Multilocus phylogeny and revised classification for mountain dragons of the genus *Japalura s.l.* (Reptilia: Agamidae: Draconinae) from Asia

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Although the genus *Japalura s.l.* has long been recognized as paraphyletic based on limited genetic sampling, its problematic taxonomy has not been revised, and phylogenetic relationships among the majority of congeners remain unknown. Here we utilize a densely sampled dataset of both multilocus genetic and morphological data to provide the first phylogenetic inference of relationships among *Japalura s.l.* species. Our results show that *Japalura s.l.* is paraphyletic, consisting of four major clades that are scattered across the phylogeny of the subfamily Draconinae: the first clade from the western, central and middle-eastern Trans-Himalayas, the second clade from the far eastern Trans-Himalayas, the third clade from East Asia and the last clade from Indochina. To address this widespread paraphyly of the genus and to stabilize the taxonomy within the family Draconinae, we revise the current taxonomy and split *Japalura s.l.* into four genera. By doing so, we recognize two existing generic names, *Japalura sensu stricto* and *Pseudocalotes*, resurrect one name available in the literature, *Diploderma*, and describe one new genus, *Cristidorsa* gen. nov. We discuss phylogenetic relationships and taxonomy within *Japalura s.l.* and present a diagnostic key to all recognized genera of the subfamily Draconinae.

ADDITIONAL KEYWORDS: China – India – integrative taxonomy – lizard – new genus – Tibetan Plateau.

INTRODUCTION

Lizards of the Old World family Agamidae have long fascinated biologists with their extreme ecological and morphological variation, from winged, gliding arboreal species to spiny, desert specialists (Smith, 1935; Moody, 1980; Stuart-Fox & Ord, 2004). Within the family, dragon lizards of the subfamily Draconinae represent a remarkable radiation of reptiles distributed throughout Asia, with more than 214 species recognized (Pyron *et al.*, 2013; Grismer *et al.*, 2016b; Uetz & Hošek, 2017). The genus *Japalura* Gray, 1853 is one of the most speciose and most widespread genera in Draconinae, containing 34 recognized species that are distributed from Pakistan along the foothills of the Himalayas to the oceanic islands of Japan (Wang *et al.*, 2016; Ananjeva *et al.*, 2017; Rao *et al.*, 2017; Fig. 1). Members of the genus display a diverse spectrum of

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species-specific, sexually selected morphological traits, including ornamentation, coloration, crest morphology, limb length, and body size (Mahony, 2009; Manthey, 2010; Denzer *et al.*, 2016). Such common and highlevels of sexual dimorphisms among congeners were hypothesized to facilitate speciation and to maintain genetic integrity of sympatric species (Wang *et al.*, 2015, 2016). Furthermore, the unique association between the microendemic distributions of congeners and the major mountains and rivers in Asia also make the genus a great system for investigating the biogeographic history of the region (Yang & Rao, 1992; Zhao *et al.*, 1999; Wang *et al.*, 2016).

Yet despite the high diversity in the genus, and its importance in evolutionary and biogeographic studies, little is known about the phylogenetic relationships of Japalura s.l. both within the genus as well as with respect to other genera in the subfamily Draconinae. Although Japalura has long been suspected to be paraphyletic, based on morphological traits (Schmidt, 1927), it was not until 2000 that phylogenetic analyses using genetic data confirmed this hypothesis (Macey et al., 2000). Using mitochondrial DNA (mtDNA) data only, Macey et al. (2000) showed that Japalura s.l. has two, well-supported, deeply divergent clades: the first clade, which is represented by the type species of the genus, Japalura variegata, and J. tricarinata, was inferred to be closely related to members of the genus Draco; and the second clade, which is represented by J. flaviceps and J. splendida, was recovered as a sistergroup to members of the genus *Pseudocalotes* (Macey et al., 2000). Subsequent studies that used the same sequence data of Japalura congeners as Macey et al. (2000) recovered similar phylogenetic relationships among draconine lizards, further highlighting the paraphyletic nature of Japalura sensu lato s.l. (Schulte et al., 2004; Zug et al., 2006).

Analysing a different mtDNA locus of another island endemic species of the genus, Japalura polygonata, Honda et al. (2000) also investigated the phylogenetic position of Japalura in the family Agamidae. Although results of this study failed to infer the exact placement of the Japalura species sequenced with strong support, Honda and colleagues showed that J. polygonata was more closely related to a clade containing the genera Calotes and Aphaniotis than to the clade containing the genera Draco and Mantheyus (Honda et al., 2000). More recently, Pyron et al. (2013) conducted a higher level study of squamate reptiles that incorporated all available mtDNA data from previous studies. Incorporating all previously included species of Japalura s.l. (J. flaviceps, J. polygonata, J. splendida, J. tricarinata and J. variegata), not only did Pyron et al. (2013) recover the same two divergent clades of Japalura observed previously by Macey et al. (2000), but also, they recovered a third distinct clade represented by *J. polygonata*, which is sister to the South-East Asian agamid *Gonocephalus robinsonii* (now recognized as *Malayodracon robinsonii*). However, given the large amount of missing data and non-overlapping gene sampling among different clades, questions remain about the phylogenetic position of this third clade represented by *J. polygonata* (Pyron *et al.*, 2013).

Currently, most authors agree that *Japalura s.l.* is paraphyletic, and the genus s.l. represents multiple, deeply divergent clades; however, no studies have addressed the taxonomic instability of the genus s.l. (Macey et al., 2000; Stuart-Fox & Owens, 2003; Mahony, 2010; Ananjeva et al., 2011; Pyron et al., 2013; Rao et al., 2017). Furthermore, although authors have given names to all three clades recovered from previous phylogenetic studies (including J. veriegata species group/Himalayan Lineage, J. splendida group/ Indochina Lineage and J. polygonata group/Island Lineage), these clade names were established based on five of 34 species of the genus s.l. only, leaving the exact group/lineage memberships of the remaining 29 congeners unknown (Stuart-Fox & Owens, 2003; Mahony, 2010; Ananjeva et al., 2011).

Fortunately, broader genetic and morphological sampling across members of the genus Japalura s.l. has now been amassed, allowing for a more robust phylogenetic assessment of monophyly of the genus and phylogenetic positions of different clades within the genus s.l. with respect to other genera in the subfamily Draconinae. In this study, we employ a multilocus phylogenetic dataset to evaluate the phylogenetic relationship of Japalura s.l. and to address long-standing taxonomic problems at the generic level. Specifically, we are interested in testing the monophyly of Japalura s.l. estimating the number of distinct clades within this genus and testing whether the three proposed species groups based on limited phylogenetic sampling from previous studies represent monophyletic groups. Based on our results, we provide a revised nomenclatural classification scheme that recognizes the species diversity of Japalura s.l. as members of one of four distinct genera, revise the morphological diagnoses for each of the four genera, and present an updated key to all genera in the subfamily Draconinae. Finally, we point to future directions for phylogenetic and taxonomic studies of Japalura s.l. and, more broadly, the subfamily Draconinae in general.

TAXONOMIC HISTORY

The genus *Japalura s.l.* has a long, convoluted taxonomic history based on morphological data. First described by Gray in 1853, the genus *Japalura* was established based on the type species, *J. variegata* from Sikkim, Himalava. Grav (1853) gave a vague definition of his new genus and stated that it was most similar to Agama, but differed by having a concealed tympanum. Ever since the initial description, different species have been placed into the genus Japalura, and numerous authors have attempted to refine the taxonomic definition of the genus based on newly included taxa. For example, Boulenger (1885) first added more diagnostic characteristics to Japalura, including having a compressed body shape, heterogeneous dorsal scales, a dorsal crest, a small gular pouch (or absent), an oblique fold, a feebly compressed tail and no pre-anal or femoral pores. Pope (1935) diagnosed Japalura from other agamid genera by having a body not depressed and possessing hidden tympana. In the same year, Smith (1935) provided a combined version of the diagnoses from Pope (1935) and Boulenger (1885) that was much more comprehensive. However, Smith's (1935) diagnosis of Japalura still cannot differentiate most of its members from other genera in the subfamily Draconinae, as many exceptions exist within the genus s.l. (e.g. J. dymondi, J. kumaonensis, J. major, J. tricarinata and J. varcoae have exposed tympana). Later, Inger (1960) further differentiated Japalura s.l. from the South-East Asian genus Phoxophrys Hubrecht, 1881 by having greatly overlapping supraciliaries, a relatively long head, the presence of hair-like sense organs on the head scales and a compressed tail at the base. However, those diagnostic characters again are not consistent among many of the congeners of Japalura (Ota, 1989b). Subsequent studies largely followed the diagnosis of the genus by Boulenger (1885), but these generic definitions remained ambiguous and impracticable, and failed to differentiate Japalura s.l. from other genera of Draconinae, such as Pseudocalotes (Zhao & Adler, 1993; Kästle & Schleich, 1998; Zhao et al., 1999; Yang & Rao, 2008; Wang et al., 2015).

Such an absence of practical morphological diagnoses of Japalura s.l. has resulted in uncertain systematic affiliations and unstable nomenclatures, not only for members of the genus s.l. but also for species in other genera (Pope, 1935; Smith, 1935; Inger, 1960; Kästle & Schleich, 1998; Yang & Rao, 2008; Mahony, 2010). Species that are currently recognized as members of Japalura s.l. have been confused and assigned to different, recognized agamid genera, including Acanthosaura Gray, 1831 (J. dymondi [Boulenger, 1906]; J. varcoae [Boulenger, 1918]; J. major [Boulenger, 1885]; J. kumaonensis [Annandale, 1907]), Calotes Cuvier, 1817 (J. tricarinata [Blyth, 1853]) and Phoxophrys (J. gramahi [Stejenger, 1924]); and species that are currently recognized in other recognized agamid genera have been placed into Japalura s.l. including Aphaniotis Boulenger, 1885 (Aphaniotis ornata [Lidth

De Jeude, 1893]), Phoxophrys (Phoxophrys nigrilabris [Boulenger, 1885]; P. robinsoni [Boulenger, 1920]; Inger, 1960), and Pseudocalotes Fitzinger, 1843 (Pseudocalotes kaulbacki, [Smith, 1937]; Mahony 2010).

Moreover, several generic names were established, but later synonymized as subjective junior synonyms of *Japalura* based on morphological data alone, including *Diploderma* Hallowell, 1861 (type species *D. polygonontum*, from the Okinawa islands of Japan), *Oriotiaris* Günther, 1864 (type species *O. tricarinata*, from north-east India), and *Oreocalotes* Jerdon, 1870 (type species *O. major*, from the western Himalayas) (Schleich & Kästle, 2002; Mahony, 2010). Nonetheless, the evolutionary independence of most of those genera represented was not examined specifically under a comprehensive morphological or molecular phylogenetic framework (Macey et al., 2000; Pyron et al., 2013; Grismer et al., 2016b).

To date, a total of 34 species are recognized as members of the genus Japalura s.l. in the recent literature (Yang & Rao, 2008; Mahony, 2010; Manthey et al., 2012; Cai et al., 2015; Wang et al., 2016; Ananjeva et al., 2017; Rao et al., 2017), including: J. andersoniana Annandale, 1905; J. bapoensis (Yang, Su & Li, 1979); J. batangensis Li, Deng, Wu & Wang, 2001; J. brevicauda Manthey, Denzer, Hou & Wang, 2012; J. brevipes Gressitt, 1936; J. chapaensis Bourret, 1937; J. dasi (Shah & Kästle, 2002); J. dymondi (Boulenger, 1906); J. fasciata Mertens, 1926; J. flaviceps Barbour & Dunn, 1919: J. grahami (Stejneger, 1924); J. hamptoni Smith, 1935; J. iadina Wang, Jiang, Siler & Che, 2016; J. kumaonensis (Annandale, 1907); J. laeviventris Wang, Jiang, Siler & Che, 2016; J. luei Ota, Chen & Shang, 1998; J. major (Jerdon, 1870); J. makii Ota, 1989; J. micangshanensis Song, 1987; J. ngoclinensis Ananjeva, Orlov & Nguyen, 2017; J. otai Mahony, 2009; J. planidorsata Jerdon, 1870; J. polygonata (Hallowell, 1861); J. sagittifera Smith, 1940: J. slowinskii Rao, Vindum, Ma, Fu & Wilkinson, 2017; J. splendida Barbour & Dunn, 1919; J. swinhonis Günther, 1864; J. tricarinata (Blyth, 1853); J. varcoea (Boulenger, 1918); J. variegata Gray, 1853; J. vela Wang, Pan, Hou, Siler & Che, 2015; J. yunnanensis Anderson, 1878; J. yulongensis Manthey, Denzer, Hou & Wang, 2012; and J. zhaoermii Gao & Hou 2002.

MATERIAL AND METHODS

TAXON SAMPLING

We collected 25 out of 34 species of the genus *Japalura s.l.* from Mainland China, Taiwan and India [Field collection permits (Mainland China: BBCJ-2014-001; Taiwan: Forest Bureau No. 1061700632); Wang *et al.*, 2015, 2016, 2017, 2018a, 2018b; Supporting

Information, Appendix S1]. Lizards were euthanized after capture in the field following the protocols of corresponding institutions in their home countries (USA and Mainland China: IACUC R17-019; Taiwan: IACUC of NTNU No. 106027), and liver or muscle tissues were collected from each individual and preserved in 95% ethanol. Specimens were fixed in buffered 10% formalin and subsequently transferred to 70% ethanol for long-term preservation. Vouchered specimens were deposited in research institutions, including the Museum of Kunming Institute of Zoology, Chinese Academy of Sciences (KIZ), National Museum of Natural Science (NMNS) and Center for Ecological Sciences, Indian Institute of Science (CESG).

Additional tissue samples were also obtained through museum loans from the California Academy of Sciences, USA (CAS), Museum of Comparative Zoology, USA (MVZ) and Sam Noble Oklahoma Museum of Natural History (OMNH) (Supporting Information, Appendix S1). Additionally, we collected novel sequence data for six outgroup species of agamids representing four genera from Mainland China and the Philippines (Supporting Information, Appendix S1), and we incorporated existing genetic sequence data of four species of Japalura s.l. and 17 species of outgroup agamids were obtained from GenBank (Supporting Information, Appendix S2).

MORPHOLOGICAL DATA COLLECTION

We examined a total of 217 specimens representing 27 species of Japalura s.l. and 31 specimens representing 11 species from seven genera of outgroup agamid diversity (Supporting Information, Appendix S3). Following previous studies (Moody, 1980; Mahony, 2010; Grismer et al., 2016a), and our prior knowledge on morphological characteristics of Draconinae (Mahony, 2010; Wang et al., 2018b), we chose a suite of pholidosis characteristics to examine for all specimens, as they are shown to be most useful in delimiting generic-level diversity (abbreviations given in parentheses): presence of modified scales on dorsal body (MDS), defined as presence of scales on dorsal body that are modified in shape, size or texture (e.g. enlarged, keeled, flat or raised in conical shape); degree of modification of dorsal modified scales (DMDS), defined as the level of modification compared to the ground or normal scales on dorsal body, defined as strongly modified or weakly modified; presence of dorsolateral ridges (PDR), defined as presence of regularly and closely arranged, lateral rows of enlarged, keeled dorsal scales on the body; presence of V-shaped ridges on dorsal body (PVR), defined as presence of enlarged, keeled scales that are arranged in V-shape ridges on dorsal body; and presence of head spines (PS), defined as presence of elongated spines on

the post-orbital, occipital and supratympanic regions of head.

In addition, standard morphometric measurements of agamid lizards were taken by KW for a subset of male specimens only (53 specimens representing 21 species of Japalura s.l. and 19 specimens representing eight species from six genera of outgroup taxa within Draconinae) (Supporting Information, Appendix S4). Male specimens only were chosen for the statistical analyses, because males are generally more diverse in morphology; and by choosing one sex only, we remove the confounding impacts from sexual dimorphism from the analyses, which is evident in most Draconinae genera (Zug et al., 2006; Manthey, 2012; Deepak et al., 2016; Wang et al., 2016). The following morphological characters and their measurement methods followed Wang et al. (2016) and included: snout-vent length (SVL), tail length (TAL), head width (HW), head length (HL), snout-eve length (SEL), fore-limb length (FLL), hindlimb length (HLL) and trunk length (TRL). In addition, length of tallest nuchal crest (CL), measured from the base to the tips of the tallest nuchal crest scale, was also recorded. All raw measurements were standardized to relative ratios by either dividing against SVL or HL, and the ratios include HL/SVL, TRL/SVL, TAL/ SVL, FLL/SVL, HLL/SVL, CL/HL, SEL/HL and HW/ HL. Only these resulting ratios were used for statistical analyses (statistical methods, see below).

STATISTICAL ANALYSES FOR MORPHOLOGICAL DATA

Principle Component Analysis (PCA) and Discriminant Analysis of Principle Components (DAPC) were performed on the relative ratios of measurements to determine whether Japalura s.l. occupied unique morphospace with respect to morphologically similar Draconinae genera, and whether the morphological clustering coincided with the phylogenetic clades recovered from our molecular dataset. PCA was performed on the relative ratios of continuous morphometric measurements, using the prcomp command in R, v.3.2.1. All PCA data were natural log transformed prior to analysis and scaled to their standard deviation in order to normalize their distribution. DAPC were performed on the principle components with eigenvalues greater than one (Kaiser, 1960), using R, v.3.2.1 (Supporting Information, Appendix S5).

MOLECULAR DATA COLLECTION

Total genomic DNA was extracted from liver or muscle tissues using Fujita's Guanidine Thyocyanate protocol (Esselstyn *et al.*, 2008). We targeted three nuclear loci, including Brain Derived Neurotrophic Factor (*BDNF*), Oocyte Maturation Factor (*CMOS*) and RNA Fingerprint Protein 35 (*R35*), and one mitochondrial gene, NADH dehydrogenase subunit 2 (*ND2*). As a result of amplifying *ND2*, we also obtained a short fragment of the cytochrome *c* oxidase subunit I (*COI*) gene and components of eight flanking transfer RNA genes (tRNA^{GIn}, tRNA^{IIe}, tRNA^{Met}, tRNA^{Trp}, tRNA^{AIa}, tRNA^{Asn}, tRNA^{Cys} and tRNA^{Tyr}). Genes were targeted and amplified using both published and newly designed primers and PCR protocols (Table 1).

Amplified products were visualized on 1.5% agarose gels, and clean, single-banded products were purified with 1 μ L of a 20% dilution of ExoSAP-IT (US78201, Amersham Biosciences). Cycle sequencing reactions were carried out using ABI Prism Big-Dye Terminator chemistry (Ver. 3.1; Applied Biosystems), and purified with Sephadex (NC9406038, Amersham Biosciences) in Centri-Sep 96 spin plates (CS-961, Princeton Separations). Purified products were sent to Eurofins Genomics for analysis. All novel sequences were deposited in GenBank (Supporting Information, Appendix S1).

ALIGNMENT AND PHYLOGENETIC ANALYSIS

Continuous gene sequences were assembled and edited using GENEIOUS, v.10.0.6. Alignments were produced initially using MUSCLE (Edgar, 2004), with minor subsequent manual adjustments. Alignments and resulting topologies are deposited in Data Dryad (doi:10.5061/dryad.683h4j5). To assess phylogenetic congruence between the mitochondrial and nuclear data, we inferred the phylogeny for each gene independently using likelihood and Bayesian analyses. As no strongly supported incongruences between mitochondrial and nuclear data were observed, we concatenated the two datasets for the final analyses.

We conducted concatenated, partitioned Bayesian analyses in the program MRBAYES, v.3.2.1 (Ronquist & Huelsenbeck, 2003). All protein-coding genes were partitioned by codon positions, except for COI, which is treated as a single partition due to its short length (<40 bp), and the mitochondrial tRNAs were combined and treated as a single partition as well (Table 2). The best models of nucleotide substitution were selected for each partition by the Akaike Information Criterion (AIC), as implemented in JMODELTEST2, v.2.1.10 (Guindon & Gascuel, 2003; Darriba et al., 2012). A rate multiplier model was used to allow substitution rates to vary among subsets. Default settings were used for all other model parameters. Two independent Markov chain Monte Carlo analyses were run, each with four Metropolis-coupled chains, a melting temperature of 0.02 and an exponential distribution with a rate parameter of 25 as the prior branch lengths (Marshall, 2010). All Bayesian analyses were run for 6000000 generations, with parameters and topologies sampled every 3000 generations. Stationarity and convergence were assessed with TRACER, v.1.6.0 (Rambaut, Suchard & Drummond, 2013), and we confirmed convergence of tree splits with AWTY (Nylander et al., 2008). Although runs reached stationarity after 5 million generations, we conservatively discarded the first 15% of samples as burn in, resulting in a total of 1700 topologies from the posterior distribution for both runs.

Partitioned Maximum Likelihood analyses were conducted using RAxML-VI-HPC, v.8.2.10 (Stamatakis, 2014) on the concatenated dataset using the same partitioning strategy as for Bayesian analysis. The more complex model (GTR + Γ) was applied for all subsets (Table 2), with 1000 replicate ML inferences run. Each

Table 1.	Summary of	primers and	annealing	temperatures	employed in	this study

Gene	Primer name	Primer sequence	Annealing tempera- ture (°C)	Referred study
ND2	L4160	CGATTCCGATATGACCARCT	52	Macey <i>et al.</i> , 2000
	H4980	ATTTTTCGTAGTTGGGTTTGRTT	52	Macey et al., 2000
	Jap_264F	GGGCCCATACCCCRAAAAC	55	Present study
	Jap_1559R	GGATTAATGCCCTCTGGATT	55	Present study
	L4437b_F	AAGCAGTTGGGCCCATACC	55	Macey et al., 2000
	$H5540_R$	TTTAGGGCTTTGAAGGC	55	Macey et al., 2000
	Jap_70F	CCACCAAACAACTACACCTA	55	Present study
	Jap_1559R	GGATTAATGCCCTCTGGATT	55	Present study
BDNF	BDNF.F	CCCCAATGAAAGAAGTGASCCTC	55	Crottini et al., 2009
	BDNF.R	TGGGTAGTTCGGCACTGAGAATTCC	55	Crottini et al., 2009
CMOS	G73.1	GGCTRTAAARCARGTGAAGAAA	55	Whiting et al., 2003
	G74.1	GARCWTCCAAAGTCTCCAATC	55	Whiting et al., 2003
R35	R35.F	GACTGTGGAYGAYCTGATCAGTGTGG	55	Whiting et al., 2003
	R35.R	GCCAAAATGAGSGAGAARCGCTTCTG	55	Whiting et al., 2003

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Gene	Partition	AIC model	Number of characters (bp)
BDNF	1st codon	GTR+Γ	176
	2nd codon	HKY	176
	3rd codon	ΗΚΥ+Γ	175
CMOS	1st codon	$JC+\Gamma$	181
	2nd codon	ΗΚΥ+Γ	180
	3rd codon	HKY	180
R35	1st codon	ΗΚΥ+Γ	213
	2nd codon	GTR	213
	3rd codon	$HKY+\Gamma$	213
tRNAs (Gln, Ile, Met, Trp, Ala, Asn, Cys, Tyr)	Combined, single partition	$GTR+\Gamma$	613
ND2	1st codon	$GTR+\Gamma$	345
	2nd codon	$GTR+\Gamma$	345
	3rd codon	$GTR+\Gamma$	345
COI	Single partition	$GTR+\Gamma$	33

Table 2. Models of evolution selected by Akaike Information Criterion (AIC) and applied for partitioned, phylogenetic analyses

inference was initiated with a random starting tree, and nodal support was assessed with 1000 bootstrap pseudoreplicates (Stamatakis, Hoover & Rougemont, 2008).

To root the resulting phylogenetic trees, *Hydrosaurus pustulatus* (subfamily Hydrosaurinae) was chosen as the outgroup of subfamily Draconinae, based on published higher level phylogenetic studies of squamate reptiles (Pyron *et al.*, 2013).

HYPOTHESIS TESTING

We tested taxonomic hypotheses proposed in previous studies to address the following questions: (1) Does our expanded dataset support the previous shown relationship between Malayodracon and island species of Japalura s.l. by Pyron et al. (2013)? (2) With expanded taxa sampling, does our dataset support the previous shown sister relationship between the genus Calotes and Salea by Grismer et al. (2016b)? (3) With the exclusion of the controversial species, J. bapoensis, do congeners from each of the three general geographic regions (the Himalayas, Indochina/Mainland East Asia and East Asian Islands) form a monophyletic group, as suggested by previously proposed species groups (e.g. Himalayan Species Group, Indochinese Species Group and Island Species Group)? (4) Does J. bapoensis form a monophyletic group with remaining species of Japalura s.l. in mainland East Asia?

To do so, we estimated posterior probabilities of different topologies that correspond to each of the above question using a Bayesian approach. We constrained corresponding topologies for each hypothetical scenario and estimated the posterior probability of supporting such hypotheses using the proportion of post-burn-in trees that are consistent with the corresponding hypothesized topology. This was accomplished by filtering the total number of post-burn-in trees for each constrained topology using PAUP*, v.4.0b10 (Swofford, 1999).

RESULTS

PHYLOGENETIC RESULTS

We did not observe any incongruent clades with strong statistical support [≥ 0.95 posterior probability (PP) or ≥ 70 bootstrap support (BS)] between nuclear and mitochondrial gene trees. Therefore, we felt justified in concatenating all data into one matrix, containing 3388 nucleotides for subsequent analyses. Characteristics of nucleotides, as well as the inferred best-fit models of sequence evolution selected by Akaike Information Criterion (AIC), are summarized in Table 2.

The results of Maximum Likelihood and Bayesian analyses are largely consistent with each other. However, the placements of an outgroup (Acanthosaura *lepidogaster*) are different between the two analyses (in ML analyses, Acanthosaura is recovered as sister to the clade containing Malayodracon, Pseudocalotes, Sitana and clade O of Japalura s.l. (BS = 53; Fig. 2). The subfamily Draconinae is recovered with strong support, Mantheyus phuwuanensis as the first branching lineage [1.00/100 (PP/BS, respectively, herein noted in the same order)]. Both analyses support the same relationships among focal clades with strong support $(\geq 0.99 \geq 80)$, with the results supporting the paraphyly of Japalura s.l. Four divergent clades are recovered within the Japalura s.l. (Fig. 2, clades C, G, N and O): (1) J. andersoniana, J. kumaonensis, J. tricarinata and

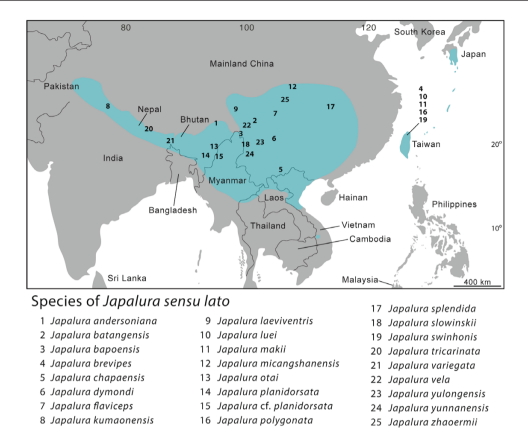


Figure 1. Distribution of *Japalura s.l.* in Asia. Shaded area (light blue) represents the estimated distribution of *Japalura s.l.* (the isolated record in south-central Vietnam represents the questionable taxon *Japalura ngoclinensis*; see Discussion); solid black lines represent national boundaries (political boundaries along the Himalaya are not included to accommodate disputed, political issues surrounding this region); and numbers represent geographic locations of sampled species from this current study, each corresponding to the listed taxon in the figure key shown below the map.

J. variegata form a monophyletic group with strong support (Fig. 2, clade C, 1.00/83), which form a strongly supported group with *Draco* and *Ptyctolaemus* (Fig. 2, clade A, 1.00/91); (2) J. otai and J. planidorsata form the second, more distantly related clade (Fig. 2, clade G, 1.00/100) that is sister to the genus *Salea* (0.99/69; Fig. 2); (3) J. bapoensis is nested within *Pseudocalotes* with strong support (Fig. 2, clade M, 1.00/100); and (4) all the remaining species from Mainland China and East Asian islands form a monophyletic group (Fig. 2, clade O, 1.00/100) that is sister to the genus *Pseudocalotes* (1.00/100). Furthermore, two well-supported sub-clades (Fig. 2, sub-clade P, 1.00/100; and sub-clade Q, 1.00/100) were recovered within clade O from Mainland China and East Asian Islands (Fig. 2).

Although population sampling is not the focus of this paper, we included multiple populations of the same species when samples were available. At the species level, most species are recovered as monophyletic lineages, except *J. planidorsata* and *J. flaviceps* in clades G and O, respectively (Fig. 2). The sampled Myanmar individuals of *J. cf. planidorsata* are paraphyletic with respect to *J. otai*, and the GenBank sequence of *J.* cf. *flaviceps* is not recovered with topotypic individuals of the species, but instead is supported as part of a monophyletic group with topotypic material of *J. zhaoermii* (Fig. 2).

MORPHOLOGICAL RESULTS

For continuous morphometric data, PCA and DAPC analyses show members of *Japalura s.l.* occupy distinct regions of morphospace compared to other genera (Fig. 3). Furthermore, four groups were recovered within *Japalura s.l.* in morphospace, which coincide with the four major clades of the genus *s.l.* recovered from the genetic data (Fig. 3). For PCA analysis, the first four components account for 78% of the total variance: PC1 loads most heavily on relative fore-limb (FLL/SVL) and hind-limb length (HLL/SVL) (25% of the total variance); PC2 loads most heavily on relative crest length (CL/HL) and snout length (SEL/HL) (23% of total variance); PC3 loads most heavily on relative trunk length (TRL/SVL) and head length (HL/SVL)

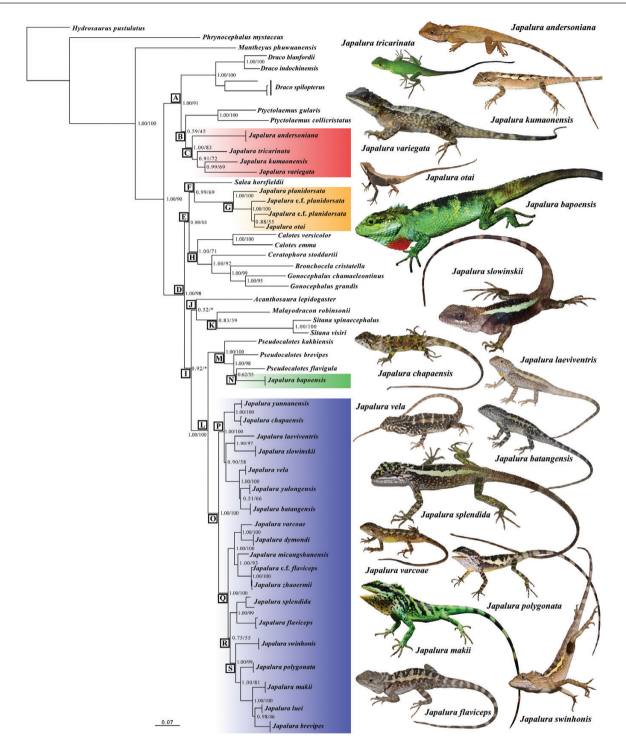


Figure 2. Phylogenetic relationships among *Japalura sensu lato* based on both Maximum Likelihood and Bayesian Analyses of two mitochondrial genes (*COI* and *ND2*), tRNAs and three nuclear genes (*BDNF*, *CMOS*, and *R35*). Maximum Likelihood bootstrap and Bayesian posterior probability values are included at all nodes, except (1) terminal nodes that unify multiple individuals of the same species [which all have 1.00/100 support (Bayesian/Maximum Likelihood)]; and (2) non-conflicting (either kind of analysis yielded significant supports) yet inconsistent nodes between two types of analyses (e.g. nodes unifying clades I and J), in which only the Bayesian posterior probability is given. Photos of selected species of *Japalura s.l.* are included (scaled to reflect relative sizes of each species) with their current taxonomic names (vs. revised taxonomic names, summarized in taxonomic accounts in the Discussion).

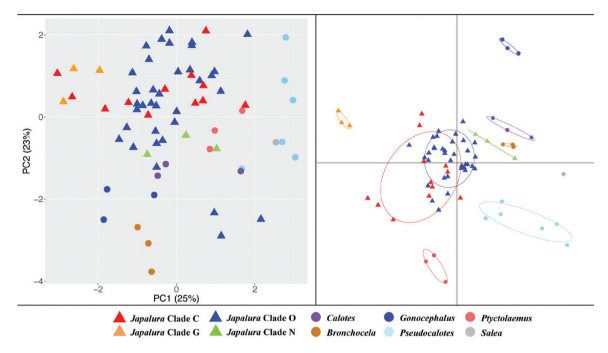


Figure 3. Principle component analysis (PCA) and discriminant analysis of principle component (DAPC) of continuous morphometric measurements and ratios of *Japalura s.l.* and morphologically similar genera. Colour code of each clade of *Japalura s.l.* coincides with the one used for the same clade in Figure 2.

(16% of the total variance); and PC4 loads most heavily on relative tail length (TAL/SVL) and head length (HL/ SVL) (14% of the total variance) (Table 3; Supporting Information, Appendix S4). The DAPC further characterize these results and show the visual clustering of four different clades of *Japalura s.l.* in morphospace (Fig. 3). In addition, each of the four genetic clades/ morphological clusters can also be diagnosed readily using a combination of discrete pholidosis characteristics (Fig. 4; Table 3).

TOPOLOGY TESTING

Our topological tests do not support the monophyly of the clade containing the East Asian Japalura s.l. and Malayodracon robinsonii (H1, PP = 0.00) and the clade containing Calotes and Salea (H2, PP = 0.00) (Table 4). For Japalura s.l. species from the three geographic regions, with the exclusion of the controversial species, J. bapoensis, the Bayesian statistical tests show no support for the monophyly of the Himalayan species group (H3, PP = 0.00), nor the monophyly of the remaining mainland Indochinese species group (H4, PP = 0.00). Only the monophyly of East Asian species (together both mainland and island species) is statistically supported (H5, PP = 1.00). As for the controversial species, J. bapoensis, we found no statistical support for its monophyly with respect to the remaining East Asian species (H6, PP = 0.00) (Table 4).

DISCUSSION

NEW CLASSIFICATION OF JAPALURA S.L.

Our morphological and phylogenetic results both support observations of paraphyly among members of *Japalura s.l.* in previous studies (Figs 2, 3; Table 3; Macey *et al.*, 2000; Mahony, 2010). We recover two previously recognized clades in similar phylogenetic positions in the inferred topologies (*J. variegata* clade and *J. splendida* clade), and we confirm the generic placement of *J. bapoensis* in the genus *Pseudocalotes* as in previous studies (Macey *et al.*, 2000; Schulte *et al.*, 2004; Zug *et al.*, 2006; Pyron *et al.*, 2013; Grismer *et al.*, 2016b; Fig. 2). Furthermore, with the exclusion of *J. bapoensis*, this study supports the previously assumed monophyly of all East Asian species, including both mainland and island lineages (H5, PP = 1.00; Fig. 2, clade O, 1.00/100; Table 4; Ananjeva *et al.*, 2011).

In addition to observing consistent patterns with previous research, our results reveal novel relationships of draconine genera with the expanded taxonomic and genetic coverage. First, analyses recover two distinct clades from the previously proposed Himalayan species group: clade C, which corresponds to the previously discovered Himalayan clade by Macey *et al.* (2000) and Pyron *et al.* (2013), and clade G, revealing relationships observed for the first time (Fig. 2). Although species of both clades are from the Himalayan region, they are recovered in distantly

Characters	Japalura s.l.	·			Outgroup genera	enera			
	Clade C	Clade G	Clade O	Clade N	Calotes	Gonocephalus	Malayodracon	Pseudocalotes	Salea
SSDB	HT	HT	HT	HT	HM	HT	HT	HT	HT
AESD	DLR & VR	DLR & VR	Irregular	DLR	/	Irregular	Irregular	Irregular	Irregular
ECSN	A	Р	A	A	A	A	A	A	A
KSL	К	К	ß	К	K	K	К	S	S
SOR	Multiple	Multiple	Singular	Multiple	Multiple	Multiple	Singular	Mostly singular	Singular
RSSOR	Subequal	Subequal	N/A	Subequal or one row	Subequal	Subequal	N/A	N/A or one row	N/A
				enlarged				enlarged	
HS	A	A	A	A	Р	A	A	A	A
SCS	LT	LT	\mathbf{TT}	LT	L	L	TT	TT or L	L
GP	FB or WD	FB	WD	FB or WD	WD	WD	WD	WD	WD

related groups within Draconinae with strong support (Fig. 2). Statistical topology tests further support the phylogenetic results, rejecting the assumed monophyly of the Himalayan congeners (PP = 0.00; Table 4; Stuart-Fox & Owens, 2003; Mahony, 2010; Ananjeva *et al.*, 2011; Wang *et al.*, 2018b). Second, we reject the previously hypothesized sister relationship between island *Japalura* species and *Malayodracon* (Pyron *et al.*, 2013). The phylogenic results support a closer relationship between East Asian species of *Japalura s.l.* (Mainland and Island) and *Pseudocalotes* (Fig. 2, clade L), and topology tests reject the null hypothesis of reciprocal monophyly between island *Japalura s.l.* and *Malayodracon* (Table 4).

The recognition of all four clades of *Japalura s.l.* as a single genus would render most genera of the subfamily Draconinae paraphyletic, thus, further complicating an existing, long-held taxonomic problem. Therefore, in an effort to revise taxonomy to best reflect evolutionary histories of each clade of *Japalura s.l.* in nomenclature, we recognize four genera that each corresponds to the evolutionary clade recovered in our phylogenetic and morphological analyses.

The first clade (Figs 2, 3, clade C) includes the type species of three currently or historically recognized generic names, namely Japalura sensu stricto Gray (1853) (type species J. variegata), Oriotiaris Günther (1864) (type species O. tricarinata) and Oreocalotes Jerdon, 1870 (type species O. major) (Fig. 2). However, we recognize Japalura sensu stricto to represent this clade for the following reasons: (1) Japalura sensu stricto has temporal priority over the other two generic names; (2) recognizing the synonym Oriotiaris as a valid genetic name would render itself and Japalura sensu stricto both paraphyletic (Fig. 2); (3) for Oreocalotes, although we do not have access to the genetic materials of the type species, the morphology of the type species (O. major or J. major as currently recognized) falls within the morpho-cluster of species of Japalura sensu stricto, and the type species is morphologically most similar to J. kumaonensis (Wang et al., 2018b), which is a confirmed member of Japalura sensu stricto (Figs 2, 3). Therefore, we treat both Oriotiaris and Oreocalotes as junior synonyms of Japalura sensu stricto.

Of the remaining three clades (clades G, N and O; Fig. 2), for clade N, we follow Mahony (2010) and transfer Japalura bapoensis to the genus Pseudocalotes, and conservatively consider it as a subspecies of P. kingdonwardi (see discussion below on its species status). Second, we resurrect the only available generic name in the literature, Diploderma, for clade O (with its type species D. polygonatum polygonatum Hallowell, 1861). Lastly, as clade G has no generic name available in the literature, we describe it as a new genus, Cristidorsa gen. nov. (described below). Based on the revised

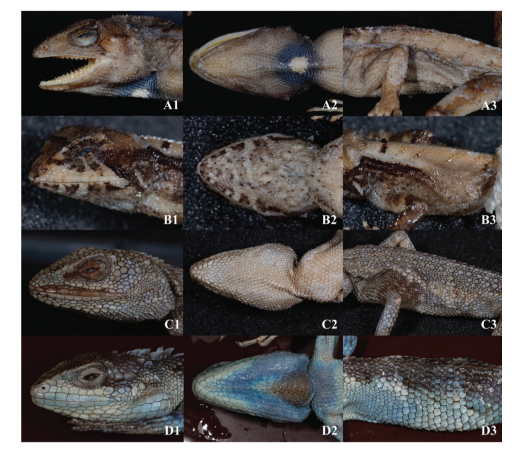


Figure 4. Close-up photographs of the head in lateral (1), and ventral (2), views, and body in lateral view (3), of representative species (A–D) of the four clades of *Japalura s.l.* recovered in phylogenetic and morphological analyses, showing diagnostic pholidosis characteristics: A, clade C: *Japalura andersoniana*, KIZ 011156; B, clade G: *Japalura cf. planidorsata*, CAS 233296; C, clade O: *Japalura polygonata*, MCZ 21244; and, D, clade N: *Japalura bapoensis*, CAS 241965.

Hypothesis	Description of constraint	Posterior probability
Does <i>Malayodracon</i> form monophyletic group w	th island Japalura s.l.?	
H1 Monophyly of island Japalura + Malayodracon	All sampled <i>Japalura</i> from Taiwan and <i>Malayodracon robinsonii</i>	0.00
Does Calotes form a sister relationship with Sal	ea?	
H2 Monophyly of the clade containing Calotes + Salea	All sampled species of <i>Calotes</i> and <i>Salea</i>	0.00
With the exclusion of Japalura bapoensis, do con	ngeners from each of three regions form a monophyl	etic group?
H3 Monophyly of Himalayan species group	All sampled species from the Himalayan region	0.00
H4 Monophyly of Indochinese species group	All sampled species from mainland east Asia	0.00
H5 Monophyly of Indochinese species + island species	All sampled species from mainland east Asia and Taiwan	1.00
Does Japalura bapoensis belong to the east Asia	n clade of <i>Japalura s.l.</i> ?	
H6 Monophyly of <i>J. bapoensis</i> + remaining Indochinese + island species	All sampled species from mainland east Asia (including <i>J. bapoensis</i>) and Taiwan	0.00

Table 4. Bayesian topology tests conducted for previously hypothesized relationships of *Japalura s.l.* and other genera within the subfamily Draconinae

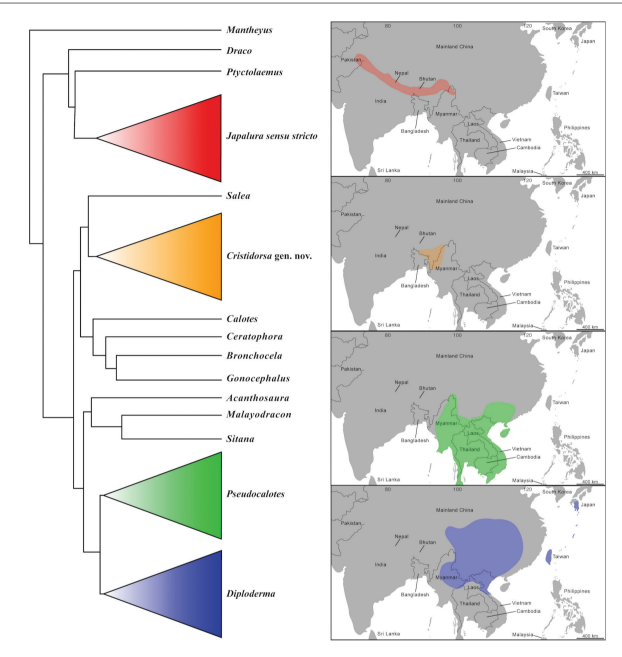


Figure 5. Simplified, inferred topology summarizing the new proposed classification scheme and recognized geographic distributions of the four focal clades of *Japalura s.l.* (*Japalura*, **Cristidorsa gen. nov.**, *Pseudocalotes* and *Diploderma*).

taxonomy and associated morphological diagnoses summarized in this study for each of the four clades, we propose a new classification scheme for all recognized members of *Japalura s.l.*

TAXONOMIC ACCOUNTS JAPALURA GRAY, 1853

Etymology: The Latin name '*Japalura*' may be derived from a locality name in India, and the term is feminine

gender. We suggest the English common name as 'Himalayan Dragon', and the Chinese name as '攀蜥' (pronounced as 'Pan-Xi').

Type species: Japalura variegata, Gray, 1853.

Diagnosis: Lizards of the genus *Japalura sensu stricto* differ from closely related genera by possessing the following morphological characteristics: (1) head width moderate, HW mostly <70% of HL; (2) nuchal and dorsal crest scales relatively low and thick, not significantly elongated into lanceolate spines, CL/HL<10%;

(3) post-orbital and post-occipital spines absent; (4) gular scales mostly homogeneous in size; (5) size of scales on lateral jaw subequal in size across gular region; (6) dorsal scale significantly heterogeneous in size and shape, not regularly imbricate; (7) pair of distinctively enlarged, conical scales on nape above shoulder absent; (8) paravertebral rows of enlarged scales present on dorsolateral body; and (9) V-shaped ridges present along dorsal midline, formed by enlarged, keeled scales.

Phylogenetic definition: We define Japalura sensu stricto using the maximum crown-clade definition, which includes species that share a more recent common ancestor to Japalura veriegata than Draco volans or Ptyctolaemus gularis.

Included species: Based on our phylogenetic results, we assign the following species to the genus Japalura sensu stricto: J. andersoniana, J. kumaonensis, J. tricarinata and J. variegata. Following our morphological results and proposed morphological diagnoses, we also assign J. dasi, J. major and J. sagittifera into this genus, pending future phylogenetic studies.

Geographic distribution: Members of the genus are distributed along the southern foothills of the Himalayas, including north-eastern Pakistan (J. kumaonensis and J. major), northern and north-eastern India (J. andersoniana, J. kumaonensis, J. major, J. sagittifera, J. tricarinata and J. variegata), Nepal (J. dasi, J. tricarinata and J. variegata), Bhutan (J. andersoniana, J. tricarinata and J. variegata), southern parts of the Tibet Autonomous Region of China (J. andersoniana and J. tricarinata), and northern part of Myanmar (J. sagittifera). Congeners may also be found in north-western Bangladesh (Fig. 5).

CRISTIDORSA

WANG, DEEPAK, DATTA-ROY, LIN, JIANG, CHE & SILER GEN. NOV.

Etymology: The Latin term '*Cristidorsa*' means 'ridged dorsum', which describes the distinct, characteristic ridges on the dorsal surface of the body in the new genus. The generic name is feminine and it consists of two parts, namely '*Cristi-*' (meaning 'ridged') and '-*dorsa*' (meaning 'dorsum'). We suggest the English common name as 'Ridged Dragons' and the Chinese name as '棱背蜥' (pronounced as 'Leng-Bei-Shi').

Type species: Cristidorsa otai (Mahony, 2009).

Diagnosis: Lizards of the genus 'Cristidorsa' differ from other closely related Draconinae genera by having the following morphological characteristics: (1) head robust and relatively wide, HW/HL mostly >70%; (2) scales of lateral head keeled; (3) nuchal and dorsal crest feeble, CL/HL<5%; (4) post-occipital and post-orbital spines absent; (5) gular scales mostly homogeneous in size; (6) size of scales on lateral jaw subequal in size across gular region; (7) single pair of distinctively enlarged conical scales present on nape above shoulder, one scale on each side of vertebral crest; (8) dorsal scale significantly heterogeneous in sizes and shapes; (9) distinct, dorsolateral rows of enlarged, keeled scales present on dorsum; and (10) V-shaped ridges present along dorsal midline, formed by enlarged, keeled scales.

Phylogenetic definition: We define *Cristidorsa* using the maximum crown-clade definition, which includes species that share a more recent common ancestor with *Cristidorsa otai* than with *Salea horsfieldii*.

Included species: Based on our phylogenetic results, we assign *C. otai* and *C. planidorsata* to the genus *Cristidorsa*.

Geographic distribution: Members of the genus *Cristidorsa* are distributed on the south-east extreme of the Himalayan foothills, including north-east India *(C. otai and C. planidorsata)*, and the north-west part of Myanmar [*C. otai*; see discussion on taxonomic status of the Myanmar population below (Fig. 5)].

DIPLODERMA HALLOWELL, 1861

Etymology: The Latin generic name 'Diploderma' consists of two parts, 'Diplo-' means 'double' or 'many', and '-derma' means 'skin', and the whole word is in a neuter gender. As the previous generic name 'Japalura' and most species names of the genus s.l. are feminine, most names of species that are now assigned to Diploderma need their gender changed to neutral (except for existing neutral-gender names like brevipes or flaviceps, Latin nouns like vela or names derived from peoples' names, i.e. dymondi, luei, makii, swinhonis, varcoae and zhaoermii). We suggest the English common name of the genus as 'Mountain Dragon', and the Chinese common name as '龙蜥' (pronounced as 'Long-Xi').

Type species: Diploderma polygonatum Hallowell, 1861.

Diagnosis: Lizards of the genus *Diploderma* differ from closely related genera by having the following morphological characteristics: (1) scales of lateral head keeled; (2) nuchal and dorsal crest scales relatively short and thick, not elongated into lanceolate spines, CL/HL mostly <10%; (3) post-occipital and post-orbital spines absent; (4) gular scales mostly homogeneous in size, not decreasing in size toward the centre; (5) scales on lateral jaw subequal in size across gular region; (6) dorsal scale significantly heterogeneous in size and shape, not regularly imbricate; (7) paravertebral dorsolateral ridges of body present in most species, formed by enlarged, keeled scales (except in *D. swinhonis* and *D. leui*); and (8) V-shaped ridges along dorsal body midline absent in all but one species (except in *D. swinhonis*).

Phylogenetic definition: We define Diploderma using the maximum crown-clade definition, which includes species that share a more recent common ancestor with Diploderma polygonatum than with Pseudocalotes tymanistriga and Acanthosaura lepidogaster.

Included species: Based on our phylogenetic results, we assign the following species into the genus Diploderma: D. batangense, D. brevipes, D. chapaense, D. dymondi, D. flaviceps, D. laeviventre, D. luei, D. makii, D. micangshanense, D. polygonatum (and all of its subspecies), D. slowinskii, D. splendidum, D. swinhonis, D. varcoae, D. vela, D. yulongense, D. yunnanense and D. zhaoermii.

According to our proposed morphological diagnoses, we also assign *Diploderma brevicaudum*, *D. fasciatum*, *D. grahami*, *D. hamptoni* and *D. iadinum* to this genus, pending on future phylogenetic confirmations. In total, 24 species of *Japalura s.l.* are reclassified into the genus *Diploderma* (for discussion on the taxonomic status of *D. ngoclinense* see below).

Geographic distribution: Members of the genus are distributed across mainland East Asia, East Asian islands and northern Indochina, including central, southern and south-western Mainland China (D. batangense, D. brevicaudum, D. chapaense, D. dymondi, D. fasciatum, D. flaviceps, D. grahami, D. iadinum, D. laeviventre, D. micangshanense, D. slowinskii, D. splendidum, D. vela, D. yulongense, D. yunnanense and D. zhaoermii) and Taiwan (D. brevipes, D. luei, D. makii, D. polygonatum and D. swinhonis), the southern islands of Japan (D. polygonatum), central and eastern Myanmar (D. hamptoni), northern Vietnam (D. fasciatum and D. chapaense) and northern Thailand (D. yunnanense). Congeners such as D. yunnanense may also be distributed in Myanmar and northern Laos close to the Chinese border (Fig. 5).

PSEUDOCALOTES FITZINGER, 1843

Type species: Pseudocalotes tympanistriga (Gray, 1831).

Diagnosis: Pseudocalotes can be diagnosed from other Draconinae genera by having: (1) scales of

lateral head smooth, especially loreal and subocular region; (2) scales of lateral jaw enlarged; (3) scales of central gular region distinctively smaller than others of throat; (4) sub-orbital scale row usually one, or multiple but with the middle one significantly enlarged; (5) post-orbital and post-occipital spines absent; (6) nuchal crest scales mostly elongated and tall, CL/HL mostly >10%; (7) dorsal crest scales feeble and low; (8) dorsal body scales heterogeneous in size and shape; (9) enlarged dorsal body scales not arranged in paravertebral, dorsolateral or V-shaped ridges.

Phylogenetic definition: We define Pseudocalotes using the maximum crown-clade definition, which includes species that share a more recent common ancestor with Pseudocalotes tympanistriga than with Diploderma polygonatum and Acanthosaura lepidogaster.

Included species: A single species of Japalura s.l. *Pseudocalotes kingdonwardi bapoensis*, is reclassified into the genus *Pseudocalotes*. Currently, the genus includes 22 recognized species in total (Grismer *et al.*, 2016a; Harvey *et al.*, 2017).

Geographic distribution: The genus is distributed in South-East Asia and Indochina, including southern Mainland China, Cambodia, Indonesia, Laos, Malaysia, Myanmar, Sumatra, Thailand and Vietnam (Fig. 5).

PHYLOGENETIC RELATIONSHIPS OF JAPALURA S.L.

Generic-level relationships

The results of this comprehensive phylogenetic analysis of the genus *Japalura s.l.* confirm previous observations of paraphyly (Fig. 2; Macey et al., 2000; Schulte et al., 2004; Zug et al., 2006; Pyron et al., 2013). Although we recover four distinct genera within Japalura sensu lato: Japalura s.l.: Cristidorsa, Pseudocalotes and Diploderma (clades C, G, N and O, respectfully; Fig. 2), and strongly support the monophyly of Japalura sensu stricto, we cannot resolve the intergeneric relationships among Japalura sensu stricto, Draco and Ptyctolaemus (clade B; 0.587/45; Fig. 2). This result is consistent with previous studies, which have recovered Japalura sensu stricto both as sister to Draco (Schulte et al., 2004) or Ptyctolaemus (Pyron et al., 2013). Therefore, at this time, no conclusion can be made with confidence regarding the intergeneric relationships among the three genera-future

studies with expanded taxonomic and genomic datasets are needed.

Without taxonomic representation of the new genus Cristidorsa, previous genomic studies supported a sister relationship between Calotes and Salea, and together, this clade was observed to be reciprocally monophyletic with a clade consisting of Bronchocela, Ceratophora, Cophotis, Gonocephalus and Lyriocephalus (Grismer et al., 2016b). In contrast, with our expanded taxon sampling, our data support the new genus Cristidorsa (Fig. 2, clade G) as the sister-group to Salea (Fig. 2, clade F, 0.99/69), and together with Bronchocela, Calotes, Ceratophora and Gonocephalus, forms a large and diverse clade (Fig. 2, clade E, 0.985/53). Furthermore, our Bayesian topology tests reject the hypothesized monophyly of a Calotes + Salea clade (Table 4, H2, PP = 0.00). We suspect that incomplete taxonomic sampling may have resulted in the relationships recovered in previous studies (Grismer et al., 2016b). However, continued research on this diverse group of agamid lizards is needed.

Using a supermatrix of 12 genetic loci, the resulting topology of Pyron et al. (2013) did not show support for the monophyly of the sampled East Asian species of Japalura s.l. The sampled East Asian mainland species (D. flaviceps and D. splendidum; both were recognized as *Japalura* at the time) were not the closest sister-group to the East Asian island species, D. polygonatum (recognized as Japalura polygonata at the time; Pyron et al., 2013). Instead, Pyron et al.'s (2013) results supported D. polygonatum as most closely related to Malayodracon robinsonii (recognized as Gonocephalus robinsonii at the time) from South-East Asia. However, it should be noted that the only available mitochondrial loci at the time for the mainland and island species were non-overlapping [ND2 for mainland species (D. flaviceps and D. splendidum) and 16S and 12S for island species (D. polygonatum and Malayodracon robinsonii); Pyron et al., 2013]. Additionally, it is recognized that D. polygonatum is morphologically distinct from Malayodracon and distributed in distinct biogeographic regions (Denzer et al., 2016; Fig. 2). Therefore, the unexpected relationships observed previously may be an artefact of incomplete gene coverage. With expanded, overlapping genetic datasets across sampled Draconinae genera, our results recover different relationships from those observed in Pyron et al. (2013). Although all island species of Diploderma from Taiwan and Japan are recovered as a monophyletic group (Fig. 2, clade S), they are nested within a larger clade containing species from mainland East Asia (Fig. 2, clade O), with the whole group as sister to *Pseudocalotes* with strong statistical support (1.00/100; Fig. 2). Additionally, our results do not support *Malayodracon* as sister to *Diploderma polygonatum*, with the null hypothesis of this sister relationship rejected with strong statistical support (Table 4).

For the generic placement of J. bapoensis (Yang, Su & Li, 1979; Yang & Rao, 2008; Mahony, 2010; Cai et al., 2015), analyses support the recent generic designation of this species in the genus *Pseudocalotes* (Mahony, 2010). First described as a subspecies of Pseudocalotes kingdonwardi (considered as Calotes kingdonwardi at the time: Smith, 1935; Yang et al., 1979; Zhao et al., 1999), P. k. bapoensis was later transferred to the genus Japalura based on morphological characteristics and subsequently elevated to full species status as J. bapoensis (Yang & Rao, 2008). However, the morphological diagnosis of the genus Japalura provided by Yang & Rao (2008) (e.g. heterogeneous dorsal scales, well-developed nuchal crests, presence of shoulder fold, feebly developed gular pouch, and small and exposed tympanum) cannot differentiate Japalura s.l. from Pseudocalotes (Mahony, 2010). In contrast, the proposed morphological characteristics of J. bapoensis match the revised diagnoses of *Pseudocalotes*, including having a single, enlarged suborbital scale row, distinctively smaller scales in the center of the throat, enlarged scales toward the lateral jaw, and distinctively large nuchal crest scales (Fig. 3; Yang & Rao, 2008; Mahony, 2010). Furthermore, the results of phylogenetic analyses in this and previous studies recover J. bapoensis as nested within the genus Pseudocalotes with strong support (J. bapoensis was treated as Pseudocalotes kingdonwardi in Grismer et al., 2016b) (Fig. 2, clade M, 1.00/100). Therefore, on the basis of morphological and phylogenetic evidence, we consider J. bapoensis as a member of the genus Pseudocalotes. As neither additional specimens nor genetic data of the nominate subspecies, P. k. kingdonwardi, are available, we cannot evaluate whether P. k. bapoensis deserves full species status at this time. Therefore, following previous published work (Mahony, 2010), we conservatively treat bapoensis as a subspecies of P. kingdonwardi, pending future analyses should specimens and genetic material of P. k. kingdonwardi become available.

Finally, it is important to note that our new classification aims to provide a suite of testable hypotheses for future phylogenetic work, and the new classifications for non-sampled species represent our best attempt at a more stable taxonomic structure. Many species, such as *Japalura sagittifera* and *Diploderma hamptoni*, are known from poorly surveyed regions, and lack vouchered specimens and genetic samples. Therefore, our new classification of these understudied taxa is based on the best available morphological data alone, which may be subject to convergence and warrants additional investigations. Additionally, future phylogenomic studies are needed to better resolve the higher level relationships of *Japalura s.l.* To facilitate future taxonomic work, we provide updated morphological diagnoses to all genera of the subfamily Draconinae below [modified from Smith (1935) and Zhao *et al.* (1999)].

	DIAGNOSTIC RET TO ALL GENERA OF SUBFAMILI DIAGONINAE	
	Femoral pores present	
	Femoral pores absent	
	Ribs much prolonged, supporting wing-like expansion Drace	
	Ribs not elongated into wing-like structure	
3a.	Four toes on hind limbs; males with prominent, retractable dewlaps	ł
3b.	Five toes on hind limbs	5
4a.	Enlarged scales present on posterior thigh, flank scales heterogeneousSitance	ı
4b.	Enlarged scales absent on posterior thigh, flank scales homogeneousSarado	ı
	Distinct bony supra-orbital arch present; globular protuberance present on snoutLyriocephalus	
5b.	Above structures all absent	3
6a.	Series of long, lanceolate spines present along medial line on gular; gular appendage projecting forward	ł
	beyond snout	3
6b.	No large spine series along medial line on gular; gular appendage absent or cannot extend forward	ł
	beyond snout	7
7a.	Distinct, elongated rostral appendage present (always in males, sometimes a mere tubercle in	ı
	females)	3
7b.	No rostral appendage, or appendage short, not elongated)
8a.	Dorsal body scales heterogeneousCeratophore	ı
	Dorsal body scales mostly homogenous	
9a.	Dorsal and nuchal crests tall; dorsal crest scales enlarged in distinct, large trianglesHarpesaurus	3
	Nuchal crest low; dorsal crest indistinct	
10a.	Three parallel longitudinal gular folds on each side of medial throat, curving and converging posteriorly	7
	into U-shape pattern	3
10b.	Gular folds not in above shape	L
	Dorsal body scales homogeneous	
	Dorsal body scales heterogeneous	
12a.	Lateral skin fold present from posterior jaw to shoulder on each side; tail long TAL >300% SVI	
	Bronchocela	
12b.	Lateral skin folds absent from posterior jaw to shoulder; tail relatively short, TAL <300% SVI	
	mostly	3
13a.	Body compressed dorsally; dorsal crest feeble Psammophilus	3
	Body not compressed dorsally; dorsal crests relatively well developed	
	Lateral gular sac pockets present; granular scutellation of gular sac Complicitus	
	Lateral gular sac pockets absent; scales of gular not in granular shape	
	Tail markedly swollen at base, flattened above, with dorsolateral keels formed by enlarged, angular	
	scales	
15b.	Tail not markedly swollen at basee, or swollen but rounded above	3
	Post-occipital and post-orbital spines present; tympanum exposed	
	Spines on dorsal head absent; tympanum exposed or concealed	
	Nuchal and dorsal crest short; tail long, TAL >185% SVL; lateral gular fold weakly developed	
	Oriocalotes	
17b.	Nuchal and dorsal crest much longer; tail short, TAL<180% SVL mostly; lateral gular fold strongly	
	developedAcanthosauro	
18a.	Enlarged dorsal body scales arranged in regular rows or ridges	
	Enlarged dorsal body scales arranged randomly	

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19a. Enlarged scales in dorsolateral ridges or rows only (except in <i>D. swinhonis</i> , which also in V-shape rows)
19b. Enlarged scales in both lateral and V-shaped ridges or rows
20a. Pair of enlarged, conical scales present on nape; head relatively wide and robust, HW >70% HL mostly
20b. Enlarged conical scale pair on nape absent; head relatively narrow and slender, HW <70% HL mostly
21a. Ventral body scales heterogeneous in size and shape
21b. Ventral body scales homogeneous in size and shape mostly
22a. Nuchal crest scales low, only slightly differentiated; body and limbs slender; neck distinct and long 23
22b. Nuchal crest scales tall, elongated in lanceolate spines; body and limbs relatively robust; neck robust and relatively short
23a. Toe V not longer than Toe I; males usually with distinctively developed dewlaps extending to anterior venter
23b. Toe V longer than Toe I; males with much smaller lateral gular folds
24a. Suborbital scales in multiple rows, subequal in size; dorsal scales fine, smaller than ventrals; distinct
transverse gular fold present; nuchal crest separated from nuchal scales by one or more rows of enlarged, supporting scales
24b. Suborbital scales in singular row, or multiple rows but one row distinctively enlarged; dorsal scales
larger or smaller than ventrals; transverse gular fold absent or indistinct; nuchal crest not separated from nuchals by enlarged, supporting scales
25a. Majority of dorsal body scales smaller than ventrals
25b. Majority of dorsal body scales larger than ventrals
26a. Lateral gular fold strongly developed with rounded tip, extending to anterior region of venter when folded; dorsal body scales muricate
26b. Lateral gular fold less developed without an obvious tip, not extending beyond throat when folded; dor- sal body scales more heterogeneous with rhombic scales
27a. Head tall, robust, distinctively raised on occipital region with distinct cranial ridges Dendragama
27b. Head more compressed dorsally, much more slender, not distinctively raised on occipital region
28a. Tympana exposed
28b. Tympana concealed
29a. Tail prehensile, short, TAL <135% SVL; tympanum concealedCophotis
29b. Tail not prehensile, long, TAL >160% SVL; tympanum exposed
30a. Dorsal body scales less heterogeneous; dorsal body scales more rounded; tail short, TAL <175% SVL
30b. Dorsal body scales more heterogeneous; dorsal body scales more elongated; tail long, TAL >200% SVL

SPECIES-LEVEL RELATIONSHIPS AND TAXONOMY

Japalura

Within Japalura, relationships of sampled taxa are largely resolved (Fig. 2). Interestingly, despite morphological similarity between J. kumaonensis and J. tricarinata (Wang et al., 2018b), J. kumaonensis is most closely related to J. variegata (0.99/69; Fig. 2). As many of the Himalayan regions lack detailed surveys for Japalura diversity, including north-east India, Bhutan, Nepal, and north-western Bangladesh, many congeners are still poorly studied. In fact, several recognized species are known from a few vouchered specimens only with no genetic samples (e.g. J. dasi, J. major and J. sagittifera). Not only may these remote regions harbour undescribed diversity, but also, given the wide variation in ornamentation observed among populations (Bhosale, Das & Manthey, 2013; Wang et al. 2018a), understudied taxa such as J. andersoniana, J. kumaonensis, J. tricarinata and J. variegata may represent species complexes and contain cryptic diversity. Future research should focus on the collection of additional, vouchered genetic data of these poorly studied lineages and close examination of Japalura populations across the Himalayan region.

Cristidorsa

For congeners from Myanmar, both our morphological and molecular data support the previous hypothesis that

Myanmar populations of Cristidorsa cf. planidorsata were misidentified, and are, in fact, C. otai. When discussing the taxonomy of Myanmar populations of C. planidorsata (considered as J. planidorsata), Mahony (2009) stated specifically that the only available specimen (juvenile) of C. planidorsata examined from Myanmar resembled the morphology of *C. otai*. However, given the lack of access to adult specimens from Myanmar, Mahony (2009) conservatively treated the Myanmar population as C. cf. planidorsata, but called for further investigation on the taxonomic status of these populations. Based on our data, the individuals of previously identified C. cf. planidorsata from Myanmar are not monophyletic with respect to the topotypic C. planidorsata, but instead, form a monophyletic group with topotypic C. otai from north-east India (Fig. 2, clade G). In addition, the morphological characteristics of the Myanmar populations match the diagnoses of C. otai and differ from C. planidorsata, including having flattened and posteriorly pointing conical scales on the temporal region of the head, shorter body lengths TRL 41.8-47.4% SVL, longer tails TAL 154.6-190.8% SVL, and shorter lip-stripes ending anterior to the limb insertion (Mahony, 2009). Therefore, we diagnose the Myanmar populations of C. cf. planidorsata as C. otai, and recommend that records of the former species in Myanmar be corrected to represent the latter.

Diploderma

Recently, Ananjeva et al. (2007) described a new species of Japalura, J. ngoclinensis, on the basis of three female specimens collected from the central highlands of Vietnam, far outside of the range of Japalura s.l. recognized at the time. Although there are no genetic data available for the species, we can assign it to the genus Diploderma based on our revised morphological diagnosis. The issue that remains is in regard to the taxonomic validity of D. ngoclinense. Despite the fact that the type specimens of the species are morphologically most similar to *D. splendidum*, a brief comparison was made to the latter species using one morphological characteristic only (Ananjeva et al., 2017). The authors argued that the new species differs from D. splendidum by the absence of a transverse gular fold (Ananjeva et al., 2017). However, based on the photographs of the type series in the original description, all type specimens of D. ngoclinense do have a shallow transverse gular fold (particularly distinct on the paratype, VNMH 3110) (Ananjeva et al., 2017: figs 2-4), identical to that seen in preserved specimens of D. splendidum [USNM 35522 (holotype), CIB 2588, 2591, 2596, 72468, 72469 from Chongqin, China]. Based on the available data, the only differentiating characteristic between D. ngoclinense from D. splendidum is the mid-dorsal scale count, which is higher in *D. ngoclinense* (54–56 vs. 44–52 in *J. splendidum*; Ananjeva *et al.*, 2017). However, such a difference is based on a small sample size of three specimens of *D. ngoclinense*, and mid-dorsal scale counts are known to vary in *Diploderma* (Ota, 1989a; Manthey *et al.*, 2012; Wang et al., 2016, 2018). In addition to the suspicious distribution of *D. ngoclinense* (isolated and 1000 km south of the most southern-known range of the genus; Ananjeva *et al.*, 2017), we question the taxonomic validity of *D. ngoclinense*. Future field confirmation, detailed morphological comparisons and phylogenetic analyses are needed to validate the taxonomic status and distribution of this species.

Historically, almost all *Diploderma* diversity in central and south-west China was thought to be a single species, D. flaviceps (Zhao et al., 1999). Although numerous new species have been described from the D. flaviceps complex, including D. batangense, D. brevicaudum, D. flaviceps, D. iadinum, D. laeviventre, D. micangshanense, D. vela, D. yulongense and D. zhaoermii (Song, 1987; Li et al., 2001; Gao & Hou, 2002; Manthey et al., 2012; Wang et al., 2015, 2016, 2017), many outlier records of D. flaviceps in south-west China have not been examined, some of which may represent either cryptic diversity in the genus or misidentifications (Wang et al., 2016). For example, the genetic sequence data of an MVZ specimen from Wenchuan, north-west Sichuan Province, China (MVZ 216622) was recorded as D. flaviceps upon deposition in GenBank, and subsequently has been used for numerous phylogenetic studies (GenBank accession no. AF128500; Macey et al., 2000; Schulte et al., 2004; Zug et al., 2006; Pyron et al., 2013; Grismer et al., 2016b). However, according to the museum record, the collection locality of this specimen is east of Wenchuan, Sichuan Province, China, which is geographically proximate to the type locality of a different congener, D. zhaoermii (Gao & Hou, 2002). Furthermore, our phylogenetic analyses recover this individual of D. cf. flaviceps as nested within topotypic material of D. zhaoermii (Fig. 2). Additionally, as D. flaviceps is not recognized to occur outside the upper Dadu River Valley and its direct tributaries (Manthey et al., 2012; Wang et al., 2016), we suspect that the Diploderma specimen (MVZ 216622, genetic sequence on GenBank AF128500) was misidentified, and it represents the species D. zhaoermii instead. Poorly studied groups and continuous taxonomic revisions create problems such as this for museum and online databases, and numerous changes to the taxonomy of the D. flavi*ceps* complex in China are no exception. We propose major natural history collections update the taxonomy of Japalura s.l., particularly members of the D. flaviceps complex from south-west China.

CONCLUSIONS

Utilizing multilocus and morphological datasets, we have addressed a long-standing issue concerning the paraphyletic nature of Japalura s.l. reclassifying the genus into four genera. Revision of this widespread group further clarifies the phylogenetic relationships among genera in the subfamily Draconinae. However, several questions remain that were beyond the scope of this study. First, interspecific phylogenetic relationships of each of the four genera within Japalura s.l. are still not well understood. Second, without a fully resolved phylogeny of these major clades within Japalura s.l. it will remain difficult to test finer scale biogeographic hypotheses concerning some of the most interesting landscapes on earth, including the trans-Himalayan Mountains and the Hengduan Mountain Region in south-west China. Third, genetic samples remain unavailable for many genera within Draconinae, including Harpesaurus, Mictopholis, Oriocalotes and Pseudocophotis. Finally, focused phylogenetic studies are still needed to resolve species-level relationships in other genera (i.e. Acanthosaura, Gonocephalus and Pseudocalotes). Together, these problems perpetuate questions about interspecific relationships and the taxonomic validity of some species recognized currently. As such, Asian Dragon Lizards continue to represent an enigmatic vertebrate radiation in need of a broad spectrum of studies aimed at better understanding everything from species-level diversity to evolutionary patterns of diversification.

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REFERENCES

- Ananjeva NB, Guo X, Wang Y. 2011. Taxonomic diversity of agamid lizards (Reptilia, Sauria, Acrodonta, Agamidae) from China: a comparative analysis. Asian Herpetological Research 2: 117–128.
- Ananjeva NB, Orlov NL, Nguyen TT. 2017. A new species of Japalura (Agamidae: Lacertilia: Reptilia) from central highland, Vietnam. Asian Herpetological Research 8: 14–21.
- Annandale N. 1905. Contributions to the Oriental herpetology II.—Notes on the Oriental lizards in the Indian Museum, with a list of the species recorded from British India and Ceylon. Journal and Proceedings of the Asiatic Society of Bengal 1: 81–93.
- Annandale N. 1907. Lacertilia. In: Boulenger GA, Annandale N, Wall F, Regan CT, eds. Report on a collection of batrachia, reptiles and fish from Nepal and the Western Himalayas. Records of the Indian Museum 1: 149–158.
- Bhosale H, Das A, Manthey U. 2013. New locality records and color variations of *Japalura andersoniana* Annandale, 1905 (Sauria: Agamidae: Draconinae). *Sauria* 35: 55–60.
- Blyth E. 1854. Notices and descriptions of various reptiles, new or little-known. *Part I. Journal of Asiatic Society of Bengal* 21: 639–655.
- **Boulenger GA. 1885.** Catalogue of lizards in the British Museum (Natural History) I. Geckonidae, Eublepharidae, Uroplatidae, Pygopodidae, Agamidae. London: Printed by the order of the Trustees.
- **Boulenger GA. 1888.** An account of the reptiles and batrachians obtained in Tenasserim by M.L. Fea. *Annali del Museo civico di storia naturale di Genova* **2:** 474–486.

- **Boulenger GA. 1906.** Descriptions of new reptiles from Yunnan. *The Annals and Magazine of Natural History* **7:** 567–568
- **Boulenger GA. 1918.** Description of a new lizard of the genus Acanthosaura from Yunnan. The Annals and Magazine of Natural History **9:** 162.
- Boulenger GA. 1920. Results of an expedition to Korinchi Peak 12.400 ft, Sumatra. III. Reptiles & batrachians collected in Korinchi, West Sumatra by Messrs. H.C. Robinson & C.B. Kloss. *Journal of the Federated Malay States Museums* 8: 285–296.
- Cai B, Wang Y, Chen Y, Li J. 2015. A revised taxonomy of Chinese reptiles. *Biodiversity Science* 23: 365–382.
- Che J, Zhou W, Hu J, Yan F, Papenfuss T, Wake D, Zhang Y. 2010. Spiny frogs (Paini) illuminate the history of the Himalayan region and Southeast Asia. *Proceedings of the National Academy of Sciences of the United States of America* 107: 13765–13770.
- Crottini A, Dordel J, Köhler J, Glaw F, Schmitz A, Vences M. 2009. A multilocus phylogeny of Malagasy scincid lizards elucidates the relationships of the fossorial genera Androngo and Cryptoscincus. Molecular Phylogenetics and Evolution 53: 345–350.
- Darriba D, Taboada GL, Doallo R, Posada D. 2012. jModel-Test 2: more models, new heuristics and parallel computing. *Nature Methods* 9: 772.
- Deepak V, Giri V, Asif M, Dutta SK, Vyas R, Zambre AM, Bhosale H, Karanth KP. 2016. Systematics and phylogeny of *Sitana* (Reptilia: Agamidae) of Peninsular India, with description of one new genus and five new species. *Contribution to Zoology* 85: 67–111.
- Denzer W, Manthey U, Wagner P, Böhme W. 2016. A critical review of Hoser's writings on Draconinae, Amphibolurinae, *Laudakia*, and Uromastycinae (Squamata: Agamidae). *Bonn Zoological Bulletin* 64: 117–138.
- Edgar RC. 2004. MUSCLE: multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Research* 32: 1792–1797.
- Esselstyn JA, Garcia HJD, Saulog MG, Heaney LR. 2008. A new species of *Desmalopex* (Pteropodidae) from the Philippines, with a phylogenetic analysis of the Pteropodini. *Journal of Mammalogy* 89: 815–825.
- Gao ZF, Hou M. 2002. Description of a new *Japalura* species from western Sichuan Province, China. *Sichuan Journal of Zoology* 21: 3–5.
- **Gray JE. 1853.** Descriptions of some undescribed species of reptiles collected by Dr Joseph Hooker in the Khassia Mountains, East Bengal, and Sikkim Himalaya. *The Annals and Magazine of Natural History* **12:** 386–392.
- Grismer LL, Quah ES, Wood PL, Anuar S, Muin A, Davis HR, Murdoch ML, Grismer JL, Cota M, Cobos AJ. 2016a. Dragons in the mist: three new species of *Pseudocalotes* Fitzinger (Squamata: Agamidae) from the sky island archipelago of Peninsular Malaysia. *Zootaxa* **4136**: 461–490.
- Grismer JL, Schulte JA II, Alexander A, Wagner P, Travers SL, Buehler MD, Welton LJ, Brown R. 2016b. The Eurasian invasion: phylogenetic data reveal multiple Southeast Asian origins for Indian Dragon Lizards. BMC Evolutionary Biology 16: 1–11.

- Guindon S, Gascuel O. 2003. A simple, fast and accurate method to estimate large phylogenies by maximum-likelihood. *Systematic Biology* **52**: 698–704.
- Harvey MB, Shaney K, Hamidy A, Kurniawan N, Smith EN. 2017. A new species of *Pseudocalotes* (Squamata: Agamidae) from the Bukit Barisan Range of Sumatra with an Estimation of its phylogeny. *Zootaxa* 4276: 215–232.
- Honda M, Ota H, Kobayashi M, Nabhitabhata J, Yong H, Sengoku S, Hikida T. 2000. Phylogenetic relationships of the family Agamidae (Reptilia: Iguania) inferred from mitochondrial DNA sequences. *Zoological Science* 17: 527–537.
- Hu S, Zhao E, Jiang Y, Fei L, Ye C, Hu Q, Huang Y, Tian W. 1987. *Amphibia-Reptilia of Xizang*. Beijing: Science Press.
- Inger RF. 1960. A review of the agamid lizards if the genus Phoxophrys Hubrecht. *Copeia* 3: 221–225.
- Kaiser HF. 1960. The application of electronic computers to factor analysis. *Education and Psychological Measurement* 20: 141–151.
- Kästle W, Schleich HH. 1998. Studies on the systematics and biology of the genus Japalura (Sauria: Agamidae). Notes on comparative ethology and taxonomy of the genus Japalura. In: Kästle W, Schleich HH, eds. The contributions to the herpetology of South Asia (Nepal, India). Wuppertal: Fuhlrott Museum, 233–246.
- Li C, Deng Q, Wu Y, Wang Y. 2001. A new species of Japalura from Sichuan (Agamidae Gray Japalura). Journal of Sichuan Teachers College (Nature Science) 22: 329–331.
- Li PP, Zhao E, Dong B. 2010. Amphibians and Reptiles of *Tibet*. Beijing: Science Press.
- Macey JR, Schulte JA 2nd, Larson A, Ananjeva NB, Wang Y, Pethiyagoda R, Rastegar-Pouyani N, Papenfuss TJ. 2000. Evaluating trans-tethys migration: an example using acrodont lizard phylogenetics. *Systematic Biology* **49**: 233–256.
- **Mahony S. 2009.** A new species of *Japalura* (Reptilia: Agamidae) from northeast India with a discussion of the similar species *Japalura sagittifera* Smith, 1940 and *Japalura planidorsata* Jerdon, 1870. *Zootaxa* **2212:** 41–61.
- Mahony S. 2010. Systematic and taxonomic revaluation of four little known Asian agamid species, *Calotes kingdonwardi* Smith, 1935, *Japalura kaulbacki* Smith, 1937, *Salea kakhienensis* Anderson, 1879 and the monotypic genus *Mictopholis* Smith, 1935 (Reptilia: Agamidae). *Zootaxa* 2514: 1–23.
- Manthey U. 2010. Agamid lizards of Southern Asia Draconinae 2, Leiolepidinae. Frankfurt: Chimaira.
- Manthey U, Denzer W, Hou M, Wang X. 2012. Discovered in historical collections: two new *Japalura* species (Squamata: Sauria: Agamidae) from Yulong Snow Mountains, Lijiang Prefecture, Yunnan, PR China. *Zootaxa* **3200**: 27–48.
- Marshall DC. 2010. Cryptic failure of partitioned Bayesian phylogenetic analyses: lost in the land of long trees. *Systematic Biology* **59**: 108–117.
- **Moody SM. 1980.** Phylogenetic relationship and historical bigeographical relationships of the genera in the family Agamidae (Reptilia: Lacertilia). Unpublished Doctoral Dissertation, University of Michigan.
- Nylander JAA, Wilgenbusch JC, Warren DL, Swofford DL. 2008. AWTY(are we there yet?): a system for graphical

exploration of MCMC convergence in Bayesian phylogenetic inference. *Bioinformatics* **24:** 581–583.

- **Ota H. 1989a.** A new species of *Japalura* (Agamidae: Lacertilia: Reptilia) from Taiwan. *Copeia* **1989**: 569–576.
- **Ota H. 1989b.** Japalura brevipes Gressitt (Agamidae: Reptilia), a valid species from high altitude area of Taiwan. *Herpetologica* **45:** 55–60.
- Pope CH. 1935. The reptiles of China: turtles, crocodilians, snakes, lizards. natural history of Central Asia. Vol. X. New York: American Museum of Natural History.
- **Pyron RA, Burbrink FT, Wiens JJ. 2013.** A phylogeny and revised classification of Squamata, including 4161 species of lizards and snakes. *BMC Evolution* **13**: 1–53.
- Rambaut A, Suchard M, Drummond A. 2013. *Tracer v. 1.6.* Available at: http://tree.bio.ed.ac.uk/software/tracer
- Rao D, Vindum JV, Ma X, Fu M, Wilkinson JA. 2017. A new species of *Japalura* (Squamata, Agamidae) from the Nu River Valley in Southern Hengduan Mountains, Yunnan, China. *Asian Herpetological Research* 8: 86–95.
- Ronquist F, Huelsenbeck JP. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* 19: 1572–1574.
- Schleich HH, Kästle W. 2002. Amphibians and reptiles of Nepal. Ruggell: Ganther Verlag KG.
- Schmidt KP. 1927. Notes on Chinese reptiles. Bulletin of the American Museum of Natural History 54: 467–551.
- Schulte JA II, Vindum JV, Win H, Thin T, Lwin KS, Shein AK. 2004. Phylogenetic relationships of the genus *Ptyctolaemus* (Squamata: Agamidae), with a description of a new species from the Chin Hills of western Myanmar. *Proceedings of the California Academy of Sciences* 55: 222–247.
- Siler CD, Oliveros CH, Santanen A, Brown RM. 2013. Multilocus phylogeny reveals unexpected diversification patterns in Asian wolf snakes (genus Lycodon). Zoologica Scripta 42: 262–277.
- Smith MA. 1935. The fauna of British India, Ceylon and Burma. Reptilia and Amphibia. Vol. II Sauria. London: Taylor & Francis.
- Smith MA. 1937. Description of a new species of agamid lizard from upper Burma. *Journal of Bombay Natural History Society* 39: 755.
- Song M. 1987. Survey of the reptiles of southern Shaanxi. Acta Herpetologica Sinica 6: 59–64.
- Stamatakis A, Hoover P, Rougemont J. 2008. A rapid bootstrap algorithm for the RAxML Web servers. Systematic Biology 57: 758–771.
- Stamatakis A. 2014. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics 30: 1312–1313.
- Stejneger L. 1924. Herpetological novelties from China. Occasional Papers of the Boston Society of Natural History 5: 119–121.
- Stuart-Fox DM, Ord TJ. 2004. Sexual selection, natural selection and the evolution of dimorphic coloration and

ornamentation in agamid lizards. *Proceedings of the Royal* Society of London B: Biological Sciences **271:** 2249–2255.

- Stuart-Fox D, Owens IP. 2003. Species richness in agamid lizards: chance, body size, sexual selection or ecology? *Journal of Evolutionary Biology* 16: 659–669.
- **Swofford DL. 1999.** *PAUP*4.0. Phylogenetic Analysis Using Parsimony (*and Other Methods).* Sunderland: Sinauer Associates.
- Uetz P, Hošek J. 2017. The reptiles database. Available at: http://www.reptile-database.org
- Wang K, Jiang K, Pan G, Hou M, Siler CD, Che J. 2015. A new species of *Japalura* (Squamata: Sauria: Agamidae) from Eastern Tibet, PR China. Asian Hepetological Research 6: 159–168.
- Wang K, Jiang K, Zou DH, Yan F, Siler CD, Che J. 2016. Two new species of *Japalura* (Squamata: Agamidae) from the Hengduan Mountain Range, China. *Zoological Research* 37: 41–56.
- Wang K, Ren J, Jiang K, Yuan Z, Che J, Siler CD. 2017. Rediscovery of the enigmatic mountain dragon, *Japalura yulongensis* (Reptilia: Sauria: Agamidae), with notes on its natural history and conservation. *Zootaxa* 4318: 351–363.
- Wang K, Jiang K, Wang YF, Poyarkov NA Jr, Che J, Siler CD. 2018a. Discovery of *Japalura chapaensis* Bourret, 1937 (Reptilia: Squamata: Agamidae) from Southeast Yunnan Province, China. *Zoological Research* 39: 105–113.
- Wang K, Jiang K, Deepak V, Abhijit D, Hou M, Che J, Siler CD. 2018b. On the occurrences of *Japalura kumaonensis* and *J. tricarinata* (Reptilia: Sauria: Draconinae) in China, with a diagnostic key to *Japalura sensu lato* from the Himalaya. *Herpetologica* 74: 181–190.
- Werner F. 1904. Beschreibung neuer Reptilien aus den Gattungen Acanthosaura, Calotes, Gastropholis und Typhlops. Zoologischer Anzeiger 27: 461–464.
- Whiting AS, Bauer AM, Sites JW Jr. 2003. Phylogenetic relationships and limb loss in sub-Saharan African scincine lizards (Squamata: Scincidae). *Molecular Phylogenetics and Evolution* 29: 582–598.
- Yang D, Rao D. 1992. The identity of reptiles and their origination and evolution in southeastern Asia and Yunnan Province of China. *Zoological Research* 13: 101–108.
- Yang D, Su C, Li S. 1979. New species and new subspecies of amphibians and reptiles from Gaoligong Shan, Yunnan. Acta Zootaxonomica Sinica 4: 185–188.
- Yang D, Rao D. 2008. *Amphibia and Reptilia of Yunnan*. Kunming: Yunnan Publishing Group Corporation.
- Zhao E, Adler K. 1993. *Herpetology of China*. Oxford: Society for the Study of Amphibians and Reptiles.
- Zhao E, Zhao K, Zhou K. 1999. Fauna Sinica, Reptilia, Vol. 2: Squamata, Lacertilia. Beijing: Science Press.
- Zug GR, Brown HHK, Schulte JA II, Vindum JV. 2006. Systematics of the garden lizards, *Calotes versicolor* group (Reptilia, Squamata, Agamidae) in Myanmar: central dry zone populations. *Proceedings of the California Academy of Sciences* 57: 35–68.

SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article at the publisher's web-site.

Appendix S1. Summary of specimens corresponding to genetic samples included in the study, including GenBank accession numbers and vouchered specimen information. CAS = California Academy of Sciences, USA; MVZ = Museum of Comparative Zoology, USA; OMNH = Sam Noble Oklahoma Museum of Natural History, USA; KIZ = Kunming Institute of Zoology, Chinese Academy of Sciences, China; ZISP = Zoological Institute, St. Petersburg, Russia; WHT = Wildlife Heritage Trust, Sri Lanka; BNHS = Bombay Natural History Society, India; TNHC = Texas Memorial Museum, USA; LSUMZ = Louisiana Museum of Natural History, USA; USNM = National Museum of Natural History, USA; FMNH = Field Museum of Natural History, USA; CES = Center for Ecological Sciences, Indian Institute of Science, India. "/" indicates only tissue samples were taken without voucher specimens. **Appendix S2.** Genbank sequences used in the phylogenetic analyses.

Appendix S3. Specimens examined. Museum abbreviations include: Museum of Natural History (BMNH), Kensington, London, UK; Museum of California Academy of Sciences (CAS), San Francisco, CA, USA; Center for Ecological Sciences, Indian Institute of Science (CESG), Bengaluru, Karnataka, India; Chengdu Institute of Biology, Chinese Academy of Sciences (CIB); Kunming Institute of Zoology, Chinese Academy of Sciences (KIZ), Kunming, Yunnan, China; University of Kansas Biodiversity Institute (KU), Lawrence, KS, USA; Field Museum of Natural History (FMNH), Chicago, IL, USA; and Museum of Comparative Zoology (MCZ), Cambridge, MA, USA. **Appendix S4.** Specimens measured for the principal component analyses of continuous morphological characteristics. Museum abbreviations as in caption of Appendix S3.

Appendix S5. Summary statistics for the principal component analyses of continuous morphological characteristics for *Japalura s.l.* and morphologically similar Draconinae genera. Abbreviations are listed in Material and Methods.