (Kimura 2-parameter)

|          | 1   | 2   | 3   | 4   | 5   | 6   |
|----------|-----|-----|-----|-----|-----|-----|
| C. albersi |     |     |     |     |     |     |
| C. elsae  |     |     |     |     |     |     |
| C. mniszechi | 0.166 |     |     |     |     |     |
| C. nagaii | 0.142 |     |     |     |     |     |
| C. scutellaris | 0.166 | 0.035 |     |     |     |     |
| C. tamdaoensis | 0.155 | 0.016 | 0.158 | 0.122 | 0.031 |     |
| C. vitalisi | 0.152 | 0.150 | 0.145 | 0.146 | 0.159 | 0.146 |

Figure 3. Maximum clade credibility time-tree obtained from BEAST based on COI and 16S rDNA. Divergence time estimates are represented next to the nodes (in millions of years) with horizontal bars indicating 95% highest posterior density intervals.
Two new synonyms of *Cyclommatus* stag beetles

**Taxonomic account**

*Cyclommatus scutellaris* Möllenkamp, 1912

*Cyclommatus scutellaris* Möllenkamp, 1912: 7.
*Cyclommatus elsae* Kriesche, 1921: 95. syn. nov.
*Cyclommatus tamdaoensis* Fujita, 2010: 14. syn. nov.
*Cyclommatus princeps* Schenk & Nguyen, 2015: 7. (Synonymized as a junior synonym of *Cyclommatus tamdaoensis* by Huang and Chen 2017: 193)

**Material examined.** China • 1 male; Yunnan Province, Shuifu County; 27 Jul. 2010; JS Xu and LX Chang leg. • 1 male; Shaanxi Province, Zhenba County; 10 Jul. 1996. • 3 males; Chongqing Province, Mt. Simianshan; 29 Aug. 2014. • 3 males; Guizhou Province, Mt. Fanjingshan; 29 Jul. 2014; YF Wu. • 1 male; Guizhou Province, Xishui County, 18 Jul. 2015, LX Zhu. • 1 male; Fujian Province, Mt. Wuyishan; 18 Jul. 2011; Q Zhang and YY Cao. • 1 male; same locality as for preceding; 18 Aug. 2010; F Zhong and XY Hu. • 3 males; Taiwan Island; 29 Aug. 1995. • 1 male; same locality as for preceding; 17 May 2008. • 1 male; same locality as for preceding; 5 Oct. 2008. • 2 males; same locality as for preceding; 20 Jul. 2010; • 2 males; same locality as for preceding; 21 Jul. 2011. • 3 males; Guangdong Province, Mt. Nanling; 18 Aug. 2010; HY Liu. • 1 male; same locality as for preceding; 9 Aug. 2011, F Zhong; • 4 males; Guangxi Autonomous Region, Jinxiu County; 18 Jul. 2012. • 1 male; same locality as for preceding; 28 Jun. 2013.

**Diagnosis.** The species is characterized in the male by the matt dorsal surface of the entire body and the long seta-range on the inner margin of the protibial, and in the female by the elytra usually without black stripes. The female can be distinguished from other members of *Cyclommatus* by the following combination of characters: 1) ground color of the dorsal surface of the body redder; 2) canthus often short, with lateral end blunt, and usually with a convex or straight outer margin; 3) central black band on the pronotum clearly defined and often narrower than the lateral orange band (Huang and Chen 2017).

**Distribution.** China (Yunnan, Shaanxi, Chongqing, Guizhou, Fujian, Taiwan Island, Guangdong, Guangxi).

**Discussion**

This study presents consistent phylogenetic relationships inferred by the BI and ML methods. The trees’ identical topology strongly supports that *C. scutellaris* is sister to the clade *C. elsae + C. tamdaoensis*. The nested structure of *C. elsae* and *C. tamdaoensis* suggests that the two nominal taxa should be treated as one species. The genetic distance also indicates that *C. elsae* and *C. tamdaoensis* are most likely two continental populations of the Taiwanese species *C. scutellaris*. 
The divergence time estimation shows that *C. scutellaris* began to diverge from the Middle Pleistocene with subsequent deep genetic isolation during the Late Pleistocene. During the glacial maxima, a land bridge facilitated the contraction of the geographical ranges of some species southwards into Taiwan (Kawamura et al. 2016). Once the glaciations ended, the rising sea levels resulted in vicariant isolation. The relict populations might have retreated to the montane habitat of Taiwan Island, causing the present disjunct distributions (Päckert et al. 2011; Wang et al. 2013). Many studies interpreted the global cooling and aridification, the Qinghai–Tibet Plateau uplift and the abrupt climate change in East Asia during the Neogene and Quaternary as responsible for the speciation, intraspecific divergence, genetic diversification, and significant phylogeographic break between the western and eastern clade in Chinese fauna (Meng et al. 2015; Qiu et al. 2017; Ye et al. 2017; Zhou et al. 2017).

Accurate species delimitations are critical in many biology areas, such as conservation biology (designating endangered species) and evolutionary biology (describing diversification patterns). Traditionally, species are identified and described by morphological characters. However, using morphological data alone may underestimate the number of species (Yang and Rannala 2010). Therefore, molecular data are used to delimitate species in the present study. However, GMYC and ABGD methods obtained different results. GMYC has been developed to delimit species according to single-locus data, which has a strong theoretical basis. Although the method is based on a reliable phylogenetic framework, it largely depends on the ultrametric gene tree's correctness. Errors in the framework underpinning the analysis may affect the final results (Kekkonen and Hebert 2014). Also, GMYC usually over-splits species, mainly due to low genetic divergence between lineages and overlap of interspecific and intraspecific divergences, or due to lack of reciprocal monophyly between sister clades (Talavera et al. 2013; Pentinsaari et al. 2016; Stokkan et al. 2018). Evaluations of the accuracy and efficiency of ABGD conducted by different researchers reached a consensus that ABGD is more conservative and faster than other methods (Puillandre et al. 2012a; Puillandre et al. 2012b; Kekkonen and Hebert 2014; Guillemin et al. 2016). Our results also produced a consistent conclusion. ABGD produced one MOTU, which agrees with the number of species defined based on multilocus phylogenetic analyses.

In summary, our study resolved a long-standing debate on species recognition in the genus *Cyclommatus* while illustrating the necessity of employing multiple data types, both morphological and molecular, for efficient and accurate species delimitation in taxa with a high degree of phenotypic convergence and variations.

**Acknowledgements**

We want to express our gratitude to Dr Jian-Yue Qiu, Dr Hao Xu, Dr Yun-Fei Wu, Prof. Ai-Min Shi and Mr Chang-Chin Chen for their kind donation of specimens. This work was supported by research grants from the National Natural Science Foundation of China (No. 31872276; 31572311).
Two new synonyms of *Cyclommatus* stag beetles

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

Bomans HE (1979) Contribution à l’étude des Coléoptères Lucanides. Description de l’allosyntype de *Cyclommatinus elsae* Kriesche. Bulletin et Annales de la Société Royale d’Entomologie de Belgique 115: 194–196.

Bouckaert R, Heled J, Kuhnert D, Vaughan T, Wu CH, Xie D, Suchard MA, Rambaut A, Drummond AJ (2014) BEAST 2: A software platform for Bayesian evolutionary analysis. PLoS Computational Biology 10: e1003537. https://doi.org/10.1371/journal.pcbi.1003537

Drummond AJ, Rambaut A (2007) BEAST: Bayesian evolutionary analysis by sampling trees. BMC Evolutionary Biology 7: e214. https://doi.org/10.1186/1471-2148-7-214

Ezard T, Fujisawa T, Barraclough T (2015) R package splits: Species’ limits by threshold statistics, version 1.0–18/r45. http://R-Forge.R-project.org/projects/splits/

Fujita H (2010) The Lucanid Beetles of the World. Mushi-sha, Tokyo, 472 pp.

Guillemin ML, Contreras-Porcia L, Ramírez ME, Macaya EC, Contador CB, Woods H, Wyatt C, Brodie J (2016) The bladed Bangiales (Rhodophyta) of the South Eastern Pacific: Molecular species delimitation reveals extensive diversity. Molecular Phylogenetics and Evolution 94: 814–826. https://doi.org/10.1016/j.ympev.2015.09.027

Han T, Lee W, Lee S, Park IG, Park H (2016) Reassessment of Species Diversity of the Subfamily Denticollinae (Coleoptera: Elateridae) through DNA Barcoding. PLoS ONE 11: e0148602. https://doi.org/10.1371/journal.pone.0148602

Holloway BA (2007) Lucanidae (Insecta: Coleoptera). Fauna of New Zealand 61: 1–254.

Hosoya T, Honda M, Araya K (2001) Genetic variations of 16S rRNA gene observed in *Ceruchus lignarius* and *Dorcus rectus rectus* (Coleoptera: Lucanidae). Entomological Science 4: 335–344.

Huang H, Chen CC (2017) Stag Beetles of China III. Formosa Ecological Company, Taiwan, 540 pp.

Kawamura A, Chang CH, Kawamura Y (2016) Middle Pleistocene to Holocene mammal faunas of the Ryukyu Islands and Taiwan: An updated review incorporating results of recent research. Quaternary International 397: 117–135. https://doi.org/10.1016/j.quaint.2015.06.044

Kekkonen M, Hebert PDN (2014) DNA barcode-based delineation of putative species: Efficient start for taxonomic workflows. Molecular Ecology Resources 14: 706–715. https://doi.org/10.1111/1755-0998.12233

Kim SI, Farrell BD (2015) Phylogeny of world stag beetles (Coleoptera: Lucanidae) reveals a Gondwanan origin of Darwin’s stag beetle. Molecular Phylogenetics and Evolution 86: 35–48. https://doi.org/10.1016/j.ympev.2015.02.015

Kraatz G (1894) *Cyclommatus albersi* nov. sp., Hirschkäfer von Birma. Deutsche Entomologische Zeitschrift 1894: 268–269. https://doi.org/10.1002/mmnd.48018940522

Kriesche R (1921) Zur Kenntnis der Lucaniden. Archiv für Naturgeschichte 86: 92–107.

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
Kurosawa Y (1974) A stag-beetle new to the Formosan fauna (Coleoptera: Lucanidae). Bulletin of the National Science Museum. Series A: Zoology 17: 103–104.

Lacroix JP (1988) Descriptions de Coleoptera Lucanidae nouveaux ou peu connus (6eme note). Bulletin de la Société Sciences Nat 1988: 5–7.

Liu DF, Jiang GF (2005) The application of nuclear genes sequences in insect molecular systematics. Zoological Systematics 30: 484–492. https://doi.org/10.3969/j.issn.1000-0739.2005.03.007

Liu J, Li C, You S, Wan X (2017) The first complete mitogenome of Cyclommatus stag beetles (Coleoptera: Lucanidae) with the phylogenetic implications. Entomotaxonomia 39: 294–299.

Maes JM (1992) Lista de los Lucanidae (Coleoptera) del Mundo. Revista Nicaraguense de Entomología 22: 1–121.

Meng HH, Gao XY, Huang JF, Zhang ML (2015) Plant phylogeography in arid Northwest China: Retrospectives and perspectives. Journal of Systematics and Evolution 53: 33–46. https://doi.org/10.1111/jse.12088

Miller MA, Pfeiffer W, Schwartz T (2011) The CIPRES Science Gateway: A community resource for phylogenetic analyses. In: Cordero Rivera A (Ed.) Proceedings of the 2011 Tera-Grid Conference: Extreme Digital Discovery. Association for Computing Machinery, New York, NY, United States, Article 41: 1–8. https://doi.org/10.1145/2016741.2016785

Möllenkamp W (1912) H. Sauter's Formosa Ausbeute. Lucanidae. (Coleoptera). Entomologische Mitteilungen 1: 6–8. https://doi.org/10.5962/bhl.part.25903

Monaghan MT, Inward DJG, Hunt T, Vogler AP (2007) A molecular phylogenetic analysis of the Scarabaenae (dung beetles). Molecular Phylogenetics and Evolution 45: 674–692. https://doi.org/10.1016/j.ympev.2007.06.009

Päckert M, Martens J, Sun YH, Severinghaus LL, Nazarenko AA, Ting J, Töpfer T, Tietze DT (2011) Horizontal and elevational phylogeographic patterns of Himalayan and Southeast Asian forest passerines (Aves: Passeriformes). Journal of Biogeography 39: 556–573. https://doi.org/10.1111/j.1365-2699.2011.02606.x

Papadopoulou A, Anastasiou I, Vogler AP (2010) Revisiting the insect mitochondrial molecular clock: The Mid-Aegean trench calibration. Molecular Biology and Evolution 27: 1659–1672. https://doi.org/10.1093/molbev/msq051

Parry FJS (1863) A few remarks upon Mr. James Thomson’ s catalogue of Lucanidae, published in the ‘Annales de la Société Entomologique de France, 1862.’ Transactions of the Royal Entomological Society of London 1: 442–452. https://doi.org/10.1111/j.1365-2311.1863.tb01291.x

Pentinsaari M, Vos R, Mutanen M (2016) Algorithmic single-locus species delimitation: Effects of sampling effort, variation and nonmonophyly in four methods and 1870 species of beetles. Molecular Ecology Resources 17: 393–404. https://doi.org/10.1111/1755-0998.12557

Pons J, Barraclough TG, Gomez-Zurita J, Cardoso A, Duran DP, Hazell S, Kamoun S, Sumlin WD, Vogler AP (2006) Sequence-Based Species Delimitation for the DNA Taxonomy of Undescribed Insects. Systematic Biology 55: 595–609. https://doi.org/10.1080/10635150600852011

Pouillaude I (1913) Note sur quelques Lucanidae d’Indo-Chine. Insecta, revue illustree d’Entomologie 3: 332–337.
Puillandre N, Lambert A, Brouillet S, Achaz G (2012a) ABGD, automatic barcode gap discovery for primary species delimitation. Molecular Ecology 21: 1864–1877. https://doi.org/10.1111/j.1365-294X.2011.05239.x

Puillandre N, Modica MV, Zhang Y, Sirovich L, Boisselier MC, Cruaud C, Holford M, Samadi S (2012b) Large-scale species delimitation method for hyperdiverse groups. Molecular Ecology 21: 2671–2691. https://doi.org/10.1111/j.1365-294X.2012.05559.x

Qiu YX, Lu QX, Zhang YH, Cao YN (2017) Phylogeography of East Asia’s Tertiary relict plants: Current progress and future prospects. Biodiversity Science 25: 136–146. https://doi.org/10.17520/biods.2016292

Rambaut A, Drummond AJ, Xie D, Baele G, Suchard MA (2018) Posterior summarization in bayesian phylogenetics using Tracer 1.7. Systematic Biology 67: 901–904. https://doi.org/10.1093/sysbio/syy032

Schenk KD, Nguyen TQ (2015) Description of two new taxa of the genus Cyclommatus Parry, 1863 from Central Vietnam (Coleoptera: Lucanidae). Beetles World 11: 7–11.

Simon C, Frati F, Beckenbach A, Crespi B, Liu H, Flook P (1994) Evolution, weighting, and phylogenetic utility of mitochondrial gene sequences and a compilation of conserved polymerase chain reaction primers. Annals of the Entomological Society of America 87: 651–701. https://doi.org/10.1093/aesa/87.6.651

Stokkan M, Jurado-Rivera JA, Oromí P, Juan C, Jaume D, Pons J (2018) Species delimitation and mitogenome phylogenetics in the subterranean genus Pseudoniphargus (Crustacea: Amphipoda). Molecular Phylogenetics and Evolution 127: 988–999. https://doi.org/10.1016/j.ympev.2018.07.002

Talavera G, Dincă V, Vila R (2013) Factors affecting species delimitations with the GMYC model: Insights from a butterfly survey. Methods in Ecology and Evolution 4: 1101–1110. https://doi.org/10.1111/2041-210X.12107

Thomson J (1856) Description de quatre Lucanides nouveaux de ma collection, précédée du catalogue des Coléoptères lucanoïdes de Hope (1845), et de l’arrangement méthodique adopté par Lacordaire pour sa famille des Pectinicornes. Revue et magasin de zoologie pure et appliquée 8: 516–528.

Thormann B, Ahrens D, Marín Armijos D, Peters MK, Wagner T, Wägele JW (2016) Exploring the Leaf Beetle Fauna (Coleoptera: Chrysomelidae) of an Ecuadorian Mountain Forest Using DNA Barcoding. PLoS ONE 11: e0148268. https://doi.org/10.1371/journal.pone.0148268

Tsai CL, Yeh WB (2016) Subspecific differentiation events of montane stag beetles (Coleoptera: Lucanidae) endemic to Formosa island. PLoS ONE 11: e0156600. https://doi.org/10.1371/journal.pone.0156600

Wang W, McKay BD, Dai C, Zhao N, Zhang R, Qu Y, Song G, Li SH, Liang W, Yang X, Pasquet E, Lei F (2013) Glacial expansion and diversification of an East Asian montane bird, the green-backed tit (Parus monticolus). Journal of Biogeography 40: 1156–1169. https://doi.org/10.1111/jbi.12055

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
Wild AL, Maddison DR (2008) Evaluating nuclear protein-coding genes for phylogenetic utility in beetles. Molecular Phylogenetics and Evolution 48: 877–891. https://doi.org/10.1016/j.ympev.2008.05.023

Yang Z, Rannala B (2010) Bayesian species delimitation using multilocus sequence data. Proceedings of the National Academy of Sciences of the United States of America 107: 9264–9269. https://doi.org/10.1073/pnas.0913022107

Ye JW, Zhang Y, Wang XJ (2017) Phylogeographic breaks and the mechanisms of their formation in the Sino-Japanese floristic region. Chinese Journal of Plant Ecology 41: 1003–1019. https://doi.org/10.17521/cjpe.2016.0388

Zhou Z, Huang J, Ding W (2017) The impact of major geological events on Chinese flora. Biodiversity Science 25: 123–135. https://doi.org/10.17520/biods.2016120

Supplementary material 1

Table S1. A list of specimens’ voucher information and GenBank accession numbers used in this study
Authors: Jiao Jiao Yuan
Data type: phylogenetic
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.1021.58832.suppl1

Supplementary material 2

Figure S1. Bayesian topology showing the relationships within C. scutellaris, C. elsae, C. tamdaoensis and outgroups
Authors: Jiao Jiao Yuan
Data type: phylogenetic
Explanation note: The figure also indicates outgroups and accession numbers. The columns on the right show numbers of entities identified by the ABGD, GMYC.
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.1021.58832.suppl2
Supplementary material 3

Figure S2. Maximum clade credibility time-tree based on COI and 16S rDNA
Authors: Jiao Jiao Yuan
Data type: phylogenetic
Explanation note: Age estimates are indicated at the nodes (Mya). The outgroups are shown in the figure.
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.1021.58832.suppl3